Ocular Differential Diagnosis

*Eighth Edition*

Frederick Hampton Roy, M.D., F.A.C.S.

*Little Rock, Arkansas*
To Mary Michelle
To my children:
Nichols, Robert, Kimberly, Frederick, Jr., Charles, and Helena
To Dr. Arlington Krause,
molder and questioner in my early formative academic life
To Dr. Philip Lewis and Dr. Roger Hiatt
for guidance and direction
Contents

Preface ix
How to Use This Book xi

I. Regional Signs and Symptoms
1. Orbit 3
2. Lids 45
3. Lacrimal System 110
4. Extraocular Muscles 127
5. Conjunctiva 183
6. Globe 220
7. Sclera 231
8. Cornea 241
9. Intraocular Pressure 305
10. Anterior Chamber 329
11. Pupil 344
12. Iris 364
13. Lens 386
14. Vitreous 421
15. Retina 433
16. Choroid 526
17. Optic Nerve 559
18. Visual-Field Defects 604

II. General Signs and Symptoms
19. Visual Disturbance 619
20. Visual Complaint 644
21. Head Position 665
Subject Index 669
Preface

The first edition of *Ocular Differential Diagnosis* was published in 1972, and various editions have been translated in Spanish, Turkish, Chinese, Portuguese, and Italian. All previous editions have been my work. Dr. Gonzolo Murillo of La Paz, Bolivia, helped in edition four and five with the diagnostic decision tables. In edition six, section editors helped to standardize the language and made helpful suggestions. I feel this is the best edition of *Ocular Differential Diagnosis* thus far. This text would not have been possible without the superb efforts of Renee Tindall, Dr. Fernando Murillo, Angie Brown, and Dr. Kae Chatman.

The *Ocular Differential Diagnosis* book provides comprehensive lists of causes for symptoms or findings. Frequently more information is needed and hopefully *Ocular Syndromes and Systemic Diseases* will furnish additional information to make a better diagnosis. I hope the ophthalmologist and optometrist using this book will bring any errors in this edition to my attention.

Frederick Hampton Roy, M.D.
Little Rock, Arkansas
How to Use This Book

This book can be used easily and quickly by following the directions presented below.

1. If the sign or symptom relates to a particular region of the eye, turn to the table of contents preceding this page to find the number of the page on which listings of the signs and symptoms pertaining to the specific region begins. This latter page (or those immediately following) will refer the user to that (or those) on which the various causes of the condition are listed. For example, let us assume that the patient has pigmentation of the cornea. The table of contents on page vii shows that the cornea section begins on page 241. Turning to page 241 the user finds references to page 248 on which the causes of corneal pigmentation are listed according to type. In the subject index, this topic is listed as Cornea, pigmentation of, 248.

2. If the symptom, such as binocular diplopia or night blindness, does not relate to a particular region of the eye, look for it either in the subject index at the back of the book or under General Signs and Symptoms beginning on page 619.

   Various features of a disease may be crosschecked. For instance, a "pulsating exophthalmos with orbital bruit and conjunctival edema" may be sought under orbit, page 3, where the user of the book is referred to exophthalmos, page 4, and orbital bruit, page 31, and under conjunctiva, page 183, where the user is referred to conjunctival edema, page 202. The terms "exophthalmos," "orbital bruit" (under orbit, bruit of) and "conjunctival edema" (under conjunctiva, edema of) may also be found in the subject index. Terms such as "secondary glaucoma" are indexed under the noun, e.g., glaucoma, secondary.

3. Following some of the differential diagnosis lists are diagnostic decision tables. These tables list the history, physical signs, and laboratory tests that differentiate each possible diagnosis. These can be identified in the subject index because they are followed by a \( t \).
Ocular Differential Diagnosis
Eighth Edition

PART I
Regional Signs and Symptoms
# Orbit

## CONTENTS

<table>
<thead>
<tr>
<th>Topic</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pseudoproptosis</td>
<td>4</td>
</tr>
<tr>
<td>Exophthalmos</td>
<td>4</td>
</tr>
<tr>
<td>Syndromes and diseases associated with exophthalmos</td>
<td>8</td>
</tr>
<tr>
<td>Specific exophthalmos</td>
<td>11</td>
</tr>
<tr>
<td>Diagnostic tables</td>
<td>16</td>
</tr>
<tr>
<td>Up to one year</td>
<td>16</td>
</tr>
<tr>
<td>One to five years</td>
<td>17</td>
</tr>
<tr>
<td>Five to ten years</td>
<td>19</td>
</tr>
<tr>
<td>Ten to thirty years</td>
<td>21</td>
</tr>
<tr>
<td>Thirty to fifty years</td>
<td>22</td>
</tr>
<tr>
<td>Fifty to seventy years</td>
<td>24</td>
</tr>
<tr>
<td>More than seventy years</td>
<td>26</td>
</tr>
<tr>
<td>Pulsating exophthalmos</td>
<td>28</td>
</tr>
<tr>
<td>Recurrent exophthalmos</td>
<td>29</td>
</tr>
<tr>
<td>Enophthalmos</td>
<td>15</td>
</tr>
<tr>
<td>Intraorbital calcifications</td>
<td>31</td>
</tr>
<tr>
<td>Orbital bruit</td>
<td>31</td>
</tr>
<tr>
<td>Diagnostic table</td>
<td>32</td>
</tr>
<tr>
<td>Orbital emphysema</td>
<td>33</td>
</tr>
<tr>
<td>Orbital pain</td>
<td>33</td>
</tr>
<tr>
<td>Shallow orbits or diminished orbital volume</td>
<td>34</td>
</tr>
<tr>
<td>Pseudohypertelorism</td>
<td>35</td>
</tr>
<tr>
<td>Hypertelorism</td>
<td>35</td>
</tr>
<tr>
<td>Hypotelorism</td>
<td>38</td>
</tr>
<tr>
<td>Deep-set eyes</td>
<td>38</td>
</tr>
<tr>
<td>Prominent supraorbital ridges</td>
<td>38</td>
</tr>
<tr>
<td>Osteolysis of bony orbit</td>
<td>39</td>
</tr>
<tr>
<td>Fossa formation of orbit</td>
<td>39</td>
</tr>
<tr>
<td>Subperiosteal orbital hemorrhage</td>
<td>39</td>
</tr>
<tr>
<td>Orbital hemorrhage</td>
<td>40</td>
</tr>
<tr>
<td>Expansion of orbital margins</td>
<td>40</td>
</tr>
<tr>
<td>Hypertrophy of orbital bones</td>
<td>40</td>
</tr>
<tr>
<td>Expansion of optic canal</td>
<td>41</td>
</tr>
<tr>
<td>Small optic canals</td>
<td>41</td>
</tr>
<tr>
<td>Erosion of optic canal</td>
<td>42</td>
</tr>
<tr>
<td>Enlargement of superior orbital fissure</td>
<td>42</td>
</tr>
<tr>
<td>Narrowing of superior orbital fissure</td>
<td>43</td>
</tr>
<tr>
<td>Small orbit</td>
<td>43</td>
</tr>
<tr>
<td>Large orbit</td>
<td>43</td>
</tr>
<tr>
<td>Hematic orbital cysts</td>
<td>43</td>
</tr>
</tbody>
</table>
**PSEUDOPROPTOSIS (APPEARANCE OF EXOPHTHALMOS)**

1. Asymmetry of bony orbits
2. Congenital cystic eyeball
3. Contralateral enophthalmos (see p. 15)
4. Facial asymmetry as progressive facial hemiatrophy (Parry-Romberg syndrome)
5. Harlequin orbit (shallow orbit with arched superior and lateral wall) as with hypophosphatasia
6. Hypoplastic supraorbital ridges as in trisomy (Edward syndrome)
7. Retraction of upper lid as with thyroid disease
8. Slight blepharoptosis as with Homer syndrome of contralateral eye
9. Shallow orbit as in Crouzon disease (craniofacial dysostosis)
10. Unilateral congenital glaucoma
11. Unilateral high-axial myopia
12. Unilateral secondary glaucoma resulting from ocular trauma during childhood


**EXOPHTHALMOS**

1. Drugs, including the following:
   - adrenal cortex injection
   - beclomethasone
   - carbimazole
   - cocaine
   - cortisone
   - desoxycorticosterone
   - dexamethasone
   - dextrothyroxine
   - fludrocortisone
   - fluorometholone
   - flu-prednisolone

2. Inflammation
   - A. Acute-orbital cellulitis
   - B. Acute suppurative-mucormycosis (diabetic or debility)
   - C. Allergic fungal sinusitis
   - D. Benign lymphoepithelial lesion (Mikulicz disease)
   - E. Chronic (nongranulomatous-pseudotumor)
   - F. Chronic (granulomatous-tuberculosis, sarcoid (Schaumann syndrome), syphilis (lues), parasites, aspergillosis)
   - G. Relapsing polychondritis

3. Injuries
A. Foreign body
B. Orbital hemorrhage
C. Orbital roof fracture
D. Secondary carotid cavernous sinus fistula
E. Thermal burns

4. Systemic disease
A. Acute intracranial hypertension
B. Amyloidosis (Lubarsch-Pick syndrome)
C. Chloroma
D. Cretinism (hypothyroidism)
E. Hydrocephalus and ventriculoperitoneal syndrome
F. Hypervitaminosis A
G. Hypophosphatasia (phosphoethanolaminuria)
H. Thyroid disorder
I. Myasthenia gravis (Erb-Goldflam syndrome)
J. Obesity

5. Tumors
A. Cartilaginous tumors
   (1) Cartilaginous hamartoma
   (2) Chondroma
   (3) Chondrosarcoma
   (4) Mesenchymal chondrosarcoma
B. Cystic lesions
   (1) Colobomatous cyst
   (2) Dermoid cyst
   (3) Hematocele
   (4) Hydatid cyst
   (5) Meningocele and meningoencephalocele
   (6) Mucocele
   (7) Optic nerve sheath cyst
   (8) Simple epithelial cyst
   (9) Teratoma
C. Fibrocytic tumors
   (1) Fibroma
   (2) Fibrosarcoma
   (3) Fibrous histiocytoma
   (4) Juvenile fibromatosis
   (5) Nodular fasciitis
D. Histiocytic lesions
   (1) Others
      a. Juvenile xanthogranuloma (JXG, nevoxanthoendothelioma)
      b. Sinus histiocytosis with massive lymphadenopathy
   (2) Systemic histiocytoses (histiocytosis X) (Hand-Schüller-Christian disease)
E. Inflammatory pseudotumor of orbit
   (1) Ectopic cerebellar tissue in orbit
(2) Local, such as fungus or foreign body
(3) Systemic such as sarcoidosis syndrome (Schaumann syndrome) or collagen disease

(4) Unknown cause

F. Lacrimal gland (fossa) lesions
(1) Epithelial tumors
   a. Adenoid cystic carcinoma
   b. Mucoepidermoid carcinoma
   c. Pleomorphic adenocarcinoma (malignant mixed tumor)
   d. Pleomorphic adenoma (benign mixed tumor)

(2) Nonepithelial lesions
   a. Infectious
   b. Inflammatory
   c. Lymphoid and leukemia
   d. Systemic (sarcoid)

G. Lipocytic and myxoid tumors
(1) Lipoma
(2) Liposarcoma
(3) Myxoid liposarcoma
(4) Myxoma

H. Lymphoid tumors and leukemias (excluding lacrimal gland lesions)
(1) Benign reactive lymphoid hyperplasia
(2) Burkitt lymphoma
(3) Lymphoblastic leukemia
(4) Myelogenous leukemia (granulocytic sarcoma)
(5) Non-Hodgkin lymphoma

I. Metastatic tumors of the orbit
(1) Malignant melanoma of skin
(2) Neuroblastoma (child)
(3) Other sites such as Ewing sarcoma
(4) Primary in breast (adult female)
(5) Primary in lung (adult male)
(6) Primary in prostate (adult male)

J. Nonepithelial lesions
(1) Benign reactive lymphoid hyperplasia
(2) Inflammatory pseudotumors (dacyroadenitis)
(3) Lymphoma
(4) Plasmacytoma

K. Optic nerve and meningeal tumors
(1) Juvenile pilocytic astrocytoma (optic nerve glioma)
(2) Meningioma
   a. Primary optic nerve sheath
   b. Secondary
(3) Malignant optic nerve glioma

L. Osseous and fibroosseous tumors
(1) Aneurysmal bone cyst
(2) Benign osteoblastoma
(3) Brown tumor of hyperparathyroidism
(4) Fibrous dysplasia (Albright syndrome)
(5) Giant cell granuloma
(6) Giant cell tumor (osteoclastoma)
(7) Infantile cortical hyperostosis
(8) Ossifying fibroma
(9) Osteoma
(10) Osteosarcoma

M. Peripheral nerve tumors
   (1) Alveolar soft-part sarcoma
   (2) Amputation neuroma
   (3) Granular cell myoblastoma
   (4) Neurilemoma
      a. Benign
      b. Malignant
   (5) Neurofibroma
      a. Plexiform
      b. Solitary
   (6) Paraganglioma (chemodectoma)

N. Primary melanocytic tumors
   (1) Blue nevus
   (2) Melanocytic hamartoma
   (3) Melanotic progonoma (retinal tumor)
   (4) Primary orbital melanoma

O. Rhabdomyoma and rhabdomyosarcoma
   (1) Rhabdomyoma
   (2) Rhabdomyosarcoma

P. Secondary orbital tumors from adjacent structures
   (1) Conjunctival origin
      a. Melanoma
      Mucoepidermoid
      c. Squamous cell carcinoma
   (2) Eyelid origin
      a. Basal cell carcinoma
      b. Melanoma
      c. Sebaceous carcinoma
      d. Squamous cell carcinoma
   (3) Intracranial origin
      a. Astrocytoma
      b. Meningioma
   (4) Intraocular origin
      a. Medulloepithelioma
      b. Neurilemoma
      c. Retinoblastoma
      d. Uveal melanoma
(5) Nasopharyngeal origin
   a. Angiofibroma
   b. Carcinoma
   c. Melanoma

(6) Paranasal sinus origin
   a. Ethmoid sinus carcinoma
   b. Inverting papilloma
   c. Maxillary sinus carcinoma
   d. Rhabdomyosarcoma

Q. Vasculogenic lesions
   (1) Capillary hemangioma
   (2) Cavernous hemangioma
   (3) Hemangiopericytoma
   (4) Hemangiosarcoma
   (5) Kaposi sarcoma
   (6) Lymphangioma
   (7) Varices
   (8) Vascular leiomyoma
   (9) Vascular leiomyosarcoma

6. Vascular disorders
   A. Allergic vasculitis
   B. Angioedema (Quincke disease)
   C. Arteriovenous aneurysm or varices
   D. Arteriovenous fistula (varicose aneurysm)
   E. Collagen disease-lupus erythematosus (Kaposi-Libman-Sacks syndrome),
      periarteritis nodosa (Kussmaul disease), or dermatomucomyositis (Wagner-Unverricht syndrome)
   F. Cranial arteritis
   G. Thrombophlebitis
   H. Scurvy causing bilateral orbital hemorrhage


**SYNDROMES AND DISEASES ASSOCIATED WITH EXOPHTHALMOS**

1. Actinomycosis
2. Albright syndrome (fibrous dysplasia)
3. Amyloidosis (Lubarsch-Pick syndrome)
4. Apert syndrome (sphenoacromiosynactyley)
5. Arteriovenous fistula (varicose aneurysm)
6. Aspergillusosis
7. Bacillus cereus
8. Bloch-Sulzberger disease (incontinentia pigment I)
9. Bonnet-Dechaume-Blanc syndrome (neuroretinaangiomaosis syndrome)
10. Bourneville syndrome (tuborous sclerosis)
11. Caffeoy syndrome (infantile cortical hyperostosis)
12. Carotid artery-cavernous sinus fistula
13. Clostridium perfringens
14. Coenureosis
15. Craniostenosis
16. Cretinism (hypothyroidism)
17. Crouzon disease (craniofacial dysostosis)
18. Cryptococcosis
19. Cushing syndrome (adrenocortical syndrome)
20. Dejean sign (orbital floor fracture)
21. de Lange syndrome (congenital muscular hypertrophy-cerebral syndrome)
22. Dermatomucoitis (polymyositis dermatomyositis)
23. Dermoid
24. Diencephalic epilepsy syndrome (autonomic epilepsy syndrome)
25. Dirofilariasis
26. Dracontiasis (Guinea worm infection)
27. Engelmann syndrome (diaphyseal dysplasia)
28. Ewing sarcoma
29. Feer disease (infantile acrodynia)
30. Fibrosarcoma
31. Fibrous dysplasia (Albright syndrome)
32. Foix syndrome (cavernous sinus thrombosis)
33. Gardner syndrome
34. Grönlad-Strandberg syndrome (pseudoxanthoma elasticum)
35. Hallermann-Streiff-François syndrome (oculomandibulofacial dyscephaly)
36. Hand-Schüller-Christian disease (histiocytosis X)
37. Heerfordt syndrome (uveoparotid fever)
38. Hemangiomas
39. Herpes zoster
40. Hodgkin disease
41. Hollenhorst syndrome (chorioretinal infarction syndrome)
42. Homer syndrome (cervical sympathetic paralysis syndrome)
43. Hunter syndrome (MPS [mucopolysaccharidosis] II)
44. Hurler (MPS I-H) syndrome
45. Hutchinson disease (adrenal cortex neuroblastoma with orbital metastasis)
46. Hydatid cyst
47. Hydrocephalus chondrodystrophicus congenita (extreme hydrocephalus syndrome)
48. Hypertension
49. Hyperthyroidism (Basedow syndrome)
50. Hypervitaminosis A
51. Hypophosphatasia (phosphoethanolaminuria)
52. Jansen disease (metaphyseal dysostosis)
53. JXG (nevoxanthoendothelioma)
54. Kleeblattschädel syndrome (cloverleaf skull)
55. Leiomyoma
56. Leopard syndrome (multiple lentigines syndrome)
57. Leprechaunism
58. Leukemia
59. Linear nevus sebaceous of Jadassohn
60. Lupus erythematosus (Kaposi-Libman-Sacks syndrome)
61. Lymphoid hyperplasia
62. Lymphangioma
63. Lymphosarcoma
64. Melnick-Needles syndrome (osteodysplasty)
65. Meningioma
66. Mikulicz syndrome (dacryosialoadenopathy)
67. Möbius disease (congenital paralysis of sixth and seventh nerves)
68. Mucocele
69. Mucormycosis
70. Multiple myeloma
71. Mumps
72. Myasthenia gravis (Erb-Goldflam syndrome)
73. Neurilemoma
74. Noonan syndrome (male Turner syndrome)
75. Osteopetrosis (Albers-Schönberg syndrome)
76. Paget syndrome (osteitis deformans)
77. Periarteritis nodosa (Kussmaul syndrome)
78. Periocular and ocular metastatic tumors
79. Pfeiffer syndrome
80. Pierre-Robin syndrome (micrognathia-glossoptosis syndrome)
81. Progeria (Hutchinson-Gilford syndrome)
82. Pyknodysostosis
83. Quincke disease (angioedema)
84. Relapsing polychondritis
85. Retinoblastoma
86. Rhabdomyosarcoma
87. Rochon-Duvigneaud syndrome (superior orbital fissure syndrome)
88. Rollet syndrome (orbital apex-sphenoidal syndrome)
89. Sarcoidosis syndrome (Schaumann syndrome)
90. Scaphocephaly syndrome
91. Scheie syndrome (MPS I-S)
92. Scurvy (avitaminosis C)
93. Sebaceous gland carcinoma
94. Seckel syndrome (bird-headed dwarf syndrome)
95. Sézary syndrome (mycosis fungoides syndrome)
96. Shy-Gonatas syndrome (orthostatic hypotension syndrome)
97. Siegrist sign (pigmented choroidal vessels)
98. Silverman syndrome (battered baby syndrome)
99. Sphenocavernous syndrome
100. Streptococcus
101. Sturge-Weber syndrome (encephalofacial angiomatosis)
102. Syphilis (lues)
103. Thermal burns
104. Trichinelllosis
105. Trisomy syndrome (Edward syndrome)
106. Tuberculosis
107. Turner syndrome (gonadal dysgenesis)
108. von Hippel-Lindau syndrome (retinocerebral angiomatosis)
109. von Recklinghausen disease (neurofibromatosis)
110. Wegener syndrome (Wegener granulomatosis)


**SPECIFIC EXOPHTHALMOS**

1. Age
   A. Newborn-most common
      *(1) Orbital sepsis
      *(2) Orbital neoplasm including congenital malignant teratoid neoplasm
   B. Neonatal-osteomyelitis of the maxilla
   C. Early childhood (up to 1 year of age-most common)
      *(1) Dermoid
      *(2) Hemangioma
      *(3) Dermolipoma
(4) Histiocytosis X including Hand-Schüller-Christian disease
*(5) Orbital extension of retinoblastoma

D. One to five years-most common
*(1) Dermoid
(2) Metastatic neuroblastoma
(3) Rhabdomyosarcoma
(4) Epithelial cyst, such as sebaceous cyst and epithelial inclusion cyst
(5) Glioma of optic nerve
(6) Sphenoid wing meningioma
*(7) Orbital extension of retinoblastoma
(8) Fibrous dysplasia (Albright syndrome)
(9) Metastatic embryonal sarcoma
*(10) Hemangioma

E. Five to ten years-most common
(1) Pseudotumor
(2) Orbital extension of retinoblastoma
(3) Malignant lymphomas and leukemias
*(4) Dermoid
*(5) Hemangioma
(6) Meningioma
(7) Fibrous dysplasia (Albright syndrome)
(8) Rhabdomyosarcoma
(9) Orbital hematoma
(10) Glioma of optic nerve

F. Ten to thirty years-most common
*(1) Pseudotumor
(2) Mucocele
(3) Meningioma
*(4) Endocrine ophthalmopathy (thyroid-related ophthalmopathy)
(5) Lacrimal gland tumor
(6) Malignant lymphomas and leukemias
(7) Dermoid
(8) Hemangioma
(9) Peripheral nerve tumors
(10) Undifferentiated sarcomas
(11) Osteoma
(12) Fibrous dysplasia (Albright syndrome)
(13) Rhabdomyosarcoma
(14) Glioma of optic nerve

G. Thirty to fifty years-most common
*(1) Pseudotumor -
(2) Mucocele
(3) Malignant lymphomas and leukemias
*(4) Hemangioma
*(5) Endocrine ophthalmopathy (thyroid-related ophthalmopathy)
(6) Lacrimal gland tumors
(7) Rhinogenic carcinoma
(8) Malignant melanoma
(9) Osteosarcoma
(10) Fibrosarcoma
(11) Metastatic carcinoma
(12) Meningioma
(13) Dermoid

H. Fifty to seventy years-most common
*(1) Pseudotumor
*(2) Mucocele
*(3) Malignant lymphomas and leukemias
(4) Dermoid
(5) Carcinoma of palpebral or epibulbar origin
*(6) Meningioma
*(7) Endocrine ophthalmopathy (thyroid-related ophthalmopathy)
(8) Lacrimal gland tumor
(9) Osteosarcoma
(10) Fibrosarcoma
(11) Undifferentiated sarcoma
(12) Metastatic carcinoma
(13) Osteoma
(14) Fibrous dysplasia (Albright syndrome)
(15) Neurofibroma
(16) Hemangioma

I. More than seventy years-most common
(1) Melanoma
(2) Pseudotumor
*(3) Lymphoma
*(4) Metastatic tumor
(5) Basal cell carcinoma
(6) Mucocele

2. Unilateral exophthalmos-most common
A. Anatomical conditions
(1) Unilateral myopia of high degree
(2) Defects in the vault of the orbit: meningocele, encephalocele, hydroencephalocele
(3) Exophthalmos associated with arterial hypertension
(4) Recurrent exophthalmos from retrobulbar hemorrhage, lymphangioma
(5) Intermittent exophthalmos associated with venous anomalies within the cranium
(6) Disease of the pituitary gland; meningiomas involving sphenoid ridge
*(7) Unilateral exophthalmos associated with endocrine or thyroid-related ophthalmopathy
B. Traumatic conditions
(1) Contralateral floor fracture with enophthalmos
(2) Fracture of the orbit with retrobulbar hemorrhage
(3) Laceration and rupture of the tissues of the orbit and the extraocular muscles
(4) Intracranial trauma sustained at birth; aneurysm in orbit
(5) Pulsating exophthalmos from carotid-cavernous aneurysm
(6) Spontaneous retrobulbar hemorrhage as seen in whooping cough
(7) Chronic subdural hematoma bulging into orbit
(8) Posterior exophthalmos (orbital apex lesion)
   a. Pseudotumor
   b. Malignant tumor
   c. Benign tumor
   d. Vascular disease
   e. Infection
C. Inflammatory conditions
   (1) Retrobulbar abscess and cellulitis
   (2) Thrombophlebitis of the orbital veins
   (3) Cavernous sinus thrombosis
   (4) Erysipelas (St. Anthony fire)
   (5) Tenonitis
   (6) Periostitis (syphilitic or tuberculous)
   (7) Orbital mucocele, pyocele; cholesteatoma
   (8) Orbital exostosis
   (9) Paget disease with hyperostosis
   (10) Actinomycosis, trichinosis, mycotic pseudotumor
   (11) Herpes (HSV-1) with acute retinal necrosis
D. Disease of blood, lymph, and hematopoietic system
   (1) Rickets (avitaminosis D)
   (2) Scurvy (avitaminosis C)
   (3) Hemophilia (factor VIII deficiency)
   (4) Lymphosarcoma
   (5) Chloroma
   (6) Hodgkin disease
E. Space-taking lesions
   (1) Vascular anomalies
      a. Congenital orbital varix (young patient with systemic abnormalities)
      b. Cavernous hemangioma (middle age)
      c. Capillary hemangioma (young children) Kasabach-Merrit syndrome
      d. Lymphangiomas
   (2) Orbital tumors: pseudotumors, orbital cysts, meningocele, lymphangioma, orbital meningocele, lacrimal gland tumor, sarcoma, metastatic carcinoma, metastatic adrenal tumors, osteomas arising in the accessory nasal sinuses, tumors of the nasopharynx (benign and malignant)
   (3) Intracranial tumor with orbital extension including chordoma and meningioma
F. Unilateral exophthalmos in children
   (1) Inflammation
   (2) Vascular disorders
   (3) Neoplasms
   (4) Metabolic diseases
   (5) Developmental anomalies
   (6) Others
   *(7) Orbital cellulitis

3. Bilateral exophthalmos-most common
   *A. Thyroid or endocrine ophthalmopathy
   B. Orbital myositis (owing to causes other than thyroid dysfunction)
   C. Cavernous sinus thrombosis (Foix syndrome)
   D. Metastatic neuroblastoma
   E. Hand-Schüller-Christian disease (histiocytosis X)
   F. Crouzon disease (craniofacial dysostosis)
   G. Paget disease (osteitis deformans)

4. Type proptosis-most common
   A. Straightforward—glioma of optic nerve, intracanal cavernous hemangioma
   B. Down and temporal-mucocele of frontal sinus
   C. Down and nasal-lacrimal gland lesion
   D. Downward—tumor of roof of orbit
   E. Upward—tumor of floor of orbit

5. Transient exophthalmos
   *A. Orbital varices
   B. Orbital varices with intracranial extension
   C. Arteriovenous malformations
   D. Cavernous hemangioma
   E. Intraorbital arteriovenous malformation

6. Pulsating exophthalmos-most common
   *A. Carotid-cavernous fistula
   B. von Recklinghausen disease associated with bony defect of skull
   C. Large frontal mucocele
   D. Meningoencephalocele
   E. Blow-in fracture of roof of orbit
   F. Neurofibromatosis
   G. Fistula
   H. Malignancies
   I. Mucoceles
   J. Orbital varix
   K. Dermoid cysts
   L. Aneurysm

7. Recurrent exophthalmos
   A. Recurrent orbital inflammation (pseudotumor) or hemorrhage
   B. Orbital cysts that rupture
   C. Lymphangioma (children)
D. Syndrome of intermittent exophthalmos-congenital venous malformations of the orbit: venous angioma and orbital varix
E. Temporal lobe tumor with orbital extension
F. Neurofibromatosis
G. Vascular neoplasm

8. Intermittent exophthalmos
   A. Orbital varices
   B. Recurrent hemorrhage
   C. Vascular neoplasm
   D. Lymphangioma

9. Exophthalmos associated with conjunctival chemosis, restricted movement of eyes because of pain-pseudotumor

10. Exophthalmos in an acutely ill patient-cavernous sinus thrombosis

11. Exophthalmos associated with engorged conjunctival episcleral vessels
    A. Nonpulsating-cerebral arteriovenous angioma, ophthalmic vein thrombosis, or cavernous sinus thrombosis
    B. Pulsating exophthalmos-carotid-cavernous sinus fistula

12. Exophthalmos associated with a palpable mass in region of the lacrimal gland
    A. Primary inflammatory exophthalmos
    B. Neoplasm
    C. Sarcoidosis syndrome (Schaumann syndrome)
    D. Hodgkin disease

13. Exophthalmos in patient with uncontrolled diabetes, usually with acidosis, who develops unilateral lid edema, ptosis, internal and external ophthalmoplegia, proptosis, and severe vision loss-orbital mucormycosis

14. Exophthalmos in an infant with ecchymosis of the eyelids
    A. Metastatic neuroblastoma
    B. Orbital leukemia infiltration

15. Bilateral exophthalmos from bilateral orbital pseudotumor
    A. Eosinophilic granuloma
    B. Retroperitoneal fibrosis
    C. Myasthenia gravis (Erb-Goldflam syndrome)


**ENOPHTHALMOS**

1. Senility (common)
2. Wasting diseases-loss of orbital fat
*3. Injury-blowout fracture of floor of orbit (most common)
4. Orbital varices-transient exophthalmos with fat atrophy

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<th>Extracted Tables:</th>
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<tbody>
<tr>
<td>Table Exophthalmos (Up to 1 year)</td>
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<td>Table Exophthalmos (1-5 years)</td>
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<tr>
<td>Table Exophthalmos (5-10 years)</td>
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<td>Table Exophthalmos (50-70 years)</td>
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<tr>
<td>Table Exophthalmos (more than 70 years)</td>
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<td>Table Pulsating exophthalmos (most common)</td>
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<td>Table Recurrent exophthalmos</td>
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5. Chronic or severe liver or gallbladder disease (usually in right eye owing to increased tone of orbicularis muscle and extraocular muscles)
6. Iatrogenic
   A. Orbital decompression
   B. Sinus surgery
7. Superior sulcus deformity
   A. Traumatic bony loss
   B. Atrophy of the orbital tissues
   C. Levator detachment with ptosis
   D. Migration of muscle cone implant
   E. Herniated orbital fat secondary to an orbital fracture
8. Associated syndromes
   A. Arthrogryposis (amyoplasia congenital)
   B. Babinski-Nageotte syndrome (medullary tegmental paralysis)
C. Cestan-Chenais syndrome (lesion in the lateral portion of medulla oblongata)
D. Cockayne syndrome (dwarfism with retinal atrophy and deafness)
E. Craniofacial syndrome (whiplash injury)
F. Cretinism (hypothyroidism)
G. Cryptophthalmia syndrome
H. Dejean syndrome (orbital floor syndrome)
I. Dejerine-Klumpke syndrome (thalamic hyperesthetic anesthesia)
J. Duane retraction syndrome
K. Freeman-Sheldon syndrome (craniocarpotarsal dysplasia)
L. General fibrosis syndrome
M. Greig syndrome (ocular hypertelorism syndrome)
N. Hemifacial microsomia syndrome (François-Haustrate syndrome)
O. Horner syndrome (cervical sympathetic paralysis syndrome)
P. Klippel-Trenaunay-Weber syndrome (angioosteohypertrophy syndrome)
Q. Krause syndrome (encephaloophthalmic syndrome)
R. Maple syrup urine disease (branched chain ketoaciduria)
S. Morquio syndrome (MPS IV)
T. Naffziger syndrome (scalenus anticus syndrome)
U. Pancoast syndrome (superior pulmonary sulcus syndrome)
V. Parry-Romberg syndrome (progressive facial hemiatrophy)
W. Passow syndrome (Bremer status dysraphicus)
X. Raeder syndrome (paratrigeminal paralysis)
Y. Retroparotid space syndrome
Z. Silent sinus syndrome
AA. Vernet syndrome (jugular foramen syndrome)
BB. von Herrenschwand syndrome (sympathetic heterochromia)
CC. Wallenberg syndrome (dorsolateral medullary syndrome)

9. Apparent enophthalmos with horizontal conjugate gaze
10. Metastatic adenocarcinoma of orbit (cicatrical)
11. Neurofibromatosis: pulsating enophthalmos
12. Typhoid fever (abdominal typhus)


**INTRAORBITAL CALCIFICATIONS**

1. Calcification of more irregular configuration and texture
A. Cysticercosis
B. Orbital hematoma
C. Plexiform neurofibroma
D. Toxoplasmosis
E. Tuberculosis
2. Calcification of orbital vessels
   A. Atheromatous plaque
   B. Monkeberg sclerosis
   C. Secondary to metabolic-endocrine disturbances such as hyperparathyroidism or hypervitaminosis
   D. Band-shaped keratopathy
3. Chronic inflammatory and parasitic disease of the orbit
4. Hemangiopericytoma
5. Intraocular calcifications following
   A. Congenital deformity
   B. Malignant lacrimal gland tumor
   C. Recurrent iritis and keratitis
   D. Retinal detachment
   E. Trauma (perforating, nonperforating, or surgical)
6. Intraocular sarcoma
7. Mucocele
8. Myositis ossificans
9. Orbital phleboliths: helical form in veins-smooth, round, or oval
10. Organized hematomas of the orbit
    * 11. Retinoblastoma
12. Retrolental fibroplasia
13. Sites of intraocular calcification
    A. Cyclitic membrane
    B. Lens
    C. Peripapillary choroid
    D. Posterior pole to ora serrata in region of choroid and pigment epithelium
    E. Retina
    F. Vitreous


**ORBITAL BRUIT (NOISE HEARD OVER ORBIT WITH STETHOSCOPE)**
**Extracted Table Orbital bruit (noise heard over orbit with stethoscope)**

1. Bilateral
   - A. Hyperthyroidism
   - B. Severe anemias
2. Unilateral
   *A. Abnormal communication in the cavernous sinus (i.e., bilateral carotid-cavernous sinus)*
   - B. Aneurysmal angioma of orbit or fundus such as in Wyburn-Mason syndrome (Bonnet-Dechaume-Blanc syndrome)
   - C. Arteriovenous aneurysm (arteriovenous fistula)
   - D. Intermittent or pulsating exophthalmos
   - E. Stenosis of carotid artery including thrombosis, sclerosis, or external pressure such as that due to an outer-ridge sphenoidmeningioma


**ORBITAL EMPHYSEMA (AIR FOUND IN ORBITAL TISSUES AND ADNEXA USUALLY DEMONSTRABLE BY PALPATION)**

*1. Due to fracture of ethmoid sinuses or orbital floor
2. Following forceful blowing of nose
3. Injury from compressed air
4. Orbital cellulitis and abscess with gas formation by infecting organism
5. Osteomyelitis and infected sinus with fistulous communication with gas formation by infecting organism
6. Resulting from use of high-speed dental drill and air-water spray during oral operation
7. Subconjunctival emphysema seen with mechanical ventilation


**ORBITAL PAIN**

1. Acute dacryoadenitis
2. Amputation neuroma of the orbit
3. Associated syndromes
   A. Cavernous sinus thrombosis syndrome
   B. Charlin syndrome (nasal nerves syndrome)
   C. Erysipelas
   D. Ophthalmoplegic migraine syndrome
   E. Raeder syndrome (paratrigeminal paralysis)
   F. Tolosa-Hunt syndrome (painful ophthalmoplegia).
4. Break-bone fever (dengue fever)
5. Clostridium perfringens
6. Eye strain from uncorrected errors of refraction
7. Myositis
   A. Collagen diseases
   B. Infectious myositis
   C. Trichinosis
8. Orbital cellulitis or abscess
9. Orbital periostitis because of injury, tuberculosis, syphilis, extension of sinus disease, or other conditions
   *10. Pseudotumor or tumor of the orbit-pain infrequently present*
11. Retrobulbar neuritis
12. Trauma
13. Tumors of cerebellopontine angle, frequent lesion of seventh nerve


**SHALLOW ORBITS OR DIMINISHED ORBITAL VOLUME (ILLUSION OF PROPTOSIS OR GLAUCOMA)**

1. Aminopterin-induced syndrome
2. Apert syndrome (acrocephalosyndactyly)
3. Carpenter syndrome
4. Cerebrohepatorenal syndrome (Smith-Lemli-Opitz syndrome)
5. Craniostenosis
6. Crouzon disease (craniofacial dysostosis)
7. Diseases of nasal passages and sinuses
   A. Dentigerous cysts
   B. Fibrous dysplasia (Albright syndrome)
   C. Hypoplasia of maxilla associated with chronic maxillary sinusitis
   D. Rhinoscleroma
8. Dubowitz syndrome
9. Early enucleation of eye
*10. Familial hypoplasia of orbital margin
11. Frontometaphyseal dysplasia (FMD)
12. Hyperostosis (hypertrophy of orbital bones)
13. Hypophosphatasia-harlequin orbit (shallow orbit with arched superior and lateral wall)
14. Kleeblattschädel syndrome
15. Lateral displacement of medial orbital wall by hypertrophic polypoid nasal sinus disease
16. Marshall-Smith syndrome
17. Oculoauriculovertebral dysplasia (Goldenhar syndrome)
18. Osteogenesis imperfecta (van der Hoeve syndrome)
19. Radiation injury of bone
20. Robert syndrome (pseudothalidomide syndrome)
21. Saethre-Chotzen syndrome
22. Secondary to fracture
23. Stanesco dysostosis syndrome
24. Trisomy 13-(trisomy D) (Patau syndrome)
25. Trisomy (Edward syndrome)
26. Zellweger syndrome
27. 6q- D syndrome
28. 9p- syndrome


**PSEUDOHYPERTELORISM (ILLUSION OF INCREASED DISTANCE BETWEEN BONY ORBITS AND INCREASED INTERPUPILLARY DISTANCE)**

1. Blepharophimosis
2. Epicanthal skin folds
3. Exotropia
4. Flat nasal bridge of nose
5. Increased distance between the inner canthi (telecanthus)
6. Widely spaced eyebrows


**HYPERTELORISM (INCREASED DISTANCE BETWEEN BONY ORBITS AND INCREASED INTERPUPILLARY DISTANCE)**

1. Aarskog syndrome (faciodigitogenital syndrome)
2. Acrocollosal syndrome
3. Acrodyssostosis syndrome
4. Albers-Schönberg disease (osteopetrosis)
5. Aminopterin-induced syndrome
6. Apert syndrome (acrodysplasia)
7. Association of hypertelorism, microtia, and facial clefting
8. Baraitser- Winter syndrome
9. BBB syndrome (hypertelorism-hypospadias syndrome)
10. Blatt syndrome (cranioorbitoocular dysraphia)
11. Blepharoonosofacial syndrome
12. Camptomelic dysplasia syndrome
13. Carpenter syndrome (acrocephalopolysyndactyly II)
14. Cat's-eye syndrome (Schachenmann syndrome)
15. Cerebral gigantism (Sotos syndrome)
16. Cerebrohepatorenal syndrome (Zellweger)
17. Cherubism
18. Chromosome partial long-arm deletion syndrome (de Grouchy syndrome)
19. Chromosome partial short-arm deletion syndrome [monosomy partial (short-arm) syndrome]
20. Chromosome short-arm deletion
21. Chondrodystrophy calcificans congenita (Conradi syndrome)
22. Cleft lip and palate sequence
23. Clefting, ectropion, and conical teeth syndrome
24. Cleidocranial dysostosis syndrome
25. Coffin-Lowry syndrome
26. Congenital hemihypertrophy
27. Cranioacarpotarsal syndrome (Freeman-Sheldon syndrome)
28. Cranioleidodysostosis syndrome (Marie-Sainton syndrome)
29. Craniosynostosis-radial aplasia (Baller-Gerold syndrome)
30. Cretinism (hypothyroidism)
31. Cri-du-chat syndrome (Cry of the cat syndrome)
32. Crouzon disease (craniofacial dysostosis)
33. Cryptophthalmos syndrome
34. Curtius syndrome (ectodermal dysplasia with ocular malformations)
35. Diamond-Blackfan syndrome
36. DiGeorge sequence
37. Down syndrome (mongolism)
38. Dubowitz syndrome (dwarfism-eczema-peculiar facies)
39. Duplication 14q syndrome
40. Ehlers-Danlos syndrome (fibrodysplasia elastica generalisata)
41. Engelmann syndrome (diaphyseal dysplasia)
42. 18p syndrome
43. Faciooculoacousticorenal syndrome
44. Familial characteristic
45. Familial metaphyseal dysplasia (Pyle disease)
46. Fetal alcohol syndrome
47. Fetal aminopterin effects
48. Fetal hydantoin effects
49. Fish odor syndrome
50. 4q syndrome
51. Frontonasal dysplasia syndrome (median cleft face syndrome)
52. Frontal encephaloceles
53. Gorlin syndrome (orodigitofacial dysostosis)
54. Greig syndrome (hypertelorism)
55. Haney-Falls syndrome (congenital keratoconus posticus circumscriptus)
56. Holt-Dram syndrome
57. Hurler syndrome (MPS I-H)
58. Hydrocephalus
59. Hypomelanosis of Ito syndrome (systematized achromic nevus)
60. Ichthyosis (collodion baby)
61. Infantile gigantism
62. Infantile hypercalcemia with supravalvular aortic stenosis (Williams-Beuren syndrome)
63. Iris dysplasia-hypertelorism-psychomotor retardation syndrome
64. Jacobs syndrome (triple X syndrome)
65. KBB syndrome (initials of family studied)
66. Kleeblattschädel syndrome (extreme hydrocephalus syndrome)
67. Klein syndrome
68. Klinefelter XXY syndrome (gynecomastia-aspermatogenesis syndrome)
69. Klippel-Feil syndrome (synostosis cervical vertebrae)
70. Larsen syndrome
71. Leprechaunism
72. Lissencephalia (Miller-Dieker syndrome)
73. Little syndrome (nail-patella syndrome)
74. Mandibulofacial dysostosis (Franceschetti syndrome)
75. Maple syrup urine disease (branched-chain ketoaciduria)
76. Marfan syndrome (arachnodactyly-dystrophia-mesodermalis congenita)
77. Marshall-Smith syndrome
78. Meckel-Gruber syndrome
79. Melnick-Needles syndrome (osteodysplasty)
80. Metaphyseal dysostosis (Jansen disease)
81. McFarland syndrome
82. Morquio-Ullrich syndrome (MPS IV)
83. Multiple basal cell nevi (Godin-Goltz syndrome)
84. Multiple lentigines syndrome (Leopard syndrome)
85. Myelomeningocele-Chiari malformations
86. Noonan syndrome (male Turner syndrome)
87. Oculodentodigital syndrome
88. Oculomandibulofacial dyscephaly (Hallermann-Streiff syndrome)
89. Optic nerve hypoplasia
90. Orofaciodigital (OFD) type I and type II (Mohr syndrome)
91. Osteogenesis imperfecta (van der Hoeves syndrome)
92. Otopalatodigital syndrome (OPD syndrome)
93. Pallister-Killian syndrome
94. Pena-shokeir type I syndrome
95. Pfeiffer syndrome
96. Potter syndrome (renofacial syndrome)
97. Ring B chromosome
98. Ring chromosome
99. Rieger syndrome (dysgenesis mesostromalis)
100. Robert syndrome (pseudothalidomide syndrome)
101. Robinow syndrome (fetal face syndrome)
102. Saethre-Chotzen syndrome (acrocephalosyndactyly type III)
103. Sjögren-Larson syndrome (oligophrenia-ichthyosis-spastic diplegia syndrome)
104. Sprengel syndrome
105. Traumatic nasoorbital fracture
106. Triploidy syndrome
107. Trisomy syndrome
108. Trisomy 6 q syndrome
109. Trisomy 9q syndrome
110. Trisomy 13- (Patau syndrome)
111. Trisomy 17p syndrome
112. Trisomy syndrome
113. Turner-Bonnevie-Ullrich-Nielsen syndrome
114. Turner syndrome (gonadal dysgenesis)
115. Waardenburg syndrome (embryonic fixation syndrome)
116. Weaver syndrome
117. Williams syndrome
118. XXXX syndrome
119. XXXXXX syndrome
120. XXXXY syndrome
121. 4p- syndrome (Wolf syndrome)
122. 4p- D syndrome
123. 6p- D syndrome
124. 9p- syndrome
125. 10q- syndrome
126. 13q- syndrome
127. 18q- syndrome


**HYPOTELORISM (DECREASED DISTANCE BETWEEN BONY ORBITS AND DECREASED INTERPUPILLARY DISTANCE)**

1. Arrhinencephaly (holoprosencephaly)
2. Cebocephalia
3. Cockayne syndrome (dwarfism with retinal atrophy and deafness)
4. Coffin-Siris syndrome
5. Ethmocephalus
*6. Familial
7. François diencephalic syndrome (Hallerman-Streiff syndrome)
8. Goldenhar syndrome (oculoauriculovertebral dysplasia)
9. Maternal phenylketonuria fetal effects
10. Meckel-Gruber syndrome
11. Median cleft lip (frontonasal dysplasia syndrome)
12. Median philtrum-premaxilla anlage
13. Ocular-dental-digital dysplasia (Meyer-Schivickerath and Weyers syndrome)
14. Ring syndrome
15. Trigonocephaly (C syndrome, Opitz trigonocephaly syndrome)
16. Trisomy 13- (Patau syndrome, trisomy D syndrome)
17. Trisomy 20p syndrome
18. Trisomy (Down syndrome, mongolism)
19. Turner syndrome (gonadal dysgenesis)
20. Williams syndrome
21. Wolf syndrome (monosomy partial syndrome)
22. 5p- D syndrome
23. 18p- syndrome


**DEEP-SET EYES**

1. Cockayne syndrome (dwarfism with retinal atrophy and deafness)
2. Craniocarotarsal syndrome (Freeman-Sheldon syndrome)
3. Familial
4. Marfan syndrome (dolichostenomelia-arachnodactylyhyperchon-droplasia-dystrophia mesodermalis congenita)
5. Mesodermal dysmorphodystrophy (Weill- Marchesani syndrome)
6. Oculocerebrorenal syndrome (Lowe syndrome)
7. Pyknody sostosis
8. Syndrome of blepharophimosis with myopathy


**PROMINENT SUPRAORBITAL RIDGES**

1. Apert syndrome (acrocephalosyndactylism syndrome)
2. Basal cell nevus syndrome (Godin-Goltz syndrome)
3. Cleidocranial dysostosis (Marie-Sainton syndrome)
4. Congenital lipodystrophy
5. Congenital syphilis (congenital lues)
6. Ectodermal dysplasia (Curtius syndrome)
*7. Frontometaphyseal dysplasia
8. Hurler syndrome (MPS I-H)
9. Marfan syndrome (arachnodactyly-dystrophia mesodermalis congenita)
10. Otopalatodigital syndrome (Taybi syndrome)
11. Pyle metaphyseal dysplasia syndrome


**OSTEOLYSIS OF BONY ORBIT**

1. Autoimmune diseases, such as Wegener granulomatosis
2. Congenital
3. Hyperparathyroidism
4. Injury, such as blowout fracture of orbital floor
5. Meningocele and encephalocele of orbit
6. Metastasis from remote primary neoplasms
7. Primary orbital disease
   A. Infectious, including tuberculosis and syphilis
   B. Neoplastic, including neurofibroma and lacrimal gland tumor
   C. Cystic, including dermoid and epidermoid cyst
8. Reticuloendotheliosis as histiocytosis X (Hand-Schüller-Christian disease)
9. Secondary extension of infectious or neoplastic disease from adjacent sinuses, brain, skin, bone, nasopharynx, and esophagus
10. Sinus disease including mucoceles


**FOSSA FORMATION OF ORBIT (LOCAL EXPANSION OF BONY ORBITAL WALL CAUSED BY PERSISTENT PRESSURE; BONY CORTEX IS INTACT)**

1. Encapsulated benign lacrimal gland tumor
2. Encapsulated malignant lacrimal gland tumor
3. Orbital dermoid


**SUBPERIOSTEAL ORBITAL HEMORRHAGE**

1. Generalized diseases with bleeding diatheses
2. Paranasal sinusitis
3. Sudden elevation of cranial venous pressure
4. Trauma


**ORBITAL HEMORRHAGE**

1. Idiopathic
2. Other
   A. General endotracheal anesthesia
   B. Late migration of orbital implant
3. Preexisting vascular tumors
   A. Cholesterol granuloma
   B. Cystic lymphangiomas
   C. Hemorrhagic varix
4. Surgery
   A. Retrobulbar injection
   B. Rhinoplasty
   C. Subtentorial infusion anesthesia
5. Systemic
   A. Heparin
   B. Thrombolytic
6. Trauma


**EXPANSION OF ORBITAL MARGINS (USUALLY ASSOCIATED WITH BENIGN TUMORS OF THE ORBIT)**

1. Dermoid
2. Hemangioma
3. Lacrimal gland tumors
4. Meningioma
5. Neurofibroma


HYPERTROPHY OF ORBITAL BONES (HYPEROSTOSIS OR SCLEROSIS OR BOTH)

1. Acromegaly
2. Anemias of childhood (severe: Cooley, sickle cell, spherocytosis, iron deficiency)
3. Cerebral atrophy (childhood)
4. Craniostenosis
5. Engelmann disease (hereditary diaphyseal dysplasia)
6. Hyperostosis frontalis interna
7. Idiopathic
8. Infantile cortical hyperostosis (Caffey disease)
9. Microcephaly
10. Myotonia atrophica (myotonic dystrophy, Curschmann-Steinert syndrome)
11. Osteopetrosis (Albers-Schönberg disease)
12. Paget disease (osteitis deformans)
13. Tumors of orbit, including osteoma, fibrous dysplasia (Albright syndrome), menigioma, metastatic neuroblastoma, mixed tumors of lacrimal gland, transitional cell carcinomas of the nasopharynx


EXPANSION OF OPTIC CANAL

1. Increased intracranial pressure
2. Inflammatory lesions
   A. Chiasmatic arachnoiditis
   B. Nonspecific granuloma
   C. Sarcoid granuloma
   D. Tuberculoma
3. Tumors
   A. Meningioma
   B. Metastatic sarcoma to choroid
C. Neurofibromatosis (von Recklinghausen syndrome)
D. Optic nerve glioma
E. Retinoblastoma

4. Vascular lesions
   A. Arteriovenous malformation
   B. Ophthalmic artery aneurysm


**SMALL OPTIC CANALS**

1. Developmental abnormalities
   A. Anophthalmos or microphthalmos
   B. Enucleation
   C. Craniosynostosis (CSO)
2. Dysostoses
   A. Osteopetrosis (Albers-Schönberg syndrome)
   B. Fibrous dysplasia (Albright syndrome)
   C. Pyle disease (craniometaphyseal dysplasia syndrome)
   D. Paget disease (osteitis deformans)
3. Inflammatory lesions-osteitis
4. Tumor-meningioma


**EROSION OF OPTIC CANAL**

1. Lateral wall
   A. Pituitary tumor
   B. Aneurysm of internal carotid artery
   C. Craniopharyngioma
   D. Tumor of orbital apex
2. Medial wall
   A. Carcinoma of sphenoid sinus
   B. Mucocele of sphenoid sinus
C. Granuloma of sphenoid sinus

3. Roof
   A. Tumor of anterior cranial fossa
   B. Surgical unroofing

4. Decrease in length
   A. Tumor of orbital apex

5. Complete destruction
   A. Malignant tumor
   B. Eosinophilic granuloma


---

**ENLARGEMENT OF SUPERIOR ORBITAL FISSURE**

1. Carotid cavernous fistula
2. Chronic increased intracranial pressure
3. Extension of infraorbital mass into fissure
4. Intracavernous carotid aneurysm
5. Intracranial chordoma
6. Masses within middle fossa
7. Metastatic carcinoma to sphenoid wings
8. Nasopharyngeal carcinoma-rare
9. Neurofibromatosis including optic nerve glioma
10. Orbital dysplasia
11. Orbital varix
12. Pituitary neoplasm-changes in sella and clinoid process
13. Posterior orbital encephalocele
14. Sarcomas, neurilemoma, or other orbital malignancies


NARROWING OF SUPERIOR ORBITAL FISSURE

1. Chronic hemolytic anemias of childhood
2. Fibrous dysplasia (Albright syndrome)
3. Meningioma
4. Osteitis
5. Osteoblastoma
6. Osteoma
7. Osteopetrosis (Albers-Schönberg syndrome)
8. Paget disease (osteitis deformans)


SMALL ORBIT

1. Anophthalmos
2. Enucleation
3. Microphthalmos
4. Mucocele


LARGE ORBIT

1. Congenital
   A. Dysplasia
   B. Glaucoma
   C. Serous cysts
2. Pseudotumor
3. Tumors within the muscle cone
   A. Hemangiomas
   B. Neurofibroma
   C. Optic glioma
   D. Orbital varix
   E. Retinoblastoma
HEMATIC ORBITAL CYSTS (BLOOD CYST OF ORBIT)

1. Blood dyscrasia
2. Cavernous hemangioma
3. Childbirth
4. Lymphangioma
5. Orbital blunt trauma
6. Spontaneous hemorrhage
7. Vascular disease


Lids

CONTENTS

Mongoloid palpebral fissure 46
Antimongoloid palpebral fissure 47
Pseudoptosis 48
Blepharoptosis 49
Syndromes and diseases associated with ptosis 52
  Ptosis-Diagnostic table 55
Specific blepharoptosis 57
Horner syndrome 59
  Horner syndrome-Diagnostic table 61
Ptosis of lower lid 62
Lagophthalmos 62
Pseudo-lid retraction 62
Lid retraction 62
Lid lag 64
Blepharospasm 65
Facial palsy 66
Infrequent blinking 68
Frequent blinking 68
Lid edema 69
Bleeding of the eyelid 77
Ectropion 78
Entropion 79
Epicanthus 80
Hypopigmentation 81
Hyperpigmentation 83
Tumors of eyelids 86
  Tumors of eyelids-Diagnostic table 87
Xanthelasma 88
Chronic blepharitis 89
Acute blepharitis 94
Thickened eyelids 97
Blepharophimosis 97
Euryblepharon 98
Lid coloboma 98
Necrosis of eyelids 99
Poliosis 99
Trichomegaly 99
Madarosis 100
  Madarosis-Diagnostic table 102
Distichiasis 104
Coarse eyebrows 105
Synophrys 105
Hertogh sign 105
Lid myokymia 105
Preseptal cellulitis of eyelid 106
Telecanthus 106
Ankyloblepharon 107
Flaring of nasal part of eyebrow 107
High arched brow 108
Absent brow hair 108
Trichiasis 108

**MONGOLOID PALPEBRAL FISSURE (TEMPORAL CANTHUS HIGHER THAN NASAL CANTHUS)**

1. Esotropia syndrome
2. Exotropia syndrome
3. Amniogenic band syndrome (Streeter dysplasia)
4. Anhidrotic ectodermal dysplasia
5. Cebocephalia (fetalis hypoplastica)
6. Chondrodystrophy (Conradi syndrome)
7. Chromosome short-arm deletion
8. Congenital spherocytic anemia
9. Crouzon syndrome (hereditary craniofacial dysostosis)
10. Duplication 14Q syndrome
11. Femoral-facial syndrome
12. Fetal hydantoin syndrome
13. Hereditary ectodermal dysplasia syndrome (Siemen syndrome)
14. Jacobs syndrome (triple X syndrome)
15. Jarcho-Levin syndrome
16. Klinefelter XXY syndrome (gynecomastia-aspermato genesis syndrome)
17. Laurence-Moon-Biedl syndrome (retinitis pigmentosa-polydactyl-adiposogenital syndrome)
18. Meckel syndrome (dysencephalial-splanchnocystic syndrome)
19. Miller-Dieker syndrome
*20. Mongoloid (trisomy or Down syndrome)
*21. Asian persons
22. Peters trisomy 5p (Peters anomaly)
23. Otopalataodigital syndrome
24. Pfeiffer syndrome
25. Pleonosteosis syndrome (Leri syndrome, carpal tunnel syndrome)
26. Prader-Willi syndrome
27. Rhizomelic chondrodysplasia punctata syndrome
28. Trisomy mosaic and 9p- syndromes
29. Trisomy syndrome
30. Trisomy 6Q syndrome
31. Trisomy 9Q syndrome
32. Trisomy and 18q syndrome
33. Trisomy 20p syndrome
34. XXXXX syndrome
35. XXXXY syndrome
36. 4p- syndrome
37. 5p- syndrome


**ANTIMONGOLOID PALPEBRAL FISSURE (DOWNWARD DISPLACEMENT OF TEMPORAL CANTHUS)**

1. Aarskog syndrome (faciodigitogenital syndrome)
2. Acrocephalosyndactylia (Apert syndrome)
3. Esotropia and exotropia
4. Bird-headed dwarf syndrome (Seckel syndrome)
5. Baraitser-Winter syndrome
6. Cardiofaciocutaneous syndrome
7. Cerebral gigantism (Sotos syndrome)
8. Cleft palate
9. Chromosome short-arm deletion
10. Cloverleaf cranium
11. Coffin-Lowry syndrome
12. Cohen syndrome
13. Congenital facial hemiatrophy (Möbius syndrome)
14. Craniofacial dysostosis (Freeman-Sheldon syndrome; whistling face syndrome)
15. Craniofacial dysostosis (Crouzon syndrome)
16. Cri-du-chat syndrome (Cry of the cat syndrome)
17. De Lange syndrome (congenital muscular hypertrophy -cerebral syndrome)
18. Di George syndrome
19. Epidermal nevus syndrome (ichthyosis hystrix)
20. Lethal multiple pterygium syndrome (LMPS)
21. Linear nevus sebaceous of Jadassohn (Jadassohn-type anetoderma)
22. Mandibulofacial dysostosis (Franceschetti syndrome and Treacher-Collins syndrome)
23. Marchesani syndrome (dystrophia mesodermalis congenita hyperplastica)
24. Maxillofacial dysostosis
25. Nager syndrome
26. Noonan syndrome (male Turner syndrome)
27. Obesity-cerebral-ocular-skeletal anomalies syndrome
28. Oculooauriculovertebral dysplasia (Goldenhar syndrome)
29. Oculomandibulofacial dyscephaly (Hallermann-Streiff syndrome)
30. Organoid nevus syndrome
31. Otopalatodigital syndrome (OPD)
32. Pyknodysostosis
33. Partial trisomy of long arm of chromosome 630. Pseudo-Ullrich-Turner syndrome
34. Ring D chromosome
35. Rubinstein-Taybi syndrome (broad thumbs syndrome)
36. Ruvalcaba syndrome
37. Saethre-Chotzen syndrome
38. 3-P syndrome
39. Trauma
40. Trisomy 9p syndrome
41. Trisomy syndrome
42. Trisomy 17p syndrome
43. Trisomy 20P syndrome
44. Trisomy syndrome (E syndrome)
45. Turner syndrome (gonadal dysgenesis)
46. Wolf syndrome (chromosome partial deletion syndrome)
47. 4q- syndrome
48. 21q syndrome
49. 9p syndrome


**PSEUDOPTOSIS**

Pseudoptosis includes conditions that simulate ptosis, but lid droop is not the result of levator malfunction, and ptosis is usually corrected when the causative factors are cleared up or removed.

1. Due to globe displacement
   A. Anophthalmia including poorly fitting prosthesis
   *B. Enophthalmos such as that resulting from blowout fracture of the floor of the orbit or atrophy of orbital fat
   *C. Microphthalmia
   *D. Phthisis bulbi
   E. Hypotony and inward collapse of eye
   F. Cornea plana
   G. Hypotropia of that eye or hypertropia of the other eye

2. Due to mechanical displacement of the lid
   A. Inflammation
      (1) Trachoma-thick, heavy lid
      *(2) Chalazion or hordeolum
      (3) Elephantiasis
      (4) Chronic conjunctivitis-conjunctival thickening
(5) Traumatic or infectious edema involving the lid
*(6) Blepharitis
(7) Corneal foreign body
(8) Contact lens
(9) Sinusitis, cellulitis

B. Tumors, especially fibromas, lipomas, or hemangiomas
C. Scar tissue due to burns, physical trauma, and lacerations that can bind the lid down
D. Tumors of lacrimal gland-S-shaped lid

*3. Dermatochalasis (ptosis adiposa, baggy lids, "puffs"-senile atrophy of the lid skin)
4. Blepharochalasis-a rare condition occurring in young persons, characterized by recurrent bouts of inflammatory lid edema with subsequent stretching of the skin
5. The oriental lid-the palpebral fissure is narrower than normal and the upper lid rarely has a furrow; hence, the fold usually hangs down to or over the lid margin.
6. Dissociated vertical deviation (DVD)
7. Duane syndrome (retraction syndrome)
8. Blepharospasm-eyebrow lower than normal, hemifacial spasm
9. Contralateral widening of the lid fissure as pseudoproptosis (see p. 4), exophthalmos, or lid retraction (see p. 62)
10. Vertical strabismus


BLEPHAROPTOSIS (PTOSIS, DROOPY UPPER LID; WEAK LEVATOR PALPEBRAE SUPERIORIS MUSCLE)

1. Congenital ptosis
   *A. Simple (most congenital ptosis)-may be the result of autosomal dominant inheritance
   B. Complicated ptosis
      (1) Ptosis with ophthalmoplegia (most congenital ptosis)-the most commonly involved muscle is the superior rectus
      (2) Ptosis with other lid deformities such as epicanthus, blepharophimosis, microphthalmia, and lid coloboma-may be hereditary
      (3) Synkinetic (paradoxical) ptosis-aberrant nervous connections from the other extrinsic muscles of the eye and jaw to the levator muscle
         a. Marcus Gunn phenomenon (jaw-winking reflex)-motor root of the fifth cranial nerve to the muscle of mastication also is misdirected through the third nerve to the levator muscle.
b. Phenomenon of Marin Amat (reverse jaw-winking reflex)
c. Misdirected third nerve syndrome-bizarre eyelid movements that may accompany various eye movements; the ptotic eyelid may rise as the medial rectus, the inferior rectus, or the superior rectus muscle contracts.

(4) Homer syndrome

C. Involutional ptosis

D. Mechanical

(1) Periorbital tumor
(2) Neuroma, neurofibroma
(3) Cicatricial skin changes

2. Acquired ptosis

A. Traumatic ptosis

(1) Eyelid laceration
(2) Postsurgical ptosis
   a. Anterior transposition of inferior oblique muscle
   b. Enucleation
   c. Orbital operation
   d. Cataract operation
   e. Radial keratotomy
(3) Foreign bodies lying in the roof of the orbit
(4) Fracture of orbital roof, also following contusion with resulting hematoma but without fracture
(5) Air-blast injury
(6) Botulinum toxin treatment of strabismus and blepharospasm
(7) Prolonged hard contact lens wear
(8) Infratemporal fossa foreign body

B. Neurogenic ptosis

(1) Peripheral involvement of the third nerve
(2) Basilar, cortical, and nuclear lesions
(3) Cerebral hemorrhages, tumors, or abscesses
(4) Multiple neuritis, nerve syphilis, or multiple sclerosis
(5) Homer syndrome-lower lid higher than other lower lid
(6) Familial dysautonomia (Riley-Day syndrome)
(7) Misdirected third nerve syndrome-following third nerve palsy the fibers do not regrow into their respective muscles.
(8) Aseptic meningitis, transient
(9) Pituitary tumor

C. Myogenic ptosis

(1) Primary muscular atrophy (late familial ptosis); ptosis is usually the only symptom.
(2) Dystrophia myotonina, in which there is dystrophia not only of the extraocular muscles but also of the face, neck, and extremities
(3) Myasthenia gravis, nonfamilial acquired ptosis
(4) The congenital fibrosis syndrome characterized by bilateral ptosis and gradual fibrosis of all the extraocular muscles
(5) Oculopharyngeal muscular dystrophy characterized by dysphagia and progressive bilateral ptosis
(6) Progressive familial myopathic ptosis and involvement of one, some, or all extraocular (and no other) muscles of one or both eyes
(7) Late spontaneous unilateral ptosis
(8) Amyloid degeneration with involvement of the levator muscle
*(9) Senility-loss of general muscle tone and atrophy of orbital fat
(10) Ptosis and normal pregnancy
(11) Hyperthyroidism and ptosis-following active stages
(12) Drugs, including the following:

<table>
<thead>
<tr>
<th>Drug</th>
<th>Drug</th>
<th>Drug</th>
</tr>
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<tbody>
<tr>
<td>adenine arabinoside</td>
<td>diphtheria and tetanus</td>
<td>nalidixic acid(?)</td>
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<td>adrenal cortex injection</td>
<td>toxoids and pertussis</td>
<td>opium</td>
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<td>(DPT)</td>
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<td>sulphiamide</td>
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<tr>
<td>carbromal</td>
<td>isosorbide dinitrate(?)</td>
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<td>measles virus vaccine</td>
<td>thiamylal</td>
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<tr>
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<tr>
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<tr>
<td>dimethyl tubocurarine</td>
<td>metocurine iodide</td>
<td>vincristine</td>
</tr>
</tbody>
</table>

(13) Corticosteroid ptosis-prolonged use of topical corticosteroid therapy
(14) Mascara ptosis-due to subconjunctival deposits of mascara
(15) Ptosis associated with chronic conjunctivitis and uveitis
(16) Use of botulinum toxin

D. Protective ptosis following injury to the eye
E. Mechanical ptosis
   (1) Tumor
      a. Benign tumor-such as neurofibroma or hemangioma
      b. Malignant tumor-such as basal cell carcinoma, squamous cell carcinoma, malignant melanoma, or rhabdomyosarcoma
      c. Metastatic lesion-such as from breast or lung
      d. Sinus extension-such as mucocele of frontal sinus
   (2) Blepharochalasis-hereditary with recurrent attacks of severe edema and residual damage to the tissues
   (3) Cicatricial ptosis-such as that secondary to cicatricial conjunctivitis (see p. 49-50) or surgical trauma to the superior fornix
   (4) Contact lens migration
   (5) Palpebral form of vernal conjunctivitis
   (6) Intracranial extension-such as chordoma


SYNDROMES AND DISEASES ASSOCIATED WITH PTOSIS

1. Aarskog syndrome (faciogenital dysplasia)-x-linked
2. Acquired immunodeficiency syndrome
3. Addison disease (idiopathic hypoparathyroidism)
4. Alacrima congenital with distichiasis, conjunctivitis, keratitis-autosomal dominant
5. Albers-Schönberg syndrome (marble bone disease)
6. Albright syndrome (osteitis fibrosa disseminata)
7. Amyloidosis (Lubarsch-Pick syndrome)
8. Apert syndrome (acrocephalosyndactyly syndrome)
9. Arteriovenous fistula
10. Axenfeld-Schurenberg syndrome (cyclic oculomotor paralysis)
11. Babinski-Nageotte syndrome (medullary tegmental paralysis)
12. Basal cell carcinoma
13. Bassen-Kornzweig syndrome (abetalipoproteinemia)
14. Bell palsy (idiopathic facial paralysis)
15. Bing-Neel syndrome (Bing disease)
16. Blepharophimosis syndrome
17. Bonnet-Dechaume-Blanc syndrome (neuroretinoangiomatosis)
18. Bonnevie-Ullrich syndrome (pterygolymphangiectasia)
19. Botulism
20. Brown syndrome (superior oblique tendon sheath syndrome)
21. Carpenter syndrome (acrocephalopolysyndactyly II)
22. Cavernous sinus syndrome (Foix syndrome)
23. Cerebral palsy
24. Cestan-Chenaïs syndrome (Cestan syndrome)
25. Chromosome long-arm deletion syndrome
26. Chromosome partial deletion (long-arm) syndrome (de Grouchy syndrome)
27. Chromosome short-arm deletion syndrome
28. Congenital fibrosis syndrome (all extraocular muscles)
29. Congenital fibrosis of the inferior rectus with ptosis-autosomal dominant
*30. Congenital ptosis
  A. Simple failure of peripheral differentiation of muscles-dominant
  B. Ptosis with blepharophimosis-dominant
  C. Ptosis due to ophthalmoplegia-autosomal dominant
31. Craniocarpotarsal dysplasia (Freeman-Sheldon syndrome)
32. Craniocervical syndrome (whiplash injury)
33. Cretinism (juvenile hypothyroidism)
34. Creutzfeldt-Jakob syndrome (spastic pseudosclerosis)
35. Cri-du-chat syndrome (Cry of the cat syndrome)
36. Crouzon syndrome (craniofacial dysostosis)
37. Cushing syndrome (2) (cerebellopontine angle syndrome)
38. Dandy-Walker syndrome (atresia of foramen Magendie)
39. Dawson disease (subacute sclerosing panencephalitis)
40. Dejerine-Klumpke syndrome (lower radicular syndrome)
41. de Lange syndrome (congenital muscular hypertrophy-cerebral syndrome)
42. Devic syndrome (ophthalmoencephalomyelopathy)
43. Diphtheria
44. Dubowitz syndrome (dwarfism-eczema-peculiar facies)
45. Duck-bill lips, low-set ears-autosomal dominant
46. Eaton-Lambert syndrome (myasthenic syndrome)
47. Eclampsia and preeclampsia
48. Ehlers-Danlos syndrome (fibrodyplasia elastic generalisata)
49. Engelmann syndrome (osteopathia hyperostotica scleroticans multiplex infantalis)
50. Epidermal nevus syndrome (ichthyosis hystrix)
51. Erb-Goldflam syndrome (myasthenia gravis)
52. Erysipelas (St. Anthony fire)
53. Fabry syndrome
54. Faciorena1 acromesometic syndrome
55. Fisher syndrome (ophthalmoplegia-ataxia-areflexia syndrome)
56. Fetal alcohol syndrome
57. Fetal trimethadione
58. Foramen lacerum syndrome (aneurysm of internal carotid artery syndrome)
59. Freeman-Sheldon syndrome
60. Garcin syndrome (half-base syndrome)
61. Gerlier disease (paralytic vertigo)
62. Gillum-Anderson syndrome (dominant blepharoptosis, high myopia)
63. Guillain-Barré syndrome (acute infectious neuritis)
64. Hairy elbow syndrome
65. Hemangiomas
66. Herpes zoster
67. Hodgkin disease
68. Horner syndrome (cervical sympathetic paralysis)
69. Hunter syndrome [mucopolysaccharidosis (MPS) II]
70. Hurler disease (MPS I)
71. Hyperammonemia
72. Hyperparathyroidism
73. Hyperthyroidism (Basedow syndrome)
74. Hypocalcemia
*75. Hypoparathyroidism
76. Hysteria
77. Infectious mononucleosis
78. Influenza
79. Jugular foramen syndrome (Vernet syndrome)
80. Kearns-Same syndrome
81. Kiloh-Nevin syndrome (muscular dystrophy of external ocular muscles)
82. Kohn-Romano syndrome (blepharoptosis, blepharophimosis, epicanthus inversus, telecanthus)
83. Komoto syndrome (congenital eyelid tetrad)
84. Krause syndrome (congenital encephaloophthalmic dysplasia)
85. Kugelberg-Welander syndrome (progressive proximal muscle atrophy)
86. Kussmaul disease (necrotizing angiitis)
87. Laurence-Moon-Bardet-Biedl syndrome (retinitis pigmentosa-polydactyly-adiposogenital syndrome)
88. Leigh disease (subacute necrotizing encephalomyelopathy)
89. Little syndrome (nail-patella syndrome)
90. Lymphangioma
91. Lymphedema
92. Malaria
93. Malignant hyperthermia syndrome
94. Maple-syrup urine disease (branched-chain ketoaciduria)
95. Marcus Gunn syndrome (jaw-winking syndrome)
96. Marin Amat syndrome (inverted Marcus Gunn syndrome)
97. MERRF syndrome
98. Micro syndrome
99. Misdirected third nerve syndrome
100. Möbius syndrome (congenital paralysis of the sixth and seventh nerves)
101. Morquio syndrome (keratosulfaturia)
102. Mucormycosis
103. Multiple sclerosis (disseminated sclerosis)
104. Myopathy, centronuclear with external ophthalmoplegia-autosomal dominant
105. Myotonic dystrophy syndrome (Curschmann-Steinert syndrome)
106. Myotubular myopathy-autosomal recessive or x-linked
107. Naffziger syndrome (scalenus anticus syndrome)
108. Neurilemoma
109. Neuroblastoma
110. Neurofibromatosis
111. Nonne-Milroy-Meige disease (congenital trophedema)
112. Noonan syndrome (male Turner syndrome)
113. Oculopharyngeal muscular dystrophy
114. Ophthalmoplegic migrane syndrome
115. Ophthalmoplegic-retinal degeneration (Kearns-Sayre syndrome)
116. Orodigital-facial syndrome (Papillon-Léage and Psaume syndrome)
117. Pachydermoperiostosis (Touraine-Solente-Gole syndrome)
118. Pancoast syndrome (superior pulmonary sulcus syndrome)
119. Parinaud syndrome (paralysis of vertical movements)
120. Parkinson syndrome (paralysis agitans)
121. Parry-Romberg syndrome (progressive facial hemiatrophy)
122. Periocular and ocular metastatic tumors
123. Pierre-Robin syndrome (micrognathia-glossoptosis syndrome)
124. Poliomyelitis
125. Progressive intracranial arterial occlusion syndrome
126. Purpura and ptosis-combined inheritance with male-to-male transmission
127. Raeder syndrome (paratrigrigeminal paralysis)
128. Retraction syndrome (Duane syndrome)-autosomal dominant
129. Retroparotid space syndrome
130. Riley-Day syndrome (congenital familial dysautonomia)
131. Ring D chromosome
132. Rollet syndrome (orbital apex-sphenoidal syndrome)
133. Rubinstein-Taybi syndrome (broad thumbs syndrome)
134. Scleroderma (progressive systemic sclerosis)
135. Scurvy (vitamin C deficiency)
136. Shy-Gonatas syndrome (similar to Hunter and Refsum syndrome)
137. Smith-Lemli-Opitz syndrome (cerebrohepatorenal syndrome)
138. Smith syndrome (facioskeletal-genital dysplasia)
139. Sparganosis


**Extracted Table Ptosis**

**SPECIFIC BLEPHAROPTOSIS**

1. Unilateral ptosis with dilated pupil-tumor or abscess of temporal lobe and third nerve palsy
2. Unilateral ptosis with miosis-midbrain lesion near the posterior commissure and Horner syndrome
3. Ptosis with disturbance of integrated ocular movement-lesion near superior colliculus
4. Bilateral ptosis with small immobile pupils and loss of upward rotation of eyeballs-lesion near posterior commissure
5. Ptosis with loss of voluntary elevation but normal involuntary elevation of the lid when the eye looks up-supranuclear lesion
6. Ptosis in repose and normal elevation with active motion-hereditary cerebellar ataxia of Pierre-Marie
7. Ptosis onset in adolescent-familial chronic external ophthalmoplegia
8. Ptosis may be early and only sign of nuclear paralysis in:
   A. Botulism
   B. Multiple sclerosis (disseminated sclerosis)
   C. Hemorrhagic superior poliomyelitis of Wernicke
   D. Tabes
   E. Vasospasm of ophthalmoplegic migraine
9. Ptosis with cranial nerve dysfunction suggests a basal lesion, such as the following:
   A. Aneurysm
   B. Epidemic paralyzed vertigo (Gerlier disease)
   *C. Herpes zoster
   D. Meningitis
   E. Polyneuritis of cranial nerves
   *F. Trauma
10. Transient ptosis
    A. Acute exanthema
    B. Acute infection such as erysipelas
    *C. Botulinum toxin injection
    D. Eclampsia
    E. Exogenous poisons such as those due to alcohol, lead, carbon monoxide, arsenic, snake venom
    F. Hematoma
    G. Influenza
    H. Scurvy (vitamin C deficiency)
11. Ptosis with orbicularis weakness-muscle disease
12. S-shaped ptosis
    A. Chronic chalazion
    B. Cyst on lateral border of tarsus
    C. Dermoid
    D. Floppy eyelid syndrome
    E. Lacrimal gland enlargement or prolapse
    F. Lateral levator palpebrae superioris muscle dehiscence
    G. Neurofibromatosis
    H. Trachoma
    I. Drugs, including the following:


HORNER SYNDROME

Homer syndrome comprises paralysis of sympathetic nerve supply with lid ptosis, miosis, apparent enophthalmos, frequently dilatation of the vessels with absence of sweating (anhidrosis) on homolateral side; the pupil demonstrates a decreased sensitivity to cocaine and hypersensitivity to adrenalin and may have heterochromia with congenital Homer syndrome.

1. Region of first neuron-lesions of hypothalamus and diencephalic region also suggest diabetes insipidus, disturbed temperature regulation, adiposogenital syndrome, and autonomic epidemic epilepsy of Penfield.
   A. Arnold-Chiari malformation
   B. Basal meningitis, such as in syphilis
   C. Base-of-skull tumors (e.g., melanoma)
   D. Multiple sclerosis
   E. Pituitary tumor
   F. Tumor of the third ventricle
   G. Midbrain, such as in syphilis
   H. Pons, such as in intrapontine hemorrhage
   I. Medulla, such as in Wallenberg syndrome (lateral medullary syndrome)-thrombosis of posterior inferior cerebellar artery
   J. Cervical region
      (1) Syringomyelia
      (2) Tumor
      (3) Injury as traumatic dislocation of cervical vertebrae or dissection of the vertebral artery
      (4) Syphilis (acquired lues)
      (5) Poliomyelitis
      (6) Meningitis
      (7) Amyotrophic lateral sclerosis
      (8) Related to scleroderma and facial hemiatrophy
      (9) Vascular malformation such as agenesis of internal carotid artery

2. Region of second neuron
   A. Spinal birth injury-Klumpke paralysis with injured lower brachial plexus
   B. Cervical rib
   C. Charcot-Tobias syndrome
   D. Thoracic lesions
      (1) Pancoast tumor-in apex of lung, such as carcinoma or tuberculosis
      (2) Aneurysm of aorta, subclavian, or carotid artery
      (3) Central venous catheterization
      (4) Mediastinal tumors
      (5) Lymphadenopathy of Hodgkin disease, leukemia, lymphosarcoma, or tuberculosis
      (6) Stellate ganglion block
(7) Tube thoracostomy

E. Neck

*(1) Enlarged lymph gland, tumors, aneurysm, and thyroid gland
(2) Carcinoma of esophagus
(3) Retropharyngeal tumors
(4) Neuroma of sympathetic chain
(5) Intraoral trauma with damage to internal carotid plexus
(6) Thin intervertebral foramina of spinal cord, such as in pachymeningitis, hypertrophic spinal arthritis, ruptured intervertebral disc, and meningeal tumors
(7) Traction of sternocleidomastoid muscle, such as from positioning on operating table
(8) Complications of tonsillectomy
(9) Mandibular tooth abscess
*(10) Lesions of middle ear, such as in acute purulent otitis media and petromastoid operation
(11) Carotid artery dissection
(12) Internal carotid artery occlusion

3. Region of third neuron

A. Aneurysm of internal carotid and its branches
B. Paratrigeminal syndrome (Raeder syndrome)
C. Cavernous sinus syndrome (Foix syndrome)
D. Tumors of cysts of orbit
E. Drugs can affect any region and include the following:

acetophenazine
alseroxylon
bupivacaine
butaperazine
carphenazine
chloroprocaine
chlorpromazine
deserpidine
diacetylmorphine
diethazine
ethopropazine
etidocaine
fluphenazine
guanethidine
influenza virus vaccine

levodopa
lidocaine
mepivacaine
mesoridazine
methilazaine
methotrimeprazine
oral contraceptives
pericyazine
perphenazine
piperacetazine
prilocaine
procaine
prochlorperazine
promazine

F. Cluster headaches (migrainous neuralgia)
G. Herpes zoster
H. Migraine
I. Fetal varicella syndrome


Extracted Table Horner Syndrome

<table>
<thead>
<tr>
<th>PTOSIS OF LOWER LID (UNCOMMON DROOPING OF LOWER LID SO THAT LID MARGIN IS ADJACENT TO GLOBE BUT BELOW LIMBUS)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Blepharophimosis syndrome</td>
</tr>
<tr>
<td>*2. Cicatricial with mechanical displacement by scar, tumor, or skin disease; may be associated with ectropion</td>
</tr>
<tr>
<td>*3. Paralytic due to lower lid lagophthalmos</td>
</tr>
<tr>
<td>4. Pseudoptosis such as in exophthalmos and higher degrees of myopia</td>
</tr>
<tr>
<td>5. Idiopathic</td>
</tr>
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</table>


LAGOPHTHALMOS (INABILITY TO CLOSE EYELIDS VOLUNTARILY)

*1. Physiologic—many people sleep with their eyes open, especially Asian people
2. Orbital—extreme proptosis
3. Mechanical—scarring of the lids or retractor muscles
4. Paralytic
   A. Seventh nerve palsy (see p. 66)
   B. Leprosy
   C. Lesions of cerebral cortex and its projections, including bilateral frontal lesions
5. Psychological
   A. Failure to comprehend the command
   *B. Unwillingness to comply with the command


PSEUDO-LID RETRACTION
1. Exophthalmos
2. Unilateral high axial myopia
3. Unilateral congenital glaucoma
4. Congenital cystic eyeball
5. Abnormalities of orbit
   A. Asymmetry
   B. Shallow such as in Crouzon disease (dysostosis craniofacialis)
   C. Harlequin-shallow orbit with arched superior and lateral wall, such as in hypophosphatasia
6. Ptosis of other eyelid


**LID RETRACTION**

Lid retraction is defined normally as more than 85% of vertical palpebral fissures and 10 mm or less with the eyelids just concealing the corneoscleral limbus at the 12 and 6 o’ clock meridians.

1. Lid retraction with upward movement of eye
   A. Congestive dysthyroid disease
   B. Deficiency in upward gaze-following rectus operation or weakness of superior rectus
   C. Excessive stimulation of levator muscles in Bell phenomenon with seventh nerve palsy
   D. Levator muscles receive excessive stimuli from nerve fiber of superior rectus
   E. Pretectal or periaqueductal lesion in midbrain
2. Lid retraction with downward movement of eye
   A. Aberrant regeneration of third nerve of inferior rectus to levator (pseudo-Graefe phenomenon) - elevation of lid in downward gaze
   B. Brown syndrome (superior oblique tendon sheath syndrome)
   C. Extrapyramidal syndrome of postencephalic parkinsonism and progressive supranuclear palsy
   D. Failure of levator to relax on downward movement of eye
      (1) Secondary neuromuscular
      *(2) Mechanical, such as from a scar
   E. Noncongestive type of dysthyroid exophthalmos (Graefe sign)-lid lag in downward gaze
3. Lid retraction with horizontal gaze
   A. Duane syndrome (retraction syndrome)
   B. Underaction of lateral rectus muscle and spillover to levator causing widening
4. Lid retraction because of supranuclear lesions-usually bilateral when due to lesion in or about posterior comissure (Collier sign, tucked lids, posterior fossa stare)
A. Bulbar poliomyelitis
B. Chorea (Huntington hereditary chorea)
C. Closed head injury associated with defective adduction of eyes, coarse nystagmus, nuclear palsy, pyramidal signs
D. Coma due to disease of ventral midbrain and pons
E. Craniostenosis
F. Epidemic encephalitis
G. Hydrocephalic infants
H. Hydrophobia
I. Hysteria
J. Malingering
K. Meningitis
L. Multiple sclerosis (disseminated sclerosis)
M. Parinaud syndrome (divergence paralysis)
N. Parkinson disease (paralysis agitans)
O. Russell syndrome
P. Sylvian aqueduct syndrome (Koerber-Solus-Elschnig syndrome)
Q. Syphilis (tabes)
R. Tumors of the midbrain; meningiomas of sphenoid wing; sellar, parasellar, and suprasellar tumors; and frontal or temporal lobe tumors
S. von Economo syndrome (encephalitis lethargica a)

5. Lid retraction because of neuromuscular disease-commonly asymmetric or unilateral
   A. Drugs
      (1) Phenylephrine and other sympathomimetics
      (2) Prostigmin and Tensilon, especially with myasthenic levator involvement
      (3) Succinylcholine, subparalytic doses
      (4) Thyroid extract
   B. Fuch phenomenon-healing of injured third nerve, previously ptotic lid has involuntarily spastic raising with movements of eyes
   C. Infant lid retraction-transient because of maternal hyperthyroidism
   D. Irritation of cervical sympathetic nerve (Homer syndrome)
   *E. Mechanical suspension of lid such as that due to scar, tumor, surgical attachment to frontalis muscle, or shortening of levator muscle or following glaucoma filtering procedures
   F. Peripheral seventh nerve paresis with loss of orbicularis oculi muscle tone

6. Lid retraction with myopathic disease
   A. Associated with hepatic cirrhosis
   B. Thyroid myopathy (Graves disease, Basedow syndrome)
      (1) Dalrymple sign-widening of palpebral fissure
      (2) Stellwag sign-retraction of upper lid associated with infrequent or incomplete blinking

7. Lid retraction following operations on vertical muscles, such as recession of superior rectus muscle or simultaneous recession and restriction of the levator by common fascial check ligament between the two muscles

8. Paradoxical lid retraction because of paradoxical levator innervation
A. Defective ocular abduction with abducens palsy
B. Lid retraction associated with ptosis of the opposite eyelid (levator denervation supersensitivity)
C. Misdirection of third nerve axons (following acquired or congenital lesions)-occurs on attempt to adduct, elevate, or depress eye
D. Movement of lower jaw
   (1) Contraction of external pterygoid muscle by opening mouth (Marcus Gunn)
   (2) Contraction of internal pterygoid muscle by closing the mouth

9. Physiologic
   A. Act of surprise
   B. Slow onset of blindness, such as that secondary to glaucoma and optic atrophy
   C. Time of attention


LID LAG

Lig lag is defined as occurring when the patient looks down and the eyelids lag behind briefly.

1. Congenital-usually in association with congenital ptosis
2. Hepatic failure
3. Iatrogenic-following ptosis surgery
4. Mechanical-scars of the upper lid
5. Myopathic disease
   *A. Graefe sign-thyroid myopathy-the upper lid pauses and then follows the eye downward (Basedow syndrome)
   B. Myotonic dystrophia
   C. Periodic myotonic lid lag-familial (hyperkalemic) myotonic periodic paralysis
6. Neuromuscular disease
   A. Excessive intake of thyroid extract
   B. Physiologic lagophthalmos-short upper tarsus in some Asian and some white persons with incomplete descent of the lid during sleep
7. Supranuclear origin-extrapyramidal syndromes have defective inhibition of lids in downward gaze
A. Congenital supranuclear lid lag
B. Guillain-Barré syndrome
C. Postencephalitic parkinsonism, Parkinson syndrome (shaking palsy)
D. Progressive supranuclear palsy


**BLEPHAROSPASM (SPASMODIC EYELID CLOSURE)**

Most common and important: Psychogenic onset commonly in children and young adults

1. Addison disease (adrenal cortical insufficiency)
2. Associated with syphilis, tetanus, and tetany
3. Basal ganglion dysfunction-onset usually after middle age; including Parkinson disease (shaking palsy)
4. Cerebral palsy
5. Cogan syndrome (nonsyphilitic interstitial keratitis) with vestibuloauditory symptoms
6. Drugs, including the following:
   - acetophenazine
   - amitriptyline
   - amodiaquine
   - amoxapine(?)
   - amphetamine
   - antazoline
   - brompheniramine
   - butaperazine
   - carbinoxamine
   - carphenezine
   - chloroquine
   - chlorpheniramine
   - chlorpromazine
   - clemastine
   - clomipramine
   - desipramine
   - dexbrompheniramine
   - dexchlorpheniramine
   - dextroamphetamine
   - dextrothyroxine
   - diethazine
   - dimercaprol
   - dimethindene
   - diphenhydramine
   - diphenylpyraline
   - doxepin
   - doxylamine
   - dronabinol
   - droperidol
   - emetine
   - ethopropazine
   - fluphenazine
   - haloperidol
   - hashish
   - hydroxychloroquine
   - imipramine
   - levodopa
   - levotheroxyine
   - liothyronine
   - liotrix
   - lorazepam
   - marihuana
   - mesoridazine
   - methamphetamine
   - methdilazine
   - methotrimeprazine
   - nortriptyline
   - pentylenetetrazol
   - perazine
   - pericyazine
   - pheniramine
   - phenmetrazine
   - phenylephrine
   - piperacetazine
   - prochlorperazine
   - promazine
   - promethazine
   - propiomazine
   - protriptyline
   - pyrilamine
   - selegiline
   - tetrahydrocannabinol
thiethylperazine  thiopropazate  thioproperazine  thioridazine  thyroglobulin
trifluoperazine  trifluperidol  triflupromazine  trimipramine  tripelennamine
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7. Electrical injury
8. Encephalitis
9. Epidemic keratoconjunctivitis
10. Hallervorden-Spatz
11. Hereditary reflex blepharospasm
12. Idiopathic (essential)
13. Leprosy (Hansen disease)
14. Meige syndrome
15. Obsessive-compulsive disorder
*16. Pain or light sensitivity following injury or inflammation or foreign bodies of lids, conjunctiva, cornea, or iris
17. Photosensitivity and sunburn
18. Poison ivy dermatitis
19. Postencephalitic blepharospasm
20. Psychogenic obsessive-compulsive disorder-onset commonly in children and young adults
21. Psychologic reflex blepharospasm-seen in premature infants with tactile stimulation of lids
22. Sparganosis
23. Systemic scleroderma (progressive systemic sclerosis)
24. Thomsen syndrome (congenital myotonia syndrome)
25. Tourette syndrome (coprolalia, generalized tic)


Facial palsy is defined as paralysis of facial muscles supplied by the seventh nerve; orbicularis oculi paralysis may result in epiphora and ectropion.

1. Congenital
2. Birth injury with nerve crushed at exit of stylomastoid foramen
3. Myogenic paralysis
   A. Myotonic atrophy
   B. Facioscapulohumeral type of muscular dystrophy
   C. Myasthenia gravis (Erb-Goldflam syndrome)
   D. Hypokalemia, periodic
   E. Curare poisoning
   F. Botulism
   G. Congenital facial diplegia (Möbius syndrome)
   H. Infants, from maternal ingestion of thalidomide
   I. Kugelberg-Welander syndrome
4. Neurologic paralysis
   A. Supranuclear paralysis-upper face, including orbicularis relatively unaffected with affected lower face
      (1) Voluntary movement-pyramidal fibers involved, such as in Weber syndrome, with contralateral hemiplegia of face and limbs and ipsilateral oculomotor paralysis
      (2) Weakness or abolition of the emotional movements of the face with retention of full voluntary activity, such as with lesion of anterior part of frontal lobe or near optic thalamus
   B. Peripheral paralysis-involvement of upper and lower face
      (1) Pontine lesion-associated structures involved include sixth nerve, conjugate ocular deviation to the same side, ipsilateral paralysis of jaw muscles, and pyramidal tract in paralysis of limb of opposite side
         a. Acute nuclear lesions, such as with anterior poliomyelitis, Landry paralysis, or degenerative conditions
         b. Foville syndrome-ipsilateral sixth nerve with loss of conjugate deviation to same side and hemiplegia of the opposite limbs
         c. Millard-Gubler syndrome-ipsilateral sixth nerve paralysis and hemiplegia of the opposite limbs
         d. Parotid gland surgery
         e. Progressive muscular atrophy
         f. Syringobulbia
         g. Tumors
         h. Vascular lesions
      (2) Posterior fossa-associated with nerve deafness, loss of taste on anterior two thirds of tongue, and occasionally diminution of tears
         a. Acoustic neuroma
         b. CHARGE (coloboma, heart disease, atresia choanae, retarded growth and retarded development or central nervous system anomalies, genital hypoplasia, and ear anomalies or deafness) syndrome association
c. Facial neuritis due to polyneuritis cranialis, beriberi, encephalitis, diabetes, or intrathecal anesthesia
d. Fracture of the skull
e. Meningitis, including syphilitic and tuberculous
f. Preauricular cyst associated with congenital cholesteatoma
g. Tumors of facial nerve

(3) Petrous temporal bone-associated with decreased lacrimation and salivary secretion, loss of taste on anterior two thirds of tongue, and intensified sensation of loud noises

*a. Arteriosclerosis
*b. Bell palsy-inflammation of facial nerve of unknown cause
c. Cephalic tetanus
d. Diabetes mellitus (Willis disease)
e. Fractures
f. Herpes zoster, spread from geniculate ganglion
g. Hypertension

* = most important

h. Nerve leprosy (Hansen disease)
i. Otitis media
j. Secondary syphilis

(4) Facial lesions at or beyond the stylomastoid foramen

a. Fracture of the ramus of the mandible
b. Melkersson-Rosenthal syndrome (Melkersson idiopathic fibroedema)
c. Neoplasia or inflammatory swelling of parotid, such as in uveoparotid fever (Heerfordt disease) and Mikulicz disease
d. Supporting lymph nodes behind the angle of the jaw


**INFREQUENT BLINKING**

*1. Contact lens use
2. Encephalitis, acute
3. Encephalitis or mild postencephalitic states
*4. Ethanol intake
5. Infants in first few months of life
6. Parkinson syndrome, including myostatic paresis of parkinsonism
7. Psychotic states
8. Progressive supranuclear palsy
9. Thyrotoxicosis including exophthalmic ophthalmoplegia (Stellwag sign)


**FREQUENT BLINKING**

*1. Reflex—strong lights, sudden approach of objects toward eyes, loud noises, and touching the cornea; reflex blinking common in albinos and light intolerance

2. Spontaneous—mental state and environment
   A. Children with habit spasm and facial tic
   B. Blepharospasm
   *C. Older persons with inadequate lacrimation and local irritation of the eyes

3. Disorders of central nervous system disease, such as parkinsonism or various forms of pseudobulbar palsy

4. Drugs, including the following:

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**LID EDEMA (PUFFINESS OR BAGGINESS OF LIDS)**

*1. Noninflammatory or minimally inflammatory swelling
A. Acosta syndrome (Mountain climbers syndrome)
B. Allergic gastroenteropathy with protein loss
C. Arteriovenous fistula
D. Cardiac and renal disease
   (1) Nephrosis and acute glomerulonephritis—early morning edema
   (2) Starvation and cachexia
E. Dermatochalasis
F. Elephantiasis
   (1) Chronic eczema or infection (erysipelas)
   (2) Hemolymphangioma
   (3) Leprosy (Hansen disease)
   (4) Lues (syphilis)
   (5) Melkersson-Rosenthal syndrome (Melkersson idiopathic fibroedema)
   (6) Nonne-Milroy-Meige disease (idiopathic hereditary lymphedema)
   (7) von Recklinghausen disease (neurofibromatosis)
   (8) Traumatic disruption of the lymph drainage system
   (9) Tuberculosis
G. Endocrine exophthalmos (hyperthyroidism)
H. Foix syndrome (cavernous sinus syndrome)
I. Granulomatous ileocolitis
J. Hutchinson syndrome (adrenal cortex neuroblastoma with orbital metastasis)
K. Infectious generalized diseases
   (1) Diphtheria
   (2) Infectious mononucleosis
   (3) Malaria
   (4) Meningococcal meningitis
   (5) Pertussis (whooping cough)
   (6) Rheumatic fever
   (7) Scarlet fever
   (8) Trypanosomiasis
   (9) Tuberculosis
   (10) Yellow fever
L. Melkersson-Rosenthal syndrome
M. Nasal nerve syndrome (Charlin syndrome)
N. Parasitic infestations
   (1) Anthrax
   (2) Ascariasis
   (3) Chlamydia
   (4) Dermatophytosis
   (5) Myiasis
   (6) Onchocerciasis syndrome (river blindness)
   (7) Tapeworms
   (8) Toxocariasis
   (9) Trichinosis
O. Protrusion of fat through orbital fascia
P. Retinoblastoma
Q. Stasis, including premenstrual edema
R. Superior vena cava syndrome
S. Systemic scleroderma (progressive systemic scleroderma)
*T. Tumors and pseudotumors
   (1) Benign and malignant ectodermal and mesodermal tumors
   (2) Brill-Symmers disease (lymphosarcoma)
   (3) Hemangiomas
   (4) Hodgkin disease
   (5) Leukemic deposit
   (6) Liposarcoma
   (7) Meningiomas of sphenoid ridge with impediment of venous circulation
       of ophthalmic veins or cavernous sinus
   (8) Neurofibromatosis
   (9) Pseudotumors
      a. Amyloid degeneration
      b. Eosinophilic or basophilic granulomas
U. Trauma
   (1) Basilar skull fractures
   (2) Injury
      *(3) Surgery
V. Angioneurotic edema caused by drugs, including the following:
   acenocoumarol  antimony sodium tartrate  bupivacaine
   acetaminophen  antimony sodium  busulfan
   acetanilid  thioglycolate  butobarbital
   acetophenazine  antipyrine  butalbital
   acetyldigitoxin  aprobarbital  butallylonal
   acyclovir  aspirin  butaperazine
   adrenal cortex injection  atropine  butethal
   albuterol  auranofin  cactinomycin
   alcohol  aurothioglucose  capreomycin
   aldosterone  aurothioglycanide  captopril
   allobarbitual  azatadine  carbamazepine
   alprazolam  bacitracin  carbencilllin
   aluminum nicotinate  barbital  carbimazole
   aminosalicylalate(?)  belladonna  carbenoxamine
   aminosalicylic acid(?)  bendroflumethiazide  carisoprodol
   amiiodarone  benzalkonium  carphenazine
   amitriptyline  benzathine penicillin G  cefaclor
   amobarbital  benzphetamine  cefadroxil
   amoxapine  benzthiazide  cefamandole
   amoxicillin  betamethasone  cefazolin
   ampicillin  betaxolol  cefonicid
   anisindicone  bleomycin  cefoperazone
   antazoline  botulinum A toxin  ceforanide
   antimony lithium thiomalate  brimonidine tartrate  cefotaxime
   antimony potassium tartrate  brompheniramine  cefotetan
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</table>
hydralazine
hydrocortisone
ibuprofen
IDU (idoxuridine)
imipramine
indomethacin
insulin
iodide and iodine solution
and compounds
iodipamide meglumine
iothalamate meglumine
iron dextran
isoniazid
isopropyl unoprostone
isosorbide
isotretinoin
ketoprofen
latanoprost
leuprolide acetate
levodopa
levothyroxine
lidocaine
lincomycin
liothyrone
liotrix
lithium carbonate
lorazepam
loxapine
mannitol
maprotiline
measles and rubella virus
vaccine (live)
measles virus vaccine (live)
measles, mumps, and rubella
virus vaccine (live)
mecamylamine
medroxyprogesterone
medrysone
mefenamic acid
melphalan
mephenytoin
mephobarbital
mepivacaine
meprednisone
meprobamate
mercuric oxide
mesoridazine
methacycline
metharbital
methdilazine
methicillin
methitural
methohexital
methotrimeprazine
methsuximide
methyl scopolamine
methyldopa
methylene blue
methylergonovine
methylprednisolone
methysergide
metoclopramide
metrizamide
mexiletine
mianserin
midazolam
minocycline
mitomycin
moxalactam
nafcillin
nalidixic acid
naltrexone
naproxen
niacin
niacinamide
nicotinic acid
nicotinyl alcohol
nifedipine
nitrazepam
nitrofurantoin
nitremersol of estrogens
and progestogens
olapataidine hydrochloride
oral contraceptives
ouabain
oxacillin
oxprenolol
oxyphenbutazone
oxytetracycline
paramethadione
paramethasone
penicillin
pentazocine
pentobarbital
perazine
pericyazine
perphenazine
phenacetin
phenobarbital
phenoxyethyl
penicillin
phensuximide
phenylbutazone
phenylmercuric acetate
phenylmercuric nitrate
pimozide
piperacazine
piperazine
piroxicam
poliovirus vaccine
polymyxin B prazosin
potassium penicillin G
potassium penicillin V
potassium phenethicillin
potassium
phenoxyethyl
prazepam
prednisolone
prednisone
primidone
probartial procaine
penicillin G
procaine
prochlorperazine
promazine
promethazine
propiomazine
propofol
propoxycline
protriptyline
quinacrine
quinidine
quinine
rabies immune globulin
rabies vaccine
2. Inflammatory edema

A. Acute hemorrhagic conjunctivitis
B. Allergic eczema (contact dermatitis)
   (1) Anesthetics
   *(2) Atropine (topical)
   (3) Eczematous keratoconjunctivitis
   (4) Iodine
   (5) Mercury
   (6) Neurodermatitis
   (7) Penicillin
   (8) Photodermatitis
   (9) Quincke edema
   (10) Tuberculosis (scrofula)
C. Anthrax
D. Bee sting of the cornea
E. Dacryoadenitis
   (1) Acute dacryoadenitis
   (2) Chronic dacryoadenitis
F. Epidemic keratoconjunctivitis
G. Erysipelas
H. Herpes simplex
I. Hollenhorst syndrome (chorioretinal infarction syndrome)
J. Hordeolum, chalazion
K. Lymphogranuloma venereum
L. Mycoses
M. Ophthalmic zoster
N. Other causes of lid edema
   (1) Conjunctival inflammations
      a. Diphtheria
      b. Newcastle disease (fowlpox)
      c. Ophthalmia neonatorum
      d. Parinaud syndrome (conjunctiva-adenitis syndrome)
   (2) Keratitis
   (3) Orbital inflammation
   (4) Periostitis
   (5) Scleritis (see p. 237-239)
      a. Posterior scleritis
      b. Scleromalacia perforans
   (6) Thermal, chemical, mechanical, or radiation injury
      a. Hypothermal injury
      b. Polychlorinated biphenyl (PCB)
O. Scalded skin syndrome (Ritter disease)
P. Serum sickness-systemic reaction to foreign serum, serum products, vaccines, penicillin, and sulfa drugs
Q. Silverman syndrome (battered-baby syndrome)
R. Spider bites
S. Urticaria due to drugs, including the following:

acenocoumarin  antinomy sodium  butallylonal
acetaminophen  thioglycollate  cactinomycin
acetanilid  antipyrine  capreomycin
acetazolamide  aprobarbital  captropil
acyclovir  aspirin  carbamazepine
albuterol  auranofin  carbenicillin
allobarbital  aurothioglucose  carbimazole
allopurinol  aurothioglycanide  carbinoxamine
alprazolam  azatadine  carisoprodol
aluminum nicotinate  Bacille Calmette-Guérin  cefaclor
amiodarone  (BCG) vaccine  cefadroxil
amitriptyline  bacitracin  cefamandole
amobarbital  barbital  cefazolin
amoxapine  bendroflumethiazide  cefonicid
amoxicillin  benzathine penicillin G  cefoperazone
ampicillin  benzphetamine  ceforanide
anisindione  benzthiazide  cefotaxime
antazoline  betamethasone  cefotetan
antimony lithium  betaxolol  cefoxitin
   thiomalate  bleomycin  cefsulodin
antimony potassium  brompheniramine  ceftazidine
tartrate  bupivacaine  cefitoxime
antimony sodium  busulfan  ceftriaxone
tartrate  butabarbital  cefuroxime
   butalbital  cephalaxin
cephaloglycin
dexchlorpheniramine
dexchlorpheniramine
flecainide
cephaloridine
dexethal
dexchlorpheniramine
fluorescein
cephalothin
dexamethasone
fluorouracil
cepapirin
dexamethasone
flurazepam
cepradine
dexamethasone
flurbiprofen
chlorambucil
dexamethasone
framycetin
chloramphenicol
dexamethasone
furosemide
chlorodiazepoxide
dexamethasone
gentamicin
chlorpromazine
dexamethasone
glutethimide
chlorothiazide
dexamethasone
gold Au 198
clophenteramine
dexamethasone
gold sodium thiomalate
clophenthixene
dexamethasone
gold sodium thiosulfate
clofibrate
dexamethasone
griseofulvin
clofibrate
dexamethasone
hetacillin
clofibrate
dexamethasone
hexethal
clofibrate
dexamethasone
hexobarbital
clofibrate
dexamethasone
hydramidine penicillin
clomiphene
dexamethasone
hydromorphone
clonazepam
dexamethasone
ibuprofen
clobazepam
dexamethasone
imipramine
clofibrate
dexamethasone
indomethacin
clofibrate
dexamethasone
influenza virus vaccine
clofibrate
dexamethasone
insulin
clofibrate
dexamethasone
interferon
clofibrate
dexamethasone
iodide and iodine
clomipramine
dexamethasone
solutions and
clonopine
dexamethasone
compounds
clophenacine
dexamethasone
iodipamide meglumine
clonazepam
dexamethasone
iodine and iodine
clonazepam
dexamethasone
iophendylate
clonazepam
dexamethasone
iothermal meglumine
clonazepam
dexamethasone
and sodium
clorazepate
dexamethasone
iothermal acid
clofibrate
dexamethasone
iron dextran
clofibrate
dexamethasone
isoniazid
clofibrate
dexamethasone
isosorbide
clofibrate
dexamethasone
isotretinoin
clofibrate
dexamethasone
ketoprofen
labetalol
levallorphan
levobunolol
lidocaine
lincomycin
lorazepam
loxapine
mannitol
maprotiline
measles and rubella virus vaccine (live)
measles virus vaccine (live)
measles, mumps, and rubella virus vaccine (live)
melphalan
meperidine
meephytoin
mephobarbital
mepivacaine
meprobamate
mercuric oxide
methacycline
methadone
methaqualone
metharbital
methazolamide
methicillin
methimazole
methitural
methocarbamol
methohexital
methotrexate
methyclothiaizide
methyldopa
methylphenidate
methylprednisolone
methylthiouracil
methyprylon
metcloprimamide
metocurine iodide
metolazone
metoprolol
metrizamide
metronidazole
mianserin
midazolam
minocycline
mitomycin
morphine
moxalactam
mumps virus vaccine (live)
nafcillin
nalidixic acid
nalorphine
naloxone
naltrexone
naproxen
neomycin
neostigmine
niacin
niacinamide
nicotinyl alcohol
nifedipine
nitrazepam
nitrofurantoin
nitromersol
nortriptyline
opium
oral contraceptives
oxacillin
oxazepam
oxymorphine
oxyphenbutazone
oxytetracycline
penicillamine
pentazocine
pentobarbital
phenacetin
phendimetrazine
phenindione
pheniramine
phenobarbital
phenprocoumon
phentermine
phenylbutazone
phenylmercuric acetate
phenylmercuric nitrate
piperazine
piroxicam
poliovirus vaccine
polythiazide
potassium penicillin G
potassium penicillin V
potassium phenethicillin
practolol
prazepam
prazosin
prilocaine
primidone
probarbital
procaine
procaine penicillin G
procarbazine
propracaicne
propranolol
propylthiouracil
protriptyline
pyrilamine
quinacrine
quinethazon
quinidine
quinine
rabies immune globulin
rabies vaccine
radioactive iodides
ranitidine
rifampin
rubella and mumps virus vaccine (live)
rubella virus vaccine (live)
secobarbital
smallpox vaccine
sodium antimonylgluconate
sodium salicylate
stibocapate
stibogluconate
stibophen
streptomycin
succinyllchoeline
sulfacetamide
sulfachlorpyridazine
sulfacytine
sulfadiazine
sulfadimethoxine  sulfamerazine  sulfameter  sulfamethazine  sulfamethizole  sulfamethoxazole  sulfamethoxypyridazine  sulfanilamide  sulfaphenazole  sulfapyridine  sulfasalazine  sulfathiazole  sulfisoxazole

T. Wegener syndrome (Wegener granulomatosis)
U. Vaccination
   (1) Ocular vaccina
   (2) Postvaccinial ocular syndrome
   (3) Variola


**BLEEDING OF THE EYELID**

1. Drugs, including the following:

   acetazolamide  aurothioglycanide  chlorpropamide
   acethohexamide  BCG vaccine  chlorthalidone
   allopurinol  bendroflumethiazide  cimetidine
   alprazolam  benzthiazide  clofibrate
   amantadine  betamethasone  clonazepam
   amitriptyline  betaxolol  clorazepate
   aspirin  carbamazepine  cyclothiazide
   auranofin  chlordiazepoxide  cytarabine
   aurothioglucose  chlorothiazide  danazol
<table>
<thead>
<tr>
<th>Drug Name</th>
<th>Other Names</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dapsone</td>
<td>Desipramine, Dexamethasone, Etomidate</td>
</tr>
<tr>
<td>Desipramine</td>
<td>Dexamethasone (live), Methaqualone</td>
</tr>
<tr>
<td>Dexamethasone</td>
<td>Desipramine (live), Dexamethasone (live)</td>
</tr>
<tr>
<td>Diazepam</td>
<td>Dexamethasone (live), Methaqualone</td>
</tr>
<tr>
<td>Dichlorphenamide</td>
<td>Methazolamide, Sulfacetamide</td>
</tr>
<tr>
<td>Diltiazem</td>
<td>Methyclothiazide, Sulfachlorpyridazine, Sulfacytine</td>
</tr>
<tr>
<td>Emetine</td>
<td>Methylprednisolone, Sulfadiazine</td>
</tr>
<tr>
<td>Ethozolamide</td>
<td>Methyprylon, Sulfadiazine</td>
</tr>
<tr>
<td>Flurazepam</td>
<td>Metolazone, Sulfadimethoxine</td>
</tr>
<tr>
<td>Furosemide</td>
<td>Metoprolol, Sulfamerazine</td>
</tr>
<tr>
<td>Glutethimide</td>
<td>Methylprednisolone, Sulfamerazine</td>
</tr>
<tr>
<td>Gold Au 198</td>
<td>Naproxen, Sulfamethazine</td>
</tr>
<tr>
<td>Gold Sodium Thrimalate</td>
<td>Nifedipine, Sulfamethoxazole</td>
</tr>
<tr>
<td>Gold Sodium Thiosulfate</td>
<td>Nitrazepam, Sulfamethoxypyrindazine</td>
</tr>
<tr>
<td>Halazepam</td>
<td>Nor triptyline, Sulfanilamide</td>
</tr>
<tr>
<td>Hydrochlorothiazide</td>
<td>Oxazepam, Sulfaphenazole</td>
</tr>
<tr>
<td>Hydrocortisone</td>
<td>Oxprenolol, Sulfapyridine</td>
</tr>
<tr>
<td>Hydroflumethiazide</td>
<td>Phenytoin, Sulfasalazine</td>
</tr>
<tr>
<td>Ibuprofen</td>
<td>Piperazine, Sulfathiazole</td>
</tr>
<tr>
<td>Imipramine</td>
<td>Piroxicam, Sulfisoxazole</td>
</tr>
<tr>
<td>Indapamide</td>
<td>Polythiazide, Temazepam</td>
</tr>
<tr>
<td>Indomethacin</td>
<td>Prazepam, Tetanus immune globulin</td>
</tr>
<tr>
<td>Influenza Virus Vaccine</td>
<td>Procarbazine, Tetanus toxoid</td>
</tr>
<tr>
<td>Interferon</td>
<td>Propranolol, Timolol</td>
</tr>
<tr>
<td>Ketoprofen</td>
<td>Protriptyline, Tolazamide</td>
</tr>
<tr>
<td>Levobunolol</td>
<td>Quinethazone, Tolbutamide</td>
</tr>
<tr>
<td>Lorazepam</td>
<td>Quinine, Triamcinolone</td>
</tr>
<tr>
<td>Measles and Rubella Virus Vaccine (Live)</td>
<td>Rifampin, Triazole</td>
</tr>
<tr>
<td>Measles Virus Vaccine (Live)</td>
<td>Rubella and Mumps Virus, Triazolome</td>
</tr>
<tr>
<td></td>
<td>vaccine (live), Verapamil</td>
</tr>
</tbody>
</table>

2. Hutchinson syndrome (adrenal cortex neuroblastoma with orbital metastasis) *3. Trauma


**ECTROPION (LID MARGIN TURNED OUTWARD FROM THE EYEBALL)**

1. Congenital ectropion
   A. With distichiasis
   B. With tight septum; microblepharon
   C. With partial coloboma
   D. With mandibulofacial dysostosis (Franceschetti syndrome)
   E. With megaloblepharon (euryblepharon)
F. With microphthalmos or buphthalmos
G. Cerebrooculofacioskeletal syndrome
H. Down syndrome (mongolism)
I. Hartnup syndrome (niacin deficiency)
J. Lowe syndrome (oculocerebrorenal syndrome)
K. Miller syndrome
L. Milroy disease (oromandibular dystonia)
M. Robinow syndrome
N. Sjögren-Larsson syndrome

2. Acquired ectropion
   A. Spastic ectropion
      *(1) Acute spastic ectropion
      (2) Blepharophimosis syndrome
      *(3) Chronic spastic ectropion becoming cicatricial ectropion
      (4) Hypothermic injury
      (5) Myasthenia gravis-afternoon onset (Erb-Goldflam syndrome)
      (6) Siemen syndrome (hereditary ectodermal dysplasia syndrome)
   B. Atonic ectropion
      (1) Anophthalmic socket
      (2) Bell palsy (Idiopathic facial paralysis)
      (2) Guillain-Barré syndrome (acute infectious neuritis)
      (3) Paralytic ectropion-lagophthalmos, such as in seventh nerve palsy
      *(4) Senile ectropion-tissue relaxation
   C. Cicatricial ectropion
      (1) Amendola syndrome
      (2) Blastomycosis
      (3) Collodion baby syndrome (congenital ichthyosis)
      (4) Chronic dermatitis
      (5) Cutaneous T-cell
      (6) Etretinate therapy
      (7) Excessive skin excision
      (8) Facial burns and scarring
      (9) Hydroa vacciniforme
      (10) Kabuki makeup syndrome
      (11) Leprosy (Hansen disease)
      (12) Orbital fracture repair
      (13) Palmoplantar keratodermia
      (14) Postblepharoplasty ectropion
      (15) Psoriasis (psoriasis vulgaris)
      (16) Radiation
      (17) Sézary syndrome (malignant cutaneous reticulosis syndrome)
      (18) Systemic fluorouracil
      (19) Thermal burns
      (20) Trauma
      (21) Transformation from chronic spastic ectropion
(22) Zinsser-Engman-Cole syndrome (dyskeratosis congenita with pigmentation)

D. Allergic ectropion-anaphylactic, contact, and microbial (usually temporary)
   (1) Danbolt-Closs syndrome (acrodermatitis enteropathica)
   (2) Elschnig syndrome

E. Mechanical
   (1) Kaposi disease (multiple idiopathic hemorrhagic sarcoma)
   (2) Leiomyoma
   (3) Lumps (chalazion, cysts, neurofibroma)


**ENTROPION (INVERSION OF LID MARGIN)**

1. Congenital, including congenital epiblepharon-inferior oblique insufficiency; ectrodactyly, ectodermal dysplasia, cleft lip-palate syndrome, including with and without lower eyelid retractor insertion
   A. Inferior oblique insufficiency syndrome
   B. Dental-ocular-cutaneous syndrome
   C. Siemen syndrome (anhidrotic ectodermal dysplasia)

2. Acquired
   *A. Spastic entropion-acute, affecting lower lid, precipitated by acute inflammation or prolonged patching
   B. Mechanical entropion
      (1) Anophthalmos
      (2) Enophthalmos
      (3) Microphthalmos
      (4) Lymphedema
   *C. Senile entropion-relative enophthalmos secondary to fat atrophy
   D. Cicatricial entropion-physical and chemical burns of conjunctiva and cicatrizing diseases, including trachoma and leprosy
      (1) Chronic cicatricial conjunctivitis
      (2) Leprosy (Hansen disease)
      (3) Radiation
      (4) Thermal burns
      (5) Trachoma
(6) Following cryosurgery of the eyelid  
(7) Amendola syndrome  
(8) Variola


**EPICANTHUS (FOLD OF SKIN OVER INNER CANTHUS OF EYE)**

1. Types  
   A. Epicanthus inversus-fold arises in the lower lid and extends upward to a point slightly above the inner canthus; it is accompanied by long medial canthal tendons, blepharophimosis, and ptosis-autosomal dominant  
   *B. Epicanthus palpebralis (common type)-epicanthal fold arises from the upper lid above the tarsal region and extends to the lower margin of the orbit  
   C. Epicanthus supraciliaris (unusual type)-epicanthal fold arises near brow and runs toward tear sac  
   *D. Epicanthus tarsalis (Mongolian eye)-epicanthal fold arises from the tarsal (lid) fold and loses itself in the skin close to the inner canthus-autosomal dominant

2. Associated conditions  
   A. Aminopterin-induced syndrome  
   B. Basal cell nevus syndrome (Gorlin syndrome)  
   C. Basset-Kornzweig syndrome (familial hypolipoproteinemia)  
   D. Bilateral renal agenesis  
   E. Blepharophimosis, ptosis, epicanthus inversus syndrome  
   F. Bonnevie-Ullrich syndrome (pterygolymphangiectasia)  
   G. Carpenter syndrome (acrocephalopolysyndactyly II)  
   H. Cat-eye syndrome (partial G-trisomy syndrome)  
   I. Cerebrofacioarticular syndrome of van Maldergen  
   J. Cerebrohepatorenal syndrome (Smith-Lemli-Opitz syndrome)  
   K. Chondrodystrophy (Conradi syndrome)  
   L. Chromosome long-arm deletion syndrome  
   M. Chromosome deletion (deletion 18)  
   N. Chromosome partial short-arm deletion syndrome (Wolf syndrome)  
   O. Chromosome short-arm deletion syndrome  
   P. Chromosome 13q partial deletion syndrome  
   Q. Congenital facial paralysis (Möbius syndrome)  
   R. Craniocarpotarsal syndrome (whistling face syndrome)  
   S. Craniosynostosis-radial aplasia (Baller-Gerold syndrome)
T. Cri-du-chat syndrome (Cry of the cat syndrome)
U. Dubowitz syndrome
V. Down syndrome (trisomy 21, mongolism)
W. Drummond syndrome (idiopathic hypercalcemia, blue diaper syndrome)
X. Ehlers-Danlos syndrome (fibrodysplasia elastica generalisata)
Y. 18q syndrome
Z. Familial blepharophimosis
AA. Fetal alcohol syndrome
BB. Freeman-Sheldon syndrome (whistling face syndrome)
CC. 4Q syndrome
DD. Gansslen syndrome (hematologic-metabolic bone disorder)
EE. Greig syndrome (ocular hypertelorism syndrome)
FF. Hurler syndrome (dysostosis multiplex)
GG. Infantile hypercalcemia
HH. Jacobs syndrome (triple X syndrome)
II. Klinefelter XXY syndrome (gynecomastia-aspermagogenesis syndrome)
JJ. Kohn-Romano syndrome (ptosis, blepharophimosis, epicanthus inversus, and telecanthus)
KK. Komoto syndrome (congenital eyelid tetrad)
LL. Laurence-Moon-Bardet-Biedl syndrome (retinitis pigmentosa-polydactyly-adiposogenital)
MM. Leopard syndrome (multiple lentigines syndrome)
NN. Leroy syndrome (mucopolysaccharide excretion)
OO. Little syndrome (nail patella syndrome)
PP. Michel syndrome
QQ. Mohr-Claussen syndrome (similar to orodigitofacial syndrome)
RR. Noonan syndrome (Turner syndrome in males)
SS. Oculocerebrorenal syndrome (Lowe syndrome)
TT. Oculodentodigital dysplasia (microphthalmos syndrome)
UU. Potter syndrome (renofacial syndrome)
VV. Ring chromosome syndrome
WW. Ring chromosome syndrome
XX. Ring chromosome (microcephaly, hypertelorism, epicanthus)
YY. Ring chromosome in the D group (13-15)
ZZ. Robinow-Silverman-Smith syndrome
AAA. Rubinstein-Taybi syndrome (broad thumbs syndrome)
BBB. Schonenberg syndrome (dwarf-cardiopathy syndrome)
CCC. Smith syndrome (facioskeletogenital dysplasia)
DDD. TAR (thrombocytopenia absent radius) syndrome
EEE. Thalassemia
FFF. Trisomy syndrome (Edward syndrome)
GGG. Turner syndrome (gonadal dysgenesis)
HHH. Waardenburg syndrome (embryonic fixation syndrome)
III. X-linked mental retardation syndrome
JJJ. XXXXX syndrome
HYPOPIGMENTATION (DEPIGMENTATION OF EYELIDS)

1. Drugs, including the following:

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dexamethasone</th>
<th>Methotrexate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adrenal cortex injection</td>
<td>dexamethasone</td>
<td>methotrexate</td>
</tr>
<tr>
<td>Alcohol</td>
<td>fludrocortisone</td>
<td>methylprednisolone</td>
</tr>
<tr>
<td>Aldosterone</td>
<td>fluorometholone</td>
<td>methylthiouracil</td>
</tr>
<tr>
<td>Beclomethasone</td>
<td>fluprednisolone</td>
<td>neostigmine</td>
</tr>
<tr>
<td>Amodiaquine</td>
<td>gentamicin(?)</td>
<td>paramethasone</td>
</tr>
<tr>
<td>Arsenic</td>
<td>hydrocortisone</td>
<td>physostigmine</td>
</tr>
<tr>
<td>Betamethasone</td>
<td>hydroquinone</td>
<td>prednisolone</td>
</tr>
<tr>
<td>Carbimazole</td>
<td>hydroxychloroquine</td>
<td>prednisone</td>
</tr>
<tr>
<td>Chloramphenicol</td>
<td>isoflurophate</td>
<td>propylthiouracil</td>
</tr>
<tr>
<td>Chloroquine</td>
<td>medrysone</td>
<td>thiopeta</td>
</tr>
<tr>
<td>Corticosteroids</td>
<td>meprednisone</td>
<td>triamcinolone</td>
</tr>
<tr>
<td>Cortisone</td>
<td>mercaptoethamine</td>
<td></td>
</tr>
<tr>
<td>Desoxycorticosterone</td>
<td>methimazole</td>
<td></td>
</tr>
</tbody>
</table>

2. Genetic factors

A. Albinism
B. Chediak-Higashi syndrome (anomalous leukocytic inclusions with constitutional stigmata)
C. Cross-McKusick-Breen syndrome
D. Fanconi syndrome (amino diabetes)
E. Hermansky-Pudlak syndrome (oculocutaneous albinism and hemorrhagic diathesis)
F. Histidinemia
G. Homocystinuria
H. Incontinentia pigmenti achromians (hypomelanosis of Ito syndrome)
I. Menkes syndrome (kinky hair syndrome)
J. Nevus depigmentosus
K. Phenylketonuria (Folling syndrome)
L. Tuberculosis (Bourneville syndrome)
M. Vitiligo
N. Vogt-Koyanagi-Harada syndrome (uveitis-vitiligo-alopecia-poliosis syndrome)
O. Waardenburg syndrome (embryonic fixation syndrome)
P. Woolf syndrome (chromosome partial deletion syndrome)
Q. Ziprkowski-Margolis syndrome

3. Following cryosurgery of the eyelid
4. Burns (thermal, ultraviolet, ionizing, radiation)
5. Trauma
6. Kwashiorkor-malnutrition in children
7. Chronic protein deficiency or loss and malabsorption of vitamin B₁₂
8. Endocrine factors
   A. Hypopituitarism (Simmond syndrome)
   B. Addison disease (adrenal cortical insufficiency)
   C. Hyperthyroidism (Graves disease)
9. Inflammation and infection
   A. Discoid lupus erythematosus
   B. Eczematous dermatitis
   C. Leprosy (Hansen disease)
   D. Onchocerciasis syndrome (river blindness)
   E. Pinta
   F. Pityriasis alba
   G. Postinflammatory hypomelanoses
   H. Post-kala-azar
   I. Psoriasis
   J. Sarcoidosis syndrome (Schaumann syndrome)
   K. Syphilis (acquired lues)
   L. Tinea versicolor
   M. Vagabond leukoderma
   N. Vitiligo
   O. Yaws
10. Scleroderma (progressive systemic sclerosis)


**HYPERPIGMENTATION (DISCOLORATION OF LIDS)**

1. Deposits of the eyelids as caused by drugs, including:
2. Hyperpigmentation as caused by drugs, including the following:
   acotophenazine   bismuth oxychloride   carphenazine
   acid bismuth sodium tartrate   bismuth sodium tartrate   chlorpromazine
   actinomycin C   thioglycollate   chlortetracycline
   Alcian blue   bismuth sodium   chrysarobin
   aluminum nicotinate   triglycollamate   colloidal silver
   aminopterin   bismuth sub salicylate   cyclophosphamide
   amiodarone   chloroquine   cytarabine
   amodiaquine   bismuth subcarbonate   dactinomycin
   amphotericin B   bleomycin   demeclocycline
   antipyrine   busulfan   diethazine
   aurothioglucose   butaperazine   doxycycline
   aurothioglycanide   calcitriol   dromostanolone
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<td>ergocalciferol</td>
<td>methylene blue</td>
<td>rifampin</td>
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<td>ethopropazine</td>
<td>mild silver protein</td>
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<td>tetracycline</td>
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<td>perazine</td>
<td>thiethylperazine</td>
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<td>pericyazine</td>
<td>thimerosal</td>
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<td>gold Au 198</td>
<td>perphenazine</td>
<td>thimerosal thiopropazate</td>
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<tr>
<td>gold sodium thiomalate</td>
<td>phenylmercuric acetate</td>
<td>thiouanine</td>
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<tr>
<td>gold sodium thiosulfate</td>
<td>phenylmercuric nitrate</td>
<td>thioperozone</td>
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<tr>
<td>hydroxychloroquine</td>
<td>pipercetazine</td>
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<td>iron dextran</td>
<td>pipobroman</td>
<td>trifluoperazine</td>
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<td>polysaccharide-iron complex</td>
<td>trimeprazine</td>
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<td>practolol</td>
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<td>procarbazine</td>
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<td>prochlorperazine</td>
<td>vitamin D₂</td>
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<td>mesoridazine</td>
<td>promazine</td>
<td>vitamin D₃</td>
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<td></td>
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<tr>
<td>methdilazine</td>
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</table>

3. Periorbital hyperpigmentation-dark circles around the eye
   *A. Allergic rhinitis
   *B. Familial (autosomal dominant)
   C. Medium and dark complexioned white persons

4. Brown hyperpigmentation
   A. Genetic factors
      (1) Acanthosis nigricans
      (2) Albright syndrome (fibrous dysplasia)
      (3) Cafe-au-lait and freckle-like macules in neurofibromatosis
      (4) Dyskeratosis congenita
      (5) Fanconi syndrome (amino diabetes)
      (6) Freckles
      (7) Lentigines
      (8) Melanocytic nevus
      (9) Neurocutaneous melanosis
      (10) Seborrheic keratosis
      (11) Xeroderma pigmentosum
   B. Metabolic factors
      (1) Gaucher syndrome (cerebrosides lipidosis)
(2) Hemochromatosis
(3) Niemann-Pick disease (essential lipid histiocytosis)
(4) Porphyria (cutanea tarda)
(5) Wilson disease (hepatolenticular degeneration)

C. Endocrine factors
   (1) adrenocorticotropic hormone (ACTH) therapy
   (2) Addison disease (adrenal cortical insufficiency)
   (3) Estrogen therapy
   (4) Melanoma
   (5) Pituitary tumors
   (6) Pregnancy

D. Nutritional factors
   (1) Kwashiorkor (hypoproteinemia syndrome)
   (2) Pellagra (avitaminosis B₂)
   (3) Sprue
   (4) Vitamin B₁₂ deficiency (Addison pernicious anemia)

E. Chemical and pharmacologic agents
   (1) Arsenic
   (2) Berlock dermatosis
   (3) Bleomycin
   (4) Busulfan
   (5) Nitrogen mustard, topical
   (6) Photochemical agents

F. Physical agents
   (1) Ionizing radiation
   (2) Thermal radiation
   *(3) Trauma
   (4) Ultraviolet light

G. Inflammation and infection
   (1) Atopic dermatitis
   (2) Lichen planus
   (3) Lichen simplex chronicus
   (4) Lupus erythematosus discoid (Kaposi-Libman-Sacks syndrome)
   (5) Psoriasis
   (6) Tinea versicolor

H. Neoplasms
   (1) Acanthosis nigricans
   (2) Malignant melanoma
   (3) Mastocytosis

I. Miscellaneous factors
   (1) Autosomal recessive ectodermal dysplasia
   (2) Catatonic schizophrenia
   (3) Chronic hepatic insufficiency
   (4) Cronkhite-Canada syndrome
   (5) Encephalitis
   (6) Erythema dyschromicum perstans
(7) Liver spots
(8) Systemic scleroderma (progressive systemic sclerosis)
(9) Whipple syndrome (intestinal lipodystrophy)

5. Blue, gray or slate hyperpigmentation
   A. Genetic factors
      (1) Blue melanocytic nevus
      (2) Dermal melanocytosis (Mongolian spot)
      (3) Franceschetti-Jadassohn syndrome (reticular pigmented dermatosis)
      (4) Incontinentia pigmenti (Bloch-Sulzberger syndrome)
      (5) Oculodermal melanocytosis
   B. Metabolic factors
      (1) Amyloidosis, cutaneous macular (Lubarsch-Pick syndrome)
      (2) Hemochromatosis
   C. Nutritional factors
      (1) Chronic nutritional insufficiency
   D. Inflammation and infection
      (1) Erythema dyschromicum perstans
      (2) Pinta
      (3) Riehl melanosis
   E. Chemical and pharmacologic agents
      (1) Chlorpromazine
      (2) Gold
      (3) Phenothiazine
      (4) Sulfonamides
      (5) Tetracycline
   F. Neoplasms
      (1) Slate-gray dermal pigmentation with metastatic melanoma and melanogenemia
   G. Other
      (1) Blue dye
      (2) Cyanosis


**TUMORS OF EYELIDS**

1. Molluscum contagiosum-small, greasy-appearing elevation that is usually umbilicated or any other granuloma
2. Neoplasm
   A. Basal cell epithelioma-common; may be a red, circumscribed, lobulated growth involving the lid margin or may have an umbilicated center (rodent ulcer)
   B. Squamous cell or Zeis cell epithelioma-hard pearly appearing lesion, usually without increased vascularity
   C. Meibomian-gland carcinoma-resembles a chalazion
   D. Metastatic tumors of the lid-respiratory tract, breast, skin (melanoma), gastrointestinal tract, or kidney
   E. Keratoacanthoma-benign, hemispherical, elevated tumor with a central keratin-filled crater; develops within several months
   F. Hemangioma - rubor of vascular tumor, usually having a smooth surface with tufts of vessels near the surface
   G. Benign mixed tumor of the lacrimal (palpebral) gland
   H. Trichilemmoma
   I. Lymphangioma
   J. Juvenile xanthogranuloma

3. Metaplasia or hyperplasia
   A. Trichoepithelioma
   B. Syringoma
   *C. Sebaceous adenoma
   *D. Papilloma-smooth, rounded, or pedunculated elevation
   *E. Nevus-usually pigmented, raised, and smooth surfaced; however, may be papillomatous or contain hair
   F. Benign calcifying epithelioma
   G. Inverted follicular keratosis
   H. Blue nevus-blue-black and velvet-like in appearance
   I. Freckles
   J. Lentigo simplex
   K. Solar lentigo
   L. Melasma

**Extracted Table Tumors of eyelids**

4. Cyst
   *A. Sebaceous
   B. Sudoriferous
   C. Traumatic
   D. Congenital inclusion
5. Lipoid proteinosis-wax-like, pearly nodules
6. Pseudotumor of lid-encysted contact lens
7. Amyloidosis (Lubarsch-Pick syndrome)

Cook BE, Bartley GB. Epidemiologic characteristics and clinical course of patients with malignant eyelid tumors in an incidence cohort in Olmsted County, Minnesota. *Ophthalmology* 1999; 106:746-750.


**XANTHELASMA (SMOOTH YELLOW DEPOSITS IN THE EYELID, ESPECIALLY THE SUPERIOR NASAL AND INFERIOR NASAL AREAS)**

1. Xanthelasma with hyperlipemia (primary or secondary)  
   *A. Type II-familial hyper-B-lipoproteinemia (familial hypercholesterolemia)*  
   *B. Type III-familial hyper-B- and hyper-pre-B-lipoproteinemia (familial hyperlipemia with hypercholesterolemia)*  
   C. Other types are infrequent, including type I, familial fat-induced hyperlipoproteinemia (hyperchylomicronemia); type IV, familial hyper-pre-B-lipoproteinemia (carbohydrate-induced hyperlipemia); type V, familial hyperchylomicronemia with hyper-pre-B-lipoproteinemia (mixed hyperlipemia), lichen sclerosis et atrophicus.

2. Xanthelasma without hyperlipemia  
   A. Generalized  
   B. Histiocytosis X (eosinophilic granuloma, Hand-Schüller-Christian disease, and Letterer-Siwe disease)  
   C. Local (no systemic disease)  
   D. Reticulohistiocytoma cutis  
   E. Xanthoma disseminatum


**CHRONIC BLEPHARITIS (INFLAMMATION OF LIDS)**

*1. Seborrheic-lid margin covered with small, white or gray scales  
   A. Associated with seborrheic dermatitis of the scalp  
   B. Aggravated by chemical fumes, smoke, and smog  
   C. May be associated with uncorrected refractive errors (especially hyperopia)  
   D. May be due to *Pityrosporon ovale*  
   E. *Aspergillus fumigatus* may be its cause  
   F. Associated with systemic diseases  
      (1) Acne rosacea  
      (2) *Acinetobacter Iwoffi*
(3) Acrodermatitis chronica atrophicans
(4) Aspergillosis
(5) Candidiasis
(6) Cretinism (hypothyroidism)
(7) Demodicosis
(8) Dermatophytosis
(9) Diphtheria
(10) Erysipelas
(11) Herpes simplex
(12) Hypocalcemia
(13) Hypoparathyroidism
(14) Listerellosis
(15) Malaria
(16) Moraxella lacunata
(17) Pellagra (avitaminosis B2)
*(18) Seborrheic dermatitis
(19) Sporotrichosis
(20) Staphylococcus
(21) Streptococcus
(22) Syphilis (acquired lues)
(23) Scleroderma (systemic scleroderma)
(24) Tuberculosis
(25) Vaccinia
(26) Xerodema pigmentosum

G. Associated with syndromes
(1) Danbolt-Closs syndrome (acrodermatitis enteropathica)
(2) Down syndrome (mongolism)
(3) Goldscheider syndrome (epidermolysis bullosa)
(4) Hand-Schüller-Christian syndrome (lipoid granuloma syndrome)
(5) Lyell syndrome (toxic epidermal necrolysis)
(6) Myotonic dystrophy syndrome (Curschmann-Steinert syndrome)
(7) Parkinson syndrome (paralysis agitans)
(8) Sézary syndrome (mycosis fungoides syndrome)
(9) Siemens syndrome (hereditary ectodermal dysplasia syndrome)
(10) Syndrome of Beal (acute follicular conjunctivitis)
(11) Wernicke syndrome
(12) Thiamine deficiency
(13) Wiskott-Aldrich syndrome
(14) Zinsser-Engman-Cole syndrome (dyskeratosis congenita with pigmentation)

H. Drugs, including the following:
acyclovir  meperidine  thimerosal
benzalkonium  mercuric oxide  trifluridine
F3T idoxuridine  nitromersol  vidarabine
isosorbide  phenylmercuric acetate
mannitol  phenylmercuric nitrate
2. Ulcerative-suppurative inflammation of the follicles of the lashes and the associated glands of Zeis and Moll
   A. *Staphylococcus aureus* or *S. albus* may be responsible
   B. Due to mixed infection of a staphylococcus and *P. ovale*
   C. Associated with vaccinia
   D. Due to *Blastomyces dermatitidis*
   E. Herpes simplex-vesicles at lash line, then ulceration

*3. Angular-inflammation of the angles of the lids, usually associated with an angular conjunctivitis
   A. *Candida albicans*
   B. *Moraxella lacunata*
   C. Stannus cerebellar syndrome (riboflavin deficiency)
   D. *Staphylococcus aureus*

4. Exfoliative dermatitis owing to drugs, including the following:
   acetohexamide bismuth sodium cefuroxime
   acetophenazine triglycollamate cephalixin
   acid bismuth sodium bismuth subcarbonate cephaloglycin
   tartrate bismuth subsalicylate cephaloridine
   adiphenine bupivacaine cephalothin
   allobarbital busulfan cepaprin
   allopurinol butabarbital cephradine
   ambutonium butalbital chlorambucil
   aminosalicylate(?) butallylonal chloroprocaine
   aminosalicylic acid butaperazine chloroquine
   amithiozone butethal chlorpromazine
   amobarbital captopril chlorpropamide
   amodiaquine carbamazepine chlorprothixene
   amoxicillin carbenicillin cimetidine
   ampicillin carbimazole ciprofloxacin
   anisindione carisoprodol clidinium
   anisotroline carphenazine clindamycin
   antipyrine cefaclor cloxacillin
   aprobarbital cefadroxil codeine
   atropine methylnitrate cefamandole cyclobarbital
   auranofin cefazolin cyclopentobarbital
   aurothioglucose cefonicid cyclophosphamide
   aurothioglycanide cefoperazone dapsone
   barbital ceforanide dicloxacillin
   bismuth carbonate cefotaxime dicyclomine
   bismuth oxychloride cefotetan diethazine
   bismuth salicylate cefoxitin diltiazem
   bismuth sodium tartrate cefsulodin diphenamid
   bismuth sodium ceftazidime diphenadione
   thioglycollate ceftizoxime diphenylhydantoin
cetrixone droperidol
enalapril
erthyrityl tetranitrate
erythromycin
ethionamide
ethopropazine
ethosuximide
ethotoin
etidocaine
fenoprofen
flecainide
fluphenazine
furosemide
glutethimide
glyburide
glycopyrrolate
gold Au 198
gold sodium hyposulfite
gold sodium thiomalate
griseofulvin
haloperidol
heptabarbital
hetacillin
hexethal
hexobarbital
hexocyclium
hydroxychloroquine
indomethacin
iodide and iodine solutions and compounds
isoniazid
isopropamide
isosorbide dinitrate
ketoprofen
lidocaine
lincomycin
mannitol hexanitrate
mechlorethamine
melphalan
mepenolate
mephenytoin
mephobarbital
mepivacaine
meprobamate
mesoridazine
methantheline
metharbital
methdilazine
methicillin
methimazole
methituril
methixene
methohexital
methotrinemepazine
methoxime
methylatropine nitrate
methylphenidate
methylthiouracil
methyprylon
moxalactam
nafcillin
naltrexone
naproxiv(?)
nitroglycerin
oxacillin
oxyphenbutazone
oxyphencyclimine
oxyphenonium
paramethadione
pentaerythritol tetranitrate
pentobarbital
perazine
pericyazine
perphenazine
phenindione
phenobarbital
phensuximide
phenylbutazone
phenytoin
pimozide
piperazinate
piperacetazine
piperidolate
piroxicam
polidine
practolol
prilocaine
primidone
probarbital
procaaine
procarbazine
prochlorperazine
promazine
promethazine
propantheline
propiomazine
propoxycaine
proproxyphene
propranolol
propylthiouracil
quinacrine
quinidine
radioactive iodides
rifampin
secobarbital
streptomycin
sulfacetamide
sulfachlorpyridazine
sulfacytine
sulfadiazine
sulfadimethoxine
sulfamerazine
sulfameter
sulfamethazine
sulfamethizole
sulfamethoxazole
sulfamethoxypyridazine
sulfanilamide
sulfaphenazole
sulfapyridine
sulfasalazine
sulfathiazole
sulfisoxazole
sulindac
talbutal
thiabendazole
thiamylal
thiethylperazine
thiogalactone
thiopental
thiopropazate
thiopropazine
thioridazine
thiosthixene
tolazamide
tolbutamide
trichloroethylene
tridihexethyl
triethylenemelamine  trimeprazine  vancomycin
trifluoperazine  trimethadione  vinbarbital
trifluperidol  trolnitate  vitamin A
triflupromazine  uracil mustard

5. Other types
   A. Due to mites (*Demodex folliculorum*)
   B. Due to pubic lice (*Phthirus pubis*)

6. Erythema due to drugs, including the following:
acebutolol  cefamandole  cyclosporin
acetaminophen  cefazolin  cyproheptadine
acetanilid  cefonicid  cytarabine
acetazolamide  cefoperazone  dacarbazine
cyclizine  ceforanide  dactinomycin
adrenal cortex injection  cefotaxrime
albuterol  cefotetan  danazol
aldoesterone  cefoxitin  dapiprazole
allopurinol  cefsulodin  hydrochloride
alprazolam  ceftizoxime  daunorubicin
amitriptyline  ceftizoxine  demeclocycline
amoxicapine  ceftiostraxone  desipramine
amantadine  cefuroxime  desoxycorticosterone
atenolol  cephalalexin  dexamethasone
auranofin  cephaloglycin  dexamethasone
aurorothio glucose  cephaloridine  dexamethasone
aurorothioglycanide  cephalothin  dexamethasone
azatadine  cephaspin  dexamethasone
BCG vaccine  cephradine  dexamethasone
beclomethasone  chlorambucil  dexamethasone
benzalkonium  chloridiazepoxide  dexamethasone
benzathine penicilllin G  chlorothiazolecyhloro- 
benzphetamine  nitratsourea (CCNU)  DIC
tetrahydroamino
betamethasone  chlorpheniramine  dichlorphenamidene  
betaxolol  chlorphenermine  diethylcarbamazine
Bis-chloroethyl-nitroso-  chlorphenermine  diethylpropion
urea (BCNU)  chlorpromazine  ditiazem
bleomycin  clofibrate  dimethindene
bromide  ciprofloxacin  dimethyl sulfoxide
bromphenraramine  cisplatin  diphenhydramine
busulphan  clemastine  diphenylypraline
cactinomycin  clofibrate  diphtheria and tetanus

doxoids (adsorbed)
captopril  clomipipramine  diphtheria and tetanus
carboxinamine  clonazepam  toxoids and pertussis
carmustine  clorazepate  vaccine (adsorbed)
cefadoril  cortisone  diphtheria toxoid
cefadroxil  cyclophosphamide (adsorbed)
disopyramide
disulfiram
DMSO
doxepin
doxorubicin
doxycycline
doxylamine
DPT vaccine
enalapril
ergonovine
ergotamine
ethionamide
ethoxzolamide
etractrate
fenfluramine
fenoprofen
flecainide
fleroxidine
fluorometholone
fluorouracil
flurazepam
flurbiprofen
framycetin
gold Au 198
gold sodium thiosulfate
halazepam
hexachlorophene
hydralazine
hydralazine V
hydralazine
hydrocortisone
hydroxyurea
ibuprofen
imipramine
influenza virus vaccine
insulin
iodipamide meglumine
iothalamate meglumine and sodium
iothalamic acid
iron dextran
isotretinoin
ketoprofen
labelalol
levallorphan
levobunolol
lomustine
lorazepam
maprotiline
measles and rubella virus vaccine (live)
measles virus vaccine (live)
measles, mumps, and rubella virus vaccine (live)
mechlorethamine
medrysone
mefenamic acid
meprilone and sodium
melphalan
meperidine
meprednisonone
mercuric oxide
methacycline
methazolamide
methicarbamol
methotrexate
methoxsalen
metylergonovine
methylprednisolone
methysergide
metocurine iodide
metoprolol
metronidazole
mexiletine
mianserine
midazolam
minocycline
minoxidil
mitomycin
mitotane
moxalactam
mumps virus vaccine (live)
nadolol
nalorphine
naloxone
naltrexone
naproxen
neomycin
neostigmine
nifedipine
nitrazepam
nitromersol
nortriptyline
oxazepam
oxprenolol
oxytetracycline
paramethasone
pentazocine
phenacetin
phenindmetrazine
pheniramine
phentermine
phenylphrine
phenylmercuric acetate
phenylmercuric nitrate
pindolol
piroxicam
poliovirus vaccine
potassium penicillin G
potassium penicillin V
potassium phenethicillin
practolol
prazepam
prazosin
prednisolone
prednisone
procaine penicillin G
procarbazine
propranolol
protriptyline
pyrilamine
rabies immune globulin
rabies vaccine
ranitidine
rifampin
rubella and mumps virus vaccine (live)
semustine
smallpox vaccine
spironolactone
streptomycin
strepotozocin
succinylcholine
sulindac
temazepam
tetanus immune globulin
tetanus toxoid
tetracycline
thimerosal
thiotepa
timolol
tocainide
trioxsalen
triprolidine
tubocurarine
uracil mustard
verapamil


**ACUTE BLEPHARITIS (INFLAMMATION OF LIDS WITH RAPID ONSET)**

1. Usual allergy to drugs, including the following:

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<th>Drug</th>
<th>Drug</th>
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<td>antazoline</td>
<td>butabarbital</td>
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<td>acetophenazine</td>
<td>antipyrine</td>
<td>butacaine</td>
</tr>
<tr>
<td>acetyldigitoxin</td>
<td>aprobarbital</td>
<td>butalbital</td>
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<tr>
<td>actinomycin C</td>
<td>aspirin</td>
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<td>acyclovir</td>
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and sodium
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isoflurophate
isoniazid
isopropamide
kanamycin
lanatoside C
levallorphan
levobunolol
levodopa
lidocaine
lincosycin
lomustine
lorazepam
measles and rubella virus vaccine (live)
measles virus vaccine (live)
measles, mumps, and rubella virus vaccine (live)
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medrysone
melphalan
mepenzolate
meperidine
mephenytoin
mepobarbital
mepivacaine
meprobamate
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methotrimethazine
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methylprednisolone
methylthioracil
methyprylon
methysgergide
metolazone
metrizamide
midazolam
mild silver protein
mitomycin
morphine
moxalactam
mumps virus vaccine (live)
nafcillin
nalorphine
naloxone
naltrexone
naphazoline
naproxen
neomycin
neostigmine
niacinamide
nicotinic acid
nicotinyl alcohol
nitrazepam
nitrofurantoin
nitromersol
nystatin
opium
oral contraceptives
ouabain
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piperazine
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pipobroman
poldine
poly thiazide
polymyxin B
potassium penicillin G
potassium penicillin V
potassium phenethicillin
potassium phenoxymethyl
prazepam
prednisolone
prilocaine
primidone
probarbital
procaine
procaine penicillin G
prochlorperazine
promazine
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propoxycaine  sulfamethoxypyridazine  thiophene
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quinethazone  sulfaphenazole  tolbutamide
quinidine  sulfapyridine  trazodone
quinine  sulfasalazine  triazolam
rabies immune globulin  sulfathiazole  trichlormethiazide
rabies vaccine  sulfisoxazole  tridihexeryl
radioactive iodides  talbutal  triethylenemelamine
scopolamine  temazepam  trifluoperazine
secobarbital  testolactone  trifluorothymidine
semustine  testosterone  trifluoperidol
silver nitrate  tetanus immune globulin  triflupromazine
silver protein  tetanus toxoid  trifluridine
sodium salicylate  tetracaine  trimeprazine
streptomycin  tetracycline  trimethadione
streptozocin  tetrahydrozoline  tropic amide
succinylcholine  thiabendazole  uracil mustard
sulfacetamide  thiamylal  vancomycin
sulfachlorpyridazine  thioethylperazine  vidarabine
sulfadiazine  thimerosal  vinbarbital
sulfadimethoxine  thiopental  warfarin

*2. Infections, such as bacterial, fungal, and viral


THICKENED EYELIDS

1. Trachoma
2. Multiple chalazia
3. Chronic conjunctivitis
*4. Blepharitis-lid margins thickened
5. Tarsitis-rare, such as in syphilis or tuberculosis
6. Trisomy (E syndrome)
7. Congenital hypothyroidism
8. Pheochromocytoma, medullary thyroid carcinoma, and neurofibromatosis
BLEPHAROPHIMOSIS (SHORT PALPEBRAL FISSURE)

1. Blepharochalasis
2. Blepharofacieskeletal syndrome
3. Blepharophimosis-amenorrhea syndrome (blepharophimosis, ptosis, epicanthus inversus syndrome)
4. Carpenter syndrome (acrocephalopolysyndactyly II)
5. Clefting syndrome with anterior chamber and lid anomalies
6. Craniofacioskeletal syndrome (Freeman-Sheldon syndrome; whistling face syndrome)
7. Down syndrome (trisomy 21, mongolism)
8. Dubowitz syndrome
9. 18P syndrome
10. Kaufman oculocerebrofacial syndrome
11. Klein-Waardenburg syndrome
12. Komoto syndrome (congenital eyelid tetrad)
13. Marden-Walker syndrome
14. Meyer-Schwickerath and Weyers syndrome
15. Michel syndrome
16. Microphthalmos
17. Mohr syndrome (orofaciocutaneous syndrome II)
18. Mohr syndrome
19. Oculopalatoskeletal syndrome
20. Ohdo blepharophimosis syndrome
21. Pena-Shokeir type II syndrome
22. Progeria (Hutchinson-Gilford syndrome)
23. Rieger syndrome (dysgenesis mesostromalis)
24. Ring chromosome in the D group (13-15) (ring D syndrome)
25. Schonberg syndrome (dwarf cardiopathy syndrome)
26. Schwartz-Jampel syndrome (osteochondromuscular dystrophy)
27. Simosa syndrome
28. Syndrome of blepharophimosis with myopathy
29. Traumatic
30. Trisomy (E syndrome) (Edward syndrome)
31. Waardenburg syndrome (embryonic fixation syndrome)
32. Young-Simpson syndrome
33. X-linked mental retardation syndrome
34. 3p- syndrome
35. 10q- syndrome


**EURYBLEPHARON**

Euryblepharon is defined as a horizontally elongated palpebral aperture (normal, 30 mm) and may be associated with ectropion and present in other family members.

1. Excessive tension of skin
2. Defective separation of the lids
3. Excessive pull of the platysma
4. Localized displacement of the lateral canthi
5. Hypoplasia of tarsus


**LID COLOBOMA**

1. Amniogenic band syndrome (amniotic bands-Streeter anomaly)
2. Epidermal nevus syndrome
3. Facial clefting syndrome (Tessier syndromes)
4. Fraser syndrome
5. Frontoanalis dysplasia syndrome
6. Goldenhar syndrome (oculoauriculovertebral dysplasia)
7. Miller syndrome
8. Nager syndrome
9. Nevus sebaceous of Jadassohn (linear sebaceous nevus syndrome)
10. Palpebral coloboma-lipoma syndrome
11. Traumatic
12. Treacher Collins-Franceschetti syndrome (mandibulofacial dysostosis)


**NECROSIS OF EYELIDS**

1. Drugs, including the following:
   - acenocoumarol
   - amphotericin B
   - anisindione
   - dicumarol
   - diphenadione
   - ethyl biscoumacetate
   - nafcillin
   - phenindione
   - phenprocoumon
   - tobramycin
   - warfarin

2. Mechanical, electrical, or thermal trauma

3. Periorbital cellulitis-periorbital necrotizing cellulitis

4. Secondary to infection


**POLIOSIS (WHITENING OF HAIR, EYEBROWS, AND EYELASHES)**

1. Albino
2. Alopecia areata
3. Aging
4. Drugs, including the following:
   - amodiaquine
   - betamethasone
   - chloroquine
   - cortisone
   - cyclosporin A
   - dexamethasone
   - epinephrine
   - fluorometholone
   - hydrocortisone
   - hydroxychloroquine
   - medrysone
   - prednisolone
   - thiotepa

5. Leprosy (Hansen disease)
6. Radiation therapy
7. Rubinstein-Taybi syndrome
8. Severe dermatitis
9. Stress
10. Unknown etiology
11. Vitiligo
13. Waardenburg syndrome (embryonic fixation syndrome)
14. Werner syndrome (progeria of adults)


TRICHOMEGALY (LONG LASHES)

1. Associated with cataract and hereditary spherocytosis
2. Congenital with pigmentary retinal degeneration, dwarfism, and mental retardation
3. Cyclosporine induced
4. De Lange syndrome (congenital muscular hypertrophy-cerebral syndrome)
5. Ectodermal dysplasia (Curtius syndrome)
6. Human immunodeficiency virus (HIV)
7. Hypertrichosis (hirsutism)
8. Isolated adrenal malfunction and ovarian atrophy
9. Noonan syndrome (male Turner syndrome)
10. Normal
11. Oliver-McFarlane syndrome
12. Rubinstein-Taybi syndrome (broad thumbs syndrome)
13. Schwartz syndrome


MADAROSIS (LOSS OF EYELASHES)

1. Chronic skin diseases, including psoriasis, neurodermatitis, exfoliative dermatitis, ichthyosis, alopecia areata, acne, lichen planus, epidermolysis bullosa, lupus erythematosus, acanthosis nigricans, dermatophytosis, hereditary ectodermal dysplasia syndrome and acrodermatitis
d2. Congenital atrichia
3. Cryptophthalmos
4. Cutaneous T cell lymphoma
5. Demodicosis
6. Drugs, including the following:

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sulfacytine(?)

7. Ehlers-Danlos syndrome, unspecified type
*8. Endocrine disease, including hypothyroidism, hyperthyroidism, pituitary insufficiency, hypoparathyroidism, and pituitary necrosis syndrome (Simmonds-Sheehan syndrome)
9. Following eyelid tattooing
10. Generalized hypotrichosis
11. HIV (human immunodeficiency virus)
12. Hypocalcemia
13. Hypothermal injury
*14. Idiopathic
15. Inflammation and infection of the lids, including seborrheic blepharitis, squamous blepharitis, herpes zoster, sebaceous gland carcinoma, vaccinia, mycotic infection, furuncles, and erysipelas
16. Intoxication with arsenic, bismuth, thallium, gold, quinine and vitamin A
17. Isolated madarosis
18. Keratosis decalvans
19. Keratosis follicularis
20. Keratosis spinulosa
21. Lid colobomas
22. Leprosy
23. Lipoid proteinosis (Urbach-Wiethe syndrome)
24. Polymorphous light eruption
25. Pseudoprogeria syndrome
26. Radiation
27. Severe debilitating systemic diseases, including tuberculosis, syphilis, sickle cell anemia, cholera, and Hansen disease (leprosy)
28. Trauma

**Extracted Table Madarosis (loss of eyelashes)**


**DISTICHIASIS (ACCESSORY ROW OF LASHES GROWING FROM OPENINGS OF MEIBOMIAN GLAND)**

1. Acquired
   A. Chemical
   B. Immunologic
   C. Physical
   *2. Congenital
      A. Anodontia-hypotrichosis syndrome
      B. Distichiasis, lymphedema syndrome
      C. Ectropion and distichiasis
      D. Idiopathic eyelid edema
E. Pierre Robin syndrome
F. Tristichiasis

3. Hereditary-autosomal dominant


**COARSE EYEBROWS**

1. Congenital hypothyroidism (cretinism)
2. CPD syndrome (chorioretinopathy and pituitary dysfunction)
3. Hunter syndrome [mucopolysaccharidosis (MPS II)]
4. Hurler syndrome (MPS I)
*5. Normal variation
6. Rubinstein-Taybi syndrome (broad thumbs syndrome)
7. Sanfilippo syndrome (MPS III)


**SYNOPHRYS (CONFLUENT EYEBROWS EXTENDING TO MIDLINE)**

1. Basal cell nevus syndrome (Gorlin syndrome)
2. Cornelia De Lange syndrome (congenital muscular hypertrophy-cerebral syndrome)
3. Deletion 3p syndrome
4. Duplication 3q syndrome
5. Frontometaphyseal dysplasia
6. Hirschhorn-Cooper syndrome (chromosome partial deletion syndrome)
7. Labard syndrome
*8. Normal variation
9. Partial trisomy chromosome 15
10. Smith-Lemli-Opitz syndrome (cerebrohepatorenal syndrome)
11. Thirteen trisomy syndrome (Patau syndrome)
12. Waardenburg syndrome (interoculoiridodermatoauditive dysplasia)

HERTOGH SIGN (LACK OF OUTER THIRD OF EYEBROWS)

1. Autonomic nervous system dysfunction
2. Diphtheria
3. Endocrinopathies
4. Hypogonadism
5. Hypothyroidism
6. Neurodermatitis
7. Scleroderma (systemic scleroderma)


LID MYOKYMIA (SPONTANEOUS FASCICULAR EYELID TREMOR WITHOUT MUSCULAR ATROPHY OR WEAKNESS)

*1. Not associated with organic disease
   A. Fatigue
   B. Lack of sleep
   C. Bright light dazzle
   D. Irritative corneal or conjunctival lesions
   E. Debility or anemia
   F. Excessive alcohol or smoking
   G. Overwork
2. Followed by spastic paretic facial contracture—in dorsal pons in adult and children
3. Multiple sclerosis (disseminated sclerosis)
4. Trigeminal neuralgia
5. Myasthenia gravis (pseudoparalytic syndrome)
6. Familial occurrence
7. Autosomal dominant familial dystonia


PRESEPTAL CELLULITIS OF EYELID

1. Eczema
2. Hordeolum
3. Neonatal conjunctivitis
4. Otitis media
5. Sinusitis
6. *S. aureus*
7. Toxic shock syndrome
8. Trauma
9. Upper respiratory tract
10. Varicella


**TELECANTHUS (DISPROPORTIONATE INCREASE IN DISTANCE BETWEEN MEDIAL CANTID; MEASUREMENTS IN INFANTS ARE 18 TO 22 MM)**

*1. Primary*

2. Secondary—may occur secondarily to an increased distance between the bony orbits (see hypertelorism)
   - A. Aarskog syndrome
   - B. Blepharonasofacial syndrome
   - C. Blepharophimosis syndrome
   - D. Camptomelic dysplasia
   - E. Carpenter syndrome
   - F. Cerebrofacioarticular syndrome of van Maldegem
   - G. Coffin-Lowry syndrome
   - H. de Lange syndrome
   - I. Deletion 5g syndrome
   - J. Dubowitz syndrome
   - K. Facial-renal acromesomelic syndrome
   - L. Faciooculoacousticorenal syndrome
   - M. Fetal alcohol syndrome
   - N. Fetal hydantoin syndrome
   - O. Frontonasal dysplasia
   - P. Lambotte syndrome
   - Q. KBG syndrome (initials of family studied)
   - R. Michel syndrome
   - S. Nasopalpebral lipoma-coloboma syndrome
   - T. Oculodentodigital syndrome
   - U. Orofaciodigital (OFD) type I and type II (Mohr syndrome)
   - V. Prader-Willi syndrome
   - W. Rieger syndrome
   - X. Simosa syndrome
   - Y. Tetra-X syndrome
   - Z. Trisomy syndrome
   - AA. Toriello-Carey syndrome
BB. Trauma
CC. Waardenburg syndrome
DD. Williams syndrome
EE. 5p- syndrome (Cri-du-chat)
FF. 10q- syndrome
GG. Fetal hydantoin syndrome


**ANKYLOBLEPHARON (PARTIAL OR COMPLETE FUSION OF UPPER TO LOWER EYELIDS)**

*1. Ablepharon, macrostomia syndrome
2. Ankyloblepharon ectodermal dysplasia, cleft lip and palate
3. Curly hair, ankyloblepharon, nail dysplasia syndrome (CHANDS)
4. Cryptophthalmos (complete fusion of lids)
5. Diphtheritic conjunctivitis
6. Ectodermal syndrome
7. Edward syndrome
8. Fraser syndrome
9. Gastrointestinal anomalies
10. Hay-Wells ectodermal pterygium syndrome
11. Popliteal pterygium syndrome
12. Smallpox
13. Trachoma
14. Trisomy 18
15. Ulcerative blepharitis


**FLARING OF NASAL PART OF EYEBROW**

1. Blepharonasofacial syndrome
2. Partial trisomy 10q syndrome
*3. Waardenburg syndrome
4. Williams syndrome

HIGH ARCHED BROW

1. Kabuki makeup syndrome
2. Shprintzen-Goldberg syndrome


ABSENT BROW HAIR

1. Cryptophthalmos
2. Duplication 14q syndrome
3. Pallister-Killian syndrome
4. Pseudoprogeria syndrome


TRICHIASIS (INWARD TURNING LASHES)

*1. Inflammation/infection
   A. Chronic blepharitis
   B. Herpes simplex or zoster
   C. Trachoma
2. Lid Tumors
   A. Basal cell carcinoma
   B. Capillary hemangioma
   C. Conjunctiva amyloidosis
3. Medications
   A. Epinephrine
   B. Idoxuridine
   C. Phospholine iodide
   D. Pilocarpine
   E. Practolol
   F. Trifluridine
   G. Vidarabine
4. Systemic/Immunologic Disorders
   A. Erythema multiforme
   B. Ocular cicatrical pemphigoid
   C. Stevens-Johnson syndrome
   D. Toxic epidermal necrolysis
   E. Vernal kertoconjunctivitis
5. Trauma
   A. Chemical injury (lye)
   B. Mechanical injury or repair of injury
      (1) Lower lid transconjunctival approach for floor fracture repair or blepharoplasty
(2) After enucleation
(3) After ectropion repair
(4) Thermal burns to face/lids

C. Surgery


3
Lacrimal System

CONTENTS

Dacryoadenitis 110
   Acute 110
   Chronic 110
   Diagnostic table 112
   Mikulicz syndrome-Diagnostic table 113
   Painless enlargement of lacrimal gland 111
   Painful enlargement of lacrimal gland 111

Bloody tears 111
   Diagnostic table 114

Excessive tears 115
   Hypersecretion of tears 115
   Inadequacy of lacrimal drainage system 118
   Diagnostic table 120

Drugs found in tears 122

Dry eye 122

Dacryocystitis 125
   Acute 125
   Chronic 125

DACRYOADENITIS (INFLAMMATION OF LACRIMAL GLAND)

1. Acute dacryoadenitis-rare catarrhal inflammation of the lacrimal gland that usually accompanies systemic disease
   A. In children-*mumps, measles, influenza, scarlet fever, erysipelas, typhoid fever
   B. In adults-gonorrhea, endogenous conjunctivitis and uveitis, infectious mononucleosis, typhoid fever, Crohn disease
   C. Secondary to inflammation from lids or conjunctiva, to include *Klebsiella pneumoniae, coliform organisms, *Staphylococcus, *Streptococcus, Aedes aegypti, *Diplococcus pneumoniae, and *Neisseria gonorrhoea

2. Chronic dacryoadenitis-proliferative inflammation of the lacrimal gland, usually because of specific granulomatous disease
   A. Boeck sarcoid (Schaumann syndrome)
   B. Heerfordt disease-chronic bilateral parotitis and uveitis, often associated with paresis of the cranial nerves, usually the seventh nerve, and other general symptoms
      *(1) Sarcoidosis syndrome (Schaumann syndrome)
      *(2) Tuberculosis
   C. Mikulicz syndrome-dacryoadenitis and parotitis manifested by chronic bilateral swelling of the lacrimal and salivary glands
      *(1) Bang disease (brucellosis)
(2) Hodgkin disease  
(3) Leukemia  
(4) Lymphoma  
(5) Lymphosarcoma (Brill-Symmers disease)  
(6) Reticuloendothelial disease  
(7) Mumps  
*(8) Sarcoidosis syndrome (Schaumann syndrome)  
(9) Syphilis  
(10) Tuberculosis  
(11) Waldenstrom macroglobulinemia  
D. Miliary tuberculosis  
E. Pseudotumor  
F. Syphilis (gumma)

3. Painless enlargement of lacrimal gland  
   A. Leukemia  
   B. Mumps

4. Painful enlargement of lacrimal gland  
   A. Autoimmunologically mediated syndrome  
   B. Lymphomatous disease (25%)  
   C. Chronic enlargement arising from sarcoid or orbital pseudotumor (25%)  
   D. Lacrimal gland neoplasm (50%)  
   (1) Benign  
      a. Adenoma  
      b. Mixed tumor  
   (2) Malignant  
      a. Carcinoma unrelated to mixed tumor  
      b. Adenocarcinoma (adenoid cystic carcinoma)  
      c. Mucoepidermoid carcinoma  
      d. Squamous cell carcinoma  
      e. Mixed tumor


BLOODY TEARS

1. Conjunctiva
   A. Application of a drug such as silver nitrate
   B. Cachectic conjunctivitis
   C. Focal dermal hypoplasia syndrome (Goltz syndrome)
   D. Fibroma
   *E. Giant papillary conjunctivitis secondary to contact lens wear or prosthesis wear
   F. Gross disturbance of autonomic nervous system
   G. Hemangioma
   H. Hereditary hemorrhagic telangiectasis
   I. Inflammatory granuloma
   J. Malignant melanoma
   K. Metastatic carcinoid tumor
   *L. Severe conjunctivitis with marked hyperemia
   *M. Subconjunctival hemorrhage following sudden venous congestion of head from stooping, coughing, choking, Valsalva trauma, hemophilia or advanced athrombia
   N. Vicarious menstruation with ectopic tissue

2. Corneal vascular lesion or pannus

3. Lid
   A. Pubic lice and nits on the lashes
   B. Trauma

4. Other
   A. Familial telangiectasis
   B. Hemophilia
   C. Hysteria
   D. Jaundice
   E. Osler-Weber-Render
   F. Pathologic process of lacrimal gland
   G. Severe anemia
   H. Severe epistaxis with regurgitation through the lacrimal passages


EXCESSIVE TEARS

1. Hypersecretion of tears—may be due to basic secretors (mucin, lacrimal, including secretion from glands of Kraus and Wolfring and oil, including secretion from Zeis, Moll, and Meibomian palpebral glands) or reflex secretors (main lacrimal glands and accessory palpebral glands)
   A. Primary (disturbance of lacrimal gland)
   B. Central or psychic
      (1) Central nervous system lues
      (2) Corticomeningeal lesions
      (3) Emotional states
      (4) Hysteria
      (5) Physical pain
      (6) Voluntary lacrimation, such as when acting
   C. Neurogenic
      (1) Ametropia, tropia, phoria, and eyestrain or fatigue
      (2) Caloric, lacrimal, and reflex tearing—bilateral lacrimation when syringing the ear with warm or cold water and during Tensilon testing
      (3) Crocodile or alligator tears—unilateral profuse tearing when eating
         a. Congenital, often associated with ipsilateral paresis of lateral rectus muscle
         b. Acquired with onset in early stage of facial palsy (Bell palsy) or sequela with parasympathetic fibers to the otic ganglion growing back into superficial petrosal nerve
         c. Duane retraction syndrome
      (4) Bell palsy (idiopathic facial paralysis)
      (5) Marin-Amat syndrome (inverted Marcus Gunn Jaw-wink phenomenon)
      (6) Melkersson-Rosenthal syndrome (Melkersson idiopathic fibro edema)
         d. Section of the greater superficial petrosal nerve
   (7) Drugs, including the following:

   | acetylbutorol | atenolol | chlordiazepoxide |
   | acetophenazine | beclomethasone | chlorpromazine |
   | acetylcholine | betamethasone | ciprofloxacin |
   | adrenal cortex injection | betahanehol | clonazepam |
   | alcohol | bishydroxycoumarin | clorazepate |
   | aldosterone | bleomycin | codeine |
   | alseroxylon | butaperazine | dactinomycin |
   | ambenonium | carbachol | dantrolene |
   | antazoline | carphenazine | desoxycorticosterone |
   | Apresoline | chloral hydrate | dexamethasone |
dextran  lithium carbonate  prazepam
diazepam  lorazepam  prochlorperazine
diazoxide  meprednisone  promazine
dicumarol  mesoridazine  promethazine
diethazine  methacholine  propiomazine
diethylcarbamazine  methaqualone  propoxyphene
diltiazem  methdilazine  propranolol
disodium pamidronate  methotrimeprazine  pyridostigmine
doxorubicin  methylprednisolone  pyridostigmine
edrophonium  metoprolol  quinidine
epinephrine  midazolam  pyrilamine
ether  mitomycin  quinidine
ethopropazine  morphine  rauwolfa serpentina
etretinate  nadolol naltrexone  rescinnamine
floxuridine  nalorphine  reserpine
fludrocortisone  naloxone  rifampin
fluorometholone  neostigmine  silydostigmine
fluourouracil  nifedipine  temazepam
fluphenazine  nitrazepam  thiethylperazine
fluprednisolone  opium  thiopropazate
flurazepam  oxazepam  thioproperazine
glycerin  paramethasone  thioridazine
halazepam  pentazocine  triazolam
heparin  perazine  trichloroethylene
hydralazine  pericystine  trifluoperazine
hydrocortisone  perphenazine  triflupromazine
hydroxyamphetamine  phenylephrine  trimeprazine
indomethacin(?)  pindolol  tripelelamine
isotretinoïn  pipercaceline  verapamil
ketarnine  piperezine  vinblastine
labetalol  piroxicam  warfarin
levallorphan  practolol

(8) Exposure to wind, cold, or bright light; photosensitivity and sunburn
(9) Glaucoma
(10) Homer syndrome (see p. 59) (cervical sympathetic paralysis syndrome)
(11) Inflammation or infection of the conjunctiva, uvea, cornea, orbit, lids, sinuses, teeth, or ears
  a. Acute hemorrhagic conjunctivitis
  b. Avitaminosis B (pellagra, niacin deficiency)
  c. Conjunctivochalasis
  d. Elschnig syndrome (I) (meibomian conjunctivitis)
  e. Epidemic keratoconjunctivitis
  f. Feer syndrome (acrodynia)
  g. Hanhart syndrome (recessive keratosis palmoplantaris)
h. Keratodermia palmaris et plantaris
i. Reiter syndrome (polyarthritis enteric)
j. Stannus cerebellar syndrome (riboflavin deficiency)
k. Thelaziasis
(12) Lesions affecting the lids
 a. Acrodermatitis chronic atrophicans
 b. Blepharoptosis
 c. Congenital distichiasis
* d. Ectropion
*e. Entropion
 f. Epiblepharon
 g. Eyelid retraction
* h. Facial paralysis
 i. Lid imbrication syndrome
 j. Papilloma
 k. Punctal apposition
 l. Trachoma
* m. Trichiasis
(13) Morquio-Brailsford syndrome (MPS IV)
(14) Myasthenia gravis-afternoon ectropion (Erb-Goldflam syndrome)
(15) Ophthalmorhinostomatohygrosis syndrome
(16) Parkinson disease-facial akinesia
(17) Reflex, such as vomiting or laughing
(18) Sjögren syndrome (secretinhibitor syndrome)
(19) Stimulation of some cortical areas-thalamus, hypothalamus, cervical sympathetic ganglia, or the lacrimal nucleus
 a. Diencephalic epilepsy syndrome (Penfield syndrome)
 b. Encephalitis
  (1) acute
  (2) hemorrhagica superior
  (3) lethargy
  (4) periaxialis diffusa
 c. Engelmann syndrome (diaphyseal dysplasia)
 d. Giant-cell arteritis (temporal arteritis)
 e. Hypothalamic tumors
 f. Meningitis
 g. Page syndrome (hypertensive diencephalic syndrome)
 h. Pseudobulbar palsy from Parkinson syndrome (shaking palsy)
 i. Sluder syndrome (lower facial neuralgia syndrome)
 j. Tic doulourex (trigeminal neuralgia syndrome)
 k. Various senile dementias
(20) Gradenigo syndrome (temporal syndrome)
(21) Raeder syndrome (paratrigeminal paralysis, cluster headache)
(22) Retroparotid space syndrome (Villaret syndrome)
(23) Rhabdomyosarcoma
(24) Rothmund syndrome (telangiectasia-pigmentation-cataract syndrome)
25. Thermal burns

D. Symptomatic
   1. Bee sting of cornea
   2. Tabes
   3. Thyrotoxicosis (Basedow syndrome)

2. Inadequacy of lacrimal drainage system
   A. Congenital anomalies of lacrimal apparatus
      1. Absence or atresia including ectrodactyly-ectodermal dysplasia-clefting syndrome
      2. Amniotocele
      *3. Fistulas of lacrimal sac and nasolacrimal duct
      4. Lateral displacement of medial canthi with lateral displacement of puncta and lengthening of canaliculi as in Waardenburg syndrome (interoculooridodermatoauditive dysplasia)
      5. Obstruction of nasolacrimal drainage system, including Walker-Clodius syndrome (lobster claw deformity with nasolacrimal obstruction)
      *6. Unformed puncta (punctal atresia)
   B. Complications from diseases such as pemphigus, Stevens-Johnson syndrome (dermatostomatitis), and lupus.
      *C. Dacryocystitis
      *D. Distended canaliculi with obstruction, such as from Actinomyces israelii (Streptothrix foersteri), papilloma, or dacryolith
   E. Because of drugs, including the following:
      acyclovir
      adenine arabinoside
      colloidal silver
      demecarium
      DEP
      echthiophate
      epinephrine
      F3T
      floxuridine
      fluorouracil
      IDU
      indoxuridine
      isoflurophate
      neostigmine
      phystostigmine
      quinacrine
      silver nitrate
      silver protein
      thiotepa
      trifluorothymidine
      trifluorothymidine (including Fuchs-Lyell syndrome) (allergic reaction due to drugs causing nasolacrimal obstruction)
      *F. Eversion of inferior lacrimal punctum, including involutional ectropion (horizontal lid laxity and retractors disinsertion)
   G. Eversion of inferior lacrimal punctum secondary to ichthyosis or scleroderma
   H. Goltz syndrome (focal dermal hypoplasia syndrome)
   I. Inadequacy of physiologic lacrimal pump
   J. Traumatic lesions of lacrimal drainage system
   K. Tumor obstruction, including polyps, papillary hypertrophy, and neurofibromas
      1. Botulinum toxin usage
      2. Dyskeratosis congenita (Zinsser-Engman-Cole syndrome)
      3. Gravity inversion
      4. Irritation from dust and gases
      5. Inflammation or destruction of turbinates
(6) Inhalation cocaine abuse
(7) Leprosy (Hansen disease)
(8) Leukemia
(9) Plasmoma
(10) Rhinosporidiosis
(11) Scleroma
(12) Tuberculosis
(13) Tumors
L. Primary neoplasms
(1) Fibroma
(2) Hemangiopericytoma
(3) Melanoma
(4) Papilloma
(5) Squamous cell carcinoma
M. Secondary involvement by neoplasms
   (1) Basal cell carcinoma
   (2) Lethal midline granuloma
   (3) Leukemia
   (4) Lymphoma
   (5) Maxillary sinus tumors
   (6) Neurofibroma
   (7) Wegener granulomatosis
N. Nasal disease
   (1) Sinusitis
   (2) Hypertrophic rhinitis
   (3) Pseudoepiphora, such as wound fistula following intraocular operation with leak of aqueous


**Extracted Table Excessive Tears**
DRUGS FOUND IN TEARS

Drugs, including the following:
alcian blue            fluorouracil            sodium salicylate
amodiaquine           hydroxychloroquine       trypan blue
aspirin               methotrexate           vitamin A
chloroquine           minocycline
fluorescein            rose bengal


DRY EYE (PAUCITY OR ABSENCE OF TEARS)

1. Xerosis-local tissue changes
   A. Cicatricial degeneration of conjunctiva and mucous tissues
      (1) General-diphtheria
      (2) Upper lid-trachoma
      (3) Lower lid
         a. Avitaminosis A
         b. Chemical irritation (especially due to alkali)
         c. Dermatitis herpetiformis (Duhring-Brocq disease)
         d. Epidermolysis bullosa (Weber-Cockayne syndrome)
         e. Erythema multiforme (Stevens-Johnson syndrome)
         f. Ocular pemphigoid
         g. Plummer-Vinson syndrome (sideropenic dysphagia syndrome)
         h. Radium burns
         i. Reiter syndrome (conjunctivourethrosynovial syndrome)
         j. Sjögren syndrome (secretoinhibitor syndrome)
         k. Uyemura syndrome (fundus albipunctatus with hemeralopia and xerosis)
   B. Exposure keratitis
      (1) Anterior lamella shortage secondary to trauma or facial bum
      *(2) Deficient lid closure as part of facial palsy
      *(3) Ectropion (see p. 81)
      (4) Eyelid retraction-Graves ophthalmopathy (incomplete blink)
      (5) Exophthalmos
      (6) Following botulism
      *(7) Infrequent blinking, such as with progressive supranuclear palsy
      (8) Lack of blinking as during coma
      (9) Levator spasm
      (10) Melkersson-Rosenthal syndrome (Melkersson idiopathic fibroedema)
(11) Methylmalonic aciduria
(12) Ocular proptosis
(13) Rapid evaporation in hot, dry areas
(14) Stiff, immobile, retracted lids, such as those occurring secondary to 
tuberculoid leprosy (Hansen disease)

2. Keratoconjunctivitis sicca—primary tear diminution of main and accessory lacrimal glands

A. Congenital
   (1) Congenital absence of lacrimal gland as in Bonnevie-Ullrich syndrome
   (2) Neurogenic
   (3) Associated with generalized disturbance
      a. Anhidrotic type of ectodermal dysplasia
      b. Familial dysautonomia (Riley-Day syndrome)
      c. Cry-du-chat syndrome (Cry of the cat syndrome)
      d. Cystic fibrosis syndrome (fibrocystic disease of pancreas)

B. Neurogenic hyposecretion
   (1) Central-aplasia of lacrimal nucleus or lesion of seventh nerve between 
nucleus and geniculate ganglion
      a. Pontine lesions
      b. Basal fractures
      c. Otitis media
   (2) Peripheral—greater superficial petrosal nerve, sphenopalatine ganglion, 
or lacrimal branch
      a. Skull fractures
      b. Associated with neoplasms
      c. Neurologic lesion of fifth nerve (neuroparalytic keratitis)
   (3) Herpes zoster of the geniculate ganglion (Ramsey-Hunt syndrome)
   (4) Parasympathetic blocking drugs, such as atropine and scopolamine, 
      may decrease an already barely adequate secretion.
   (5) Botulism
   (6) Deep anesthesia
   (7) Debilitating disease, such as typhus and cholera, and high temperature
   (8) Allergy

C. Systemic disease
   (1) Allgrove syndrome
   (2) Acquired immunodeficiency syndrome (AIDS)
   (3) Amyotrophic lateral sclerosis
   (4) Danbolt-Closs syndrome (acrodermatitis enteropathica)
   (5) Disseminated lupus erythematosus (Kaposi-Libman-Sacks syndrome)
   (6) Drugs, including the following:
      acebutolol      astemizole      benzalkonium
      acetophenazine  atenolol      benzthiazide
      albuterol       atropine      brimonidine
      aluminum nicotinate(?) azatadine      brompheniramine
      amitriptyline   belladonna   busulfan
      antazoline      bendroflumethiazide  butaperazine
<table>
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<tr>
<th>Drug Name</th>
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(7) Felty syndrome (uveitis-rheumatoid arthritis syndrome)
(8) Gougerot-Sjögren syndrome (oligophrenia-ichthyosis-spastic diplegia syndrome)
(9) Heerfordt syndrome (uveoparotitis)
(10) Lubarsch-Pick syndrome (primary amyloidosis)
(11) Mikulicz syndrome-acryoadenitis and parotitis
a. Hodgkin disease
b. Leukemia
c. Lymphoma
d. Lymphosarcoma
e. Mumps
f. Sarcoidosis syndrome (Schaumann syndrome)
g. Syphilis
h. Tuberculosis
i. Waldenstrom macroglobulinemia
(12) Pancreatitis  
(13) Pheochromocytoma, medullary thyroid carcinoma, and multiple mucosal neuromas  
(14) Polyarteritis nodosa (Kussmaul disease)  
(15) Relapsing polychondritis  
(16) Rheumatoid arthritis (adult)  
(17) Scleroderma (progressive systemic sclerosis)  


DACRYOCYSTITIS (INFECTION OF THE LACRIMAL SAC)  

1. Acute dacryocystitis  
   *A. Beta-hemolytic streptococcus  
   B. Corynebacterium diphtheriae  
   C. Dacryolith  
   D. Erysipelothrix insidiosa  
   E. Friedlander bacillus  
   F. Fusobacterium (canaliculitis and dacryocystitis)  
   G. Granulomatous "pseudotumor"  
   *H. Haemophilus aegyptius (Koch-Weeks bacillus)  
   I. Infectious mononucleosis  
   J. Influenza  
   K. Lymphocytic neoplasia  
   L. Neisseria catarrhalis  
   M. Pasteurella multocida  
   *N. Pneumococcus  
   *O. Pseudomonas aeruginosa  
   P. Rhinosporidiosis  
   Q. Rubeola (measles)  
   R. Serratia marcescens - gram-negative coccobacillus  
   *S. Staphylococcus  
   T. Streptococcus  
   U. Tularemia  
   V. Variola  

2. Chronic dacryocystitis
A. Associated with osteopoikilosis
B. Actinomyces israelii
C. Aspergillus
D. Bacillus fusiformis
E. Candida albicans
F. Escherichia coli
G. Lymphoma of the lacrimal sac
H. Mycobacterium fortuitum and Mycobacterium chelonei
I. Nocardia asteroides
J. Francisella tularensis
K. Mycobacterium leprae
L. Proteus vulgaris
M. Sporotrichosis
N. Syphilis (acquired lues)
O. Systemic sarcoidosis
P. Thermal burns
Q. Trachoma
R. Treponema vincentii
S. Tuberculosis (Mycobacterium tuberculosis)
T. Wegener granulomatosis


4

Extraocular Muscles

CONTENTS

Pseudoesotropia  128
Esophoria and esotropia  128
Pseudoexotropia  129
Exophoria and exotropia  130
Pseudohypertropia  130
Hyperphoria and hypertropia  130
Brown superior oblique tendon sheath syndrome  132
Duane syndrome  133
Oculomotor apraxia  133
Monocular limitation of elevation of adducted eye with forced duction test  134
Cyclic, recurrent, repetitive, episodic disorders of extraocular muscles  134
Syndromes and diseases associated with strabismus  135
Horizontal gaze palsy  139
Oscillations of eyes  140
Cogwheel eye movements  141
Pendular nystagmus  142
Horizontal jerk nystagmus  142
Vertical nystagmus  143
Rotary nystagmus  144
See-saw nystagmus  144
Retraction nystagmus  145
Monocular nystagmus  145
Periodic alternating nystagmus  145
Positional nystagmus  146
Optokinetic nystagmus  148
Syndromes and diseases associated with nystagmus  148
Oculogyric crisis  152
Ocular bobbing  153
Paralysis of third nerve  153
  Diagnostic table  154
  Childhood causes of third nerve (oculomotor) palsy  157
Paralysis of fourth nerve  158
  Diagnostic table  159
  Childhood causes of fourth nerve (trochlear) palsy  160
Pseudoabducens palsy  160
Paralysis of sixth nerve  161
  Diagnostic tables  162
  Paralysis of sixth nerve  162
  Paralysis of third, fourth, and sixth cranial nerves  164
  Childhood causes of sixth nerve (abducans) palsy  167
  Childhood causes of third, fourth, and sixth nerve palsy  167
Acute ophthalmoplegia 168
Chronic ophthalmoplegia 171
Bilateral complete ophthalmoplegia 171
External ophthalmoplegia 172
Internuclear ophthalmoplegia 173
        Diagnostic table 174
Painful ophthalmoplegia 173
        Diagnostic table 175
Transient ophthalmoplegia 176
Painful ocular movements 176
Poor convergence 177
Spasm of convergence 178
Divergence paralysis 178
Oculocardiac reflex 179
Retraction of the globe 179
Forced duction test 180
Double elevator palsy 181
Ocular neuromyotonia 181
Extraocular muscle enlargement 181

**PSEUDESOTROPIA (OCULAR APPEARANCE OF ESOTROPIA WHEN NO MANIFEST DEVIATION OF VISUAL AXIS IS PRESENT)**

* 1. Abnormal shape of skull or abnormal thickness of skin surrounding the orbits
2. Enophthalmos
3. Entropion (p. 79)
4. Hypotelorism with narrow interpupillary distance
5. Lateral displacement of the concavity of the upper eyelid margin from the center of the pupil
*6. Negative-angle kappa-pupillary light reflex displaced temporally (see Decentered Pupillary Light Reflex, p. 361)
*7. Prominent epicanthal fold
8. Telecanthus-the orbits are normally placed, but the medial canthi are far apart secondary to lateral displacement of the soft tissues


**ESOPHORIA AND ESOTROPIA (VISUAL AXIS DEVIATED INWARD; MAY BE LATENT OR MANIFEST)**

1. Comitant (nonparalytic)-angle of deviation is constant in all directions of gaze
   A. Accommodative-hyperopic refractive error
   B. Nonaccommodative-refractive error not cause of deviation
(1) Anomalous insertion of horizontally acting muscles
(2) Abnormal check ligaments
(3) Faulty innervational development
(4) Autosomal recessive trait
(5) Idiopathic
(6) Tumor of the brain
   a. Cerebellar astrocytoma
   b. Pontine glioma

2. Noncomitant-the angle of deviation varies in different directions of gaze
   A. Abducens palsy (p. 161)
   B. Accommodative spasm
   C. Blowout fracture
   D. Divergence paralysis
   E. Drug use (marihuana)
   F. Duane syndrome
   G. Myasthenia gravis
   H. Thyroid myopathy

3. "V" pattern esotropia-deviation greater in downward gaze
   A. Underaction-superior oblique muscles
   B. Overaction-inferior oblique muscles

4. "A" pattern esotropia
   A. Underaction-inferior oblique muscles

5. Monocular esotropia-one eye may be used to the exclusion of the other; amblyopia is usual in the deviating eye

6. Esotropia-near/distance disparity
   A. High accommodation convergence-accommodation (AC/A) ratio-greater convergence for near than for distance, causing greater esodeviation for near than for distance
   B. Convergence excess-greater esodeviation for near than for distance
   C. Divergence insufficiency-greater esodeviation for distance than for near


**PSEUDOEXOTROPIA (OCULAR APPEARANCE OF EXOTROPIA WHEN NO MANIFEST DEVIATION OF VISUAL AXIS IS PRESENT)**

1. Displaced macula (heterotopia of the macula; see p. 451)
2. Heterochromia when the lighter-colored eye appears to diverge (see p. 368-372)
3. Hypertelorism with wide interpupillary distance
4. Exophthalmos
5. Positive-angle kappa-pupillary light reflex displaced nasally (see decentered papillary light reflex, p. 361)
6. Narrow lateral canthus
7. Wide palpebral fissure


EXOPHORIA AND EXOTROPIA (VISUAL AXIS DEVIATED OUTWARD; MAY BE LATENT OR MANIFEST)

1. Comitant
   A. Refractive-myopic refractive error cause of deviation (low AC/A ratio)
   B. Nonrefractive-refractive error not cause of deviation
   C. Anomalous insertion of horizontally acting muscles
   D. Abnormal check ligaments
   E. Faulty innervational development
   F. Autosomal-dominant trait
   G. Idiopathic

2. Noncomitant
   A. Convergence insufficiency
   B. Divergence excess
   C. Duane type II, III
   D. Internuclear ophthalmoplegia
   E. Myasthenia gravis
   F. Third nerve palsy
   G. Thyroid myopathy

3. Pattern exotropia
   A. V exotropia-deviation greater in upward than in downward gaze
      (1) Underaction superior oblique
      (2) Overaction inferior oblique
   B. Exotropia-deviation greater in downward than in upward gaze
      (1) Underaction inferior oblique muscle
      (2) Overaction superior oblique muscle


**PSEUDOHYPERTROPIA**

1. Facial asymmetry with one eye placed higher than the other
2. Unilateral coloboma of lid
3. Unilateral ptosis


**HYPERPHORIA AND HYPERTROPIA (VISUAL AXIS DEVIATED UPWARD; MAYBE MANIFEST OR LATENT)**

1. Nonparalytic hypertropia  
   A. Abnormal insertion of muscles  
   B. Abnormal fascial attachments  
   C. Complications of systemic diseases, such as myasthenia gravis, thyrotoxicosis, and orbital tumors

2. Paralytic hypertropia-isolated cyclovertical muscle palsy  
   A. Brainstem disease  
   B. Fourth nerve palsy  
   C. Multiple sclerosis  
   D. Skew deviation  
   E. Third nerve palsy

3. Double hyperphoria (alternating circumduction)-fuses, but cover test shows alternating hyperphoria

4. Apparent paralysis of elevation of one eye  
   A. Local neuromuscular and orbital causes  
      (1) Dysthyroid ophthalmoplegia (noncongestive and congestive form)  
      (2) Myasthenia gravis (Erb-Goldflam syndrome)  
      (3) Orbital floor fracture  
      (4) Progressive supra nuclear ophthalmoplegia  
      (5) Oculomotor nerve paresis superior division  
      (6) Unilateral double-elevator palsy, congenital dysfunction of superior rectus and inferior oblique muscles  
      (7) Myositis  
         a. "Collagen diseases"  
         b. Infectious myositis  
         c. Trichinosis  
      (8) Orbital tumors  
         a. Dermoid cyst  
         b. Hemangioma  
         c. Lymphoma  
         d. Meningioma  
         e. Optic nerve glioma  
         f. Previous strabismus surgery
g. Rhabdomyosarcoma
(9) Systemic amyloidosis with ocular muscle infiltration
(10) Vertical retraction syndrome (Parinaud syndrome)
(11) Superior oblique tendon sheath syndrome (Brown syndrome)

B. Skew deviation due to a central nervous system lesion—one eye is above the other; may be the same for all directions of gaze or vary in different directions of gaze
(1) Unilateral labyrinthine disease
(2) Cerebellar tumors, such as astrocytomas and medulloblastomas
(3) Acoustic neuromas
(4) Vascular accidents of pons and cerebellum, such as thrombosis of cerebellar and pontine arteries
(5) Unilateral internuclear ophthalmoplegia and less frequently bilateral internuclear ophthalmoplegia
(6) Compressive lesions, such as platybasia and Arnold-Chiari malformation
(7) Brain-stem arteriovenous malformations
(8) Aberrant regeneration of third nerve

C. Central nervous system lesions
(1) Arteriosclerosis, thrombosis, arteritis (syphilitic), or embolus of fine vessels to midbrain

5. Apparent paralysis of elevation of both eyes
A. Physiologic in older persons
B. Parinaud syndrome (divergence paralysis)
C. Chronic progressive external ophthalmology (CPEO)
D. Progressive supranuclear palsy
E. Myasthenia gravis
F. Midbrain lesion
   (1) Upgaze center
   (2) Bilateral third nerve palsy
   (3) Dorsal midbrain syndrome
G. Congenital fibrous syndrome
H. Thyroid myopathy
I. Metastatic tumor (breast cancer)

6. Paralysis of downward gaze
A. Reverse Parinaud syndrome
B. Associated with choreoathetotic syndromes
C. Parkinsonian syndromes
D. Myasthenia gravis
E. Miscellaneous


BROWN SUPERIOR OBLIQUE TENDON SHEATH SYNDROME
(LIMITATION OF ELEVATION IN ADDUCTION THAT RESEMBLES AN UNDERACTION OF INFERIOR OBLIQUE MUSCLE)

1. Congenital onset
   A. Congenital Brown syndrome
   B. Inelastic muscle-tendon complex
2. Anomalies of Superior Oblique Tendon fibers
3. Congenital pseudo-Brown Syndrome-anomalous inferior orbital adhesions
4. Posterior orbital bonds
5. Acquired onset
   A. Acquired Brown syndrome
   B. Peritrochlear scarring and adhesions
   C. Chronic sinusitis
6. Trauma-superior temporal orbit
7. Blepharoplasty and fat removal
8. Lichen sclerosis at astrophicus and morphea
9. Tendon-trochlear inflammation and edema
10. Idiopathic inflammatory (pain and click)
11. Trochlearitis with superior oblique myositis
12. Acute sinusitis
13. Adult rheumatoid arthritis
14. Juvenile rheumatoid arthritis
15. Systemic lupus erythematosus
16. Possibly distant trauma (CPR [cardiopulmonary resuscitation] and long-bone fractures)
17. Possibly postpartum hormonal change
18. Superior nasal orbital mass
19. Glaucoma implant
20. Neoplasm
21. Tight or inelastic superior oblique muscles
22. Thyroid disease (inelastic muscles)
23. Peribulbar anesthesia (inelastic tendon)
24. Hurler Scheie syndrome (inelastic tendon)
25. Superior oblique tuck (short tendon)
26. Idiopathic
27. Acquired pseudo-Brown syndrome
28. Orbital fracture
29. Retinal band around inferior oblique muscle
30. Inferior temporal adhesion
31. Following double plate Molteno implantation


**DUANE SYNDROME**
Congenital ocular motility disorder is characterized by limited abduction or limited adduction. The palpebral fissure narrows on attempted adduction.

1. Type 1 characteristics
   a. A or V phenomena
   b. Defective abduction
   c. Palpebral fissure narrowing on adduction
   d. Retraction of the globe
   e. Updrift or downdrift of the affected eye on adduction or attempted abduction

2. Type 2 characteristics
   a. Abduction appears to be normal or only slightly limited
   b. Distinct narrowing of the palpebral fissure and retraction of the globe on attempted adduction.
   c. Limitation or complete palsy of adduction with exotropia of the paretic eye.

3. Type 3 characteristics
   a. Limitation or absence of both abduction and adduction of the affected eye
   b. Globe retraction and narrowing of the palpebral fissure on attempted adduction


**OCULOMOTOR APRAXIA**
Oculomotor apraxia is defined as defective or absent horizontal voluntary eye movements and includes head thrusting to look at objects to the side.

1. Ataxia-telangiectasia syndrome
2. Brain tumor
   A. Astrocytoma
   B. Lipoma
3. Isolated
4. Male predominance
5. Neurofibromatosis
6. Oral-facial-digital syndrome type II
7. Post cardiac surgery


**MONOCULAR LIMITATION OF ELEVATION OF ADDUCTED EYE WITH FORCED DUCTION TEST [IN ELEVATION AND ADDUCTION STRABISMUS WITH RESTRICTED MOTILITY]**

1. Acquired
   A. Thyroid myopathy
   *B. Excessive recession or resection of muscle
   C. Orbital fracture
   D. Retinal detachment operation
   E. Strabismus surgery complicated by adhesions
   F. CPEO
2. Congenital
   A. Congenital fibrous syndrome
   B. Neurogenic paralysis with secondary contracture of antagonist muscle
   C. Duane retraction syndrome
   D. Brown superior oblique tendon sheath syndrome
   E. Strabismus fixus


**CYCLIC, RECURRENT, REPETITIVE, EPISODIC DISORDERS OF EXTRAOCULAR MUSCLES**

1. Cyclic strabismus
   A. Associated with frontoorbital fibrous dysplasia
   B. Associated with Graves disease
   C. Associated with optic atrophy
D. Cyclic superior oblique palsy
E. Cyclic third nerve palsy
F. Esotropia, vertical
   (1) Comitant
   (2) Noncomitant

2. Cyclic vertical deviation
3. Diabetic nerve palsies
4. Myasthenia gravis
5. Oculogyric crisis (see p. 152)
6. Periodic alternating gaze deviation
7. Periodic alternating nystagmus
8. Periodic vertical nystagmus
   A. Associated with potassium abnormality
   B. Familial
9. Petit mal epilepsy
   A. Exotropia
   B. Upward deviation
10. Ping-pong gaze
11. Recurrent sixth nerve paralysis in children (see p. 169)
12. Spasmus nutans
13. Twitch of lids (orbicularis)

Hamed L. *Cyclic periodic disorders in diagnostic problems in clinical ophthalmology.*


**SYNDROMES AND DISEASES ASSOCIATED WITH STRABISMUS**

1. Esotropia syndrome
2. Exotropia syndrome
3. Aarskog syndrome (facial-digital-genital syndrome)
4. Aberfeld syndrome (congenital blepharophimosis associated with generalized myopathy)
5. Achondroplasia
6. Addison pernicious anemia
7. African eyeworm disease
8. Albinism
9. Albright hereditary osteodystrophy (pseudohypoparathyroidism)
10. Amyloidosis
11. Apert syndrome (acrocephalosyndactylism syndrome)
12. Arnold-Chiari syndrome (platybasia syndrome)
13. Arylsulfatase A deficiency syndrome
14. Aspergillosis
15. Axenfeld-Schurenberg syndrome (cyclic oculomotor paralysis)
16. Bacterial endocarditis
17. Bang disease (brucellosis)
18. Behçet syndrome (oculobuccogenital syndrome)
19. Benedikt syndrome (tegmental syndrome)
20. Best disease (vitelliform dystrophy)
22. Bing-Neel syndrome (associated with macroglobulinemia and central nervous system symptoms)
23. Bloch-Sulzberger disease (incontinentia pigmenti)
24. Blocked nystagmus syndrome (nystagmus blockage syndrome)
25. Bonnet-Dechaume-Blanc syndrome (neuroretinoangiomatosis syndrome)
26. Bonnevie-Ullrich syndrome (pterygolymphangieciasia)
27. Botulism
28. Brown-Marie syndrome (hereditary ataxia syndrome)
29. Canine tooth syndrome (class VII superior oblique palsy)
30. Cerebral palsy
31. Chediak-Higashi syndrome (anomalous leukocytic inclusions with constitutional stigmata)
32. Chromosome partial deletion (short-arm) syndrome (Wolf syndrome)
33. Chromosome 13q partial deletion (long-arm) syndrome (thirteen Q syndrome)
34. Chromosome partial deletion (long-arm) syndrome (DeGrouchy syndrome)
35. Chromosome partial (short-arm) partial deletion syndrome
36. Congenital syphilis
37. Convergence insufficiency syndrome
38. Craniofacial dysplasia (Freeman-Sheldon syndrome; whistling face syndrome)
39. Craniosenosis
40. Cri-du-chat syndrome (Cry of the cat syndrome)
41. Crohn disease (granulomatous ileocolitis)
42. Crouzon disease (craniofacial dysostosis)
43. Cushing syndrome (II) (cerebellopontine angle syndrome)
44. Cysticercosis
45. Cytomegalic inclusion disease, congenital
46. Dawson disease (subacute sclerosing panencephalitis)
47. De Lange syndrome (congenital muscular hypertrophy-cerebral syndrome)
48. Dengue fever
49. Devic syndrome (ophthalmoencephalomyelopathy)
50. Diabetes mellitus
51. Diphtheria
52. Diamond-Blackfan syndrome
53. Down disease (mongolism, trisomy 21)
54. Drugs, including the following:
   alcohol        chloroform        iothalamate meglumine
   baclofen       cholecalciferol    and sodium
   calcitriol     ergocalciferol     iothalamic acid
   chloramphenicol(?)  insulin  isocarboxazid
measles and rubella virus vaccine (live)  mumps virus vaccine (live)  rubella virus vaccine (live)
measles virus vaccine (live)  nialamide  tranlycypromine
measles, mumps, and rubella virus vaccine (live)  pemoline  tripelennamine
metoclopramide  pentylenotetrazol  vitamin A
metrizamide  phenelzine  vitamin D₂
rubella and mumps virus vaccine (live)  vitamin D₃

54. Drummond syndrome (idiopathic hypercalcemia)
55. Duane syndrome (retraction syndrome)
56. Ectrodactyly-ectodermal dysplasia clefting syndrome (EEC syndrome)
57. Ehlers-Danlos disease (fibrodysplasia elastica generalisata)
58. Electrical injury
59. Ellis-van Creveld syndrome (chondroectodermal dysplasia)
60. Encephalitis, acute
61. Engelmann syndrome (osteopathia hyperostotica scleroticans multiplex infantalis)
62. Epidermal nevus syndrome (ichthyosis hystrix)
63. Erb-Goldflam disease
64. Fetal alcohol syndrome
65. Fibrosarcoma
66. François dyscephalic syndrome
67. Gaucher syndrome (glucocerebrosidase storage disease)
68. Gangliosidosis
   A. Infantile (GM1)
   B. Juvenile (GM2)
69. Goltz syndrome (focal dermal hypoplasia syndrome)
70. Gorlin-Goltz syndrome (multiple basal cell nevi syndrome)
71. Greig syndrome (ocular hypertelorism syndrome)
72. Grönlund-Strandberg syndrome (systemic elastodystrophy)
73. Hemangiomias
74. Hemifacial hyperplasia with strabismus (Bencze syndrome-autosomal dominant)
75. Hemifacial microsomia (otomandibular dysostosis)
76. Homocystinuria
77. Hurler disease (mucopolysaccharidoses type I)
78. Hutchinson syndrome (adrenal cortex neuroblastoma with orbital metastasis)
79. Hydrocephalus, congenital
80. Hydrophobia (rabies)
81. Hyperthyroidism
82. Hypocalcemia
83. Hypomelanosis of Ito syndrome (incontinentia pigamenti a-chromians)
84. Hypothermal injury
85. Hysteria
86. Infectious mononucleosis
87. Influenza
88. Jacobs syndrome (triple X syndrome)
89. Johnson syndrome (adherence syndrome)
90. Klippel-Feil syndrome (congenital brevicollis)
91. Koerber-Salus-Elschnig syndrome (nystagmus retractorius syndrome)
92. Kohn-Romano syndrome (telecanthus, ptosis, epicantus inversus, blepharophimosis)
93. Krause syndrome (congenital encephaloophthalmic dysplasia)
94. Kugelberg-Welander syndrome (progressive muscle atrophy)
95. Kussmaul disease (necrotizing angitis)
96. Larsen syndrome (hypertelorism, microtia, and facial clefting)
97. Laurence-Moon-Bardet-Biedl syndrome (retinitis pigmentosa-polydactyly-adiposogenital syndrome)
98. Leigh disease (subacute necrotizing encephalomyelopathy)
99. Leukemia
100. Linear nevus sebaceous of Jadassohn
101. Lowe syndrome (oculocerebroprenal syndrome)
102. Lymphangioma
103. Lymphedema
104. Lymphoid hyperplasia (Burkitt lymphoma)
105. Malaria
106. Malignant hyperpyrexia syndrome
107. Malignant hyperthermia syndrome
108. Maple syrup urine disease
109. Marcus Gunn syndrome (jaw-winking syndrome)
110. Marfan syndrome (arachnодactyly-dystrophia mesodermalis congenita)
111. Measles
112. Melnick-Needles syndrome (osteodysplasty)
113. Mieten syndrome (corneal opacity, nystagmus, flexion contracture, growth failure)
114. Millard-Gubler syndrome (abducens-facial hemiplegia alternans)
115. Möbius syndrome (congenital paralysis of sixth or seventh nerves)
116. Monofixation syndrome (blind-spot syndrome)
117. Morning glory syndrome (hereditary central glial anomaly of the optic disk)
118. Mucocele
119. Mucormycosis
120. Mulibrey nanism syndrome (Perheentupa syndrome)
121. Multiple lentigines syndrome (leopard syndrome)
122. Multiple sclerosis
123. Mumps
124. Myasthenia gravis (Erb-Goldflam syndrome)
125. Naegeli syndrome (melanophoric nevus syndrome)
126. Nematode ophthalmia syndrome (toxocariasis)
127. Neonatal hemolytic disease of hyperbilirubinemia
128. Neuroblastoma
129. Nevus sebaceous of Jadassohn
130. Nevoid basal cell carcinoma syndrome
131. Nielsen syndrome (exhaustive psychosis syndrome)
132. Noonan syndrome (male Turner syndrome)
133. Noone-Milroy-Meige disease (congenital trophedema)
134. Nothnagel syndrome (ophthalmoplegia cerebellar ataxia syndrome)
135. Nystagmus compensation syndrome
136. Obesity-cerebrolocular-skeletal anomalies syndrome
137. Ocular vaccinia
138. Oculocerebellar tegmental syndrome
139. Oculo-oto-ororenoerythropoietic syndrome
140. Ophthalmoplegic retinal degeneration syndrome
141. Orbital floor syndrome (Dejean syndrome)
142. Paget syndrome (osteitis deformans)
143. Pallister-Killian syndrome
144. Papillon-Léage and Psauame syndrome (orodigital facial syndrome)
145. Parkinson syndrome
146. Parry-Romberg disease (progressive facial hemiatrophy)
147. Periocular and ocular metastatic tumors
148. Pertussis (whooping cough)
149. Pierre Robin syndrome (micrognathia-glossoptosis syndrome)
150. Polymyalgia rheumatica
151. Postvaccinial ocular syndrome
152. Pseudoophthalmoplegia syndrome (Roth-Bielschowsky syndrome)
153. Prader-Willi syndrome (hypotonia-obesity syndrome)
154. Pseudohypoparathyroidism (Seabright-Bantam syndrome)
155. Reiter syndrome (conjunctivourethrosynovial syndrome)
156. Relapsing fever
157. Retinoblastoma
158. Ring chromosome 18
159. Ring D chromosome
160. Ring dermoid syndrome
161. Rocky Mountain spotted fever
162. Rubella, congenital
163. Rubinstein-Taybi syndrome (broad-thumb syndrome)
164. Sabin-Feldman syndrome
165. Sandifer syndrome (hiatus hernia-torticollis syndrome)
166. Schilder syndrome (encephalitis periaxialis diffusa)
167. Seckel bird-headed dwarfism
168. Skew deviation syndrome
169. Smallpox
170. Smith-Lemli-Opitz syndrome (cerebrohepatorenal syndrome)
171. Spongy degeneration of the white matter
172. Streptococcus
173. Superior oblique tendon sheath syndrome (Brown syndrome)
174. Supravalvular aortic stenosis syndrome (infantile hypercalcemia with mental retardation)
175. Tay-Sachs syndrome (familial amaurotic idiocy)
176. Temporal arteritis syndrome (cranial arteritis syndrome)
177. Terson syndrome (subarachnoid hemorrhage syndrome)
Horizontal gaze palsy comprises an inability to look horizontally in a given direction; analysis includes optically induced movement, voluntary or command movement, pursuit movement, or vestibular movement.

1. Horizontal palsy of voluntary and command movement-frontal lobe gaze center (second frontal gyrus, Brodmann area 8) or in the corresponding internal capsule gaze palsy of side opposite lesion; it may be associated with facial palsy as well as hemiparesis or hemiplegia toward the side of the gaze palsy, caloric ocular movement intact, and doll’s head intact.

2. Horizontal palsy of command and pursuit movements, optically induced movements, and vestibular movements-pons and posterior longitudinal bundle; the gaze palsy is toward the side of the lesion, facial palsy often present, caloric and doll’s head responses are absent.

Oscillations of eyes (involuntary, rapid, to-and-fro movement of eyes having no rhythm or regularity)

1. Ocular dysmetria- "overshooting" of the eyes with attempted fixation; horizontal ocular dysmetria is associated with lesions of the cerebellum or its pathways as in Friedreich ataxia, Huntington chorea, spinocerebellar degeneration, internuclear ophthalmoplegia, manic depression, alcoholism, schizophrenia, severe diffuse brain damage, cerebellopontine angle tumors, hereditary ectodermal dysplasia with olivopontocerebellar degeneration, Fabry disease (glycosphingolipid lipidosis), vestibulocerebellar ataxia, and toluene abuse

2. Ocular flutter-flutter-like oscillations that are intermittent, rapid, to-and-fro motions, or motions of equal amplitude, interrupt maintained fixation; horizontal ocular flutter is associated with lesions of the cerebellum or its pathways as in limb ataxia, multiple sclerosis, poliomyelitis, neoplasms, or vascular accident.
3. Opsoclonus—irregular, hyperkinetic, multidirectional, spontaneous eye movement that persists in sleep
   A. Infections
      (1) Coxsackie B3
      (2) Encephalitis—mild, severe, viral or post infections (including St. Louis encephalitis)
      (3) Haemophilus influenzae
      (4) Meningitis
      (5) Paratyphi A
      (6) Psittacosis
      (7) Salmonella
   B. Tumors
      (1) Breast malignancy
      (2) Bronchogenic carcinoma
      (3) Glioblastoma
      (4) Neuroblastoma
      (5) Thyroid carcinoma
      (6) Uterine carcinoma
   C. Toxins and drugs
      (1) Amitriptyline
      (2) Chlordecone
      (3) Lithium-haloperidol (Haldol)
      (4) Thallium
      (5) Toluene abuse
   D. Other
      (1) Acute cerebellar ataxia
      (2) Friedreich ataxia
      (3) Multiple sclerosis (disseminated sclerosis) (rare)
      (4) Nonketotic coma
      (5) Sign of "myoclonic encephalopathy of infancy"
      (6) Vascular accidents
      (7) Vertebrobasilar insufficiency
4. Lightning eye movements (ocular myoclonus)—rapid to-and-fro movements of small conjugate saccades; probably because of bilateral abnormality of a pontine paramedical zone and pretectal lesions, such as vascular, inflammatory, neoplastic, demyelinating, or trauma of tegmentum as thyroid, lung or uterus carcinoma, neuroblastoma, Menzel hereditary ataxia, pontine myelinolysis, coxsackie B infection, cherry-red spot myoclonus syndrome, Ramsay-Hunt syndrome, and L-tryptophan


**COGWHEEL EYE MOVEMENTS (JERKY INACCURATE PURSUIT MOVEMENTS)**

1. Basal ganglia disease
   A. Anoxia
   B. Carbon disulfide poisoning
   C. Carbon monoxide poisoning
   D. Drugs, including:
   - acetophenazine
   - alcohol
   - allobarbital
   - alprazolam
   - alseroxylon
   - amitriptyline
   - amobarbital
   - aprobarbital
   - barbital
   - bromide
   - bupivacaine
   - butabarbital
   - butalbital
   - butallylonal
   - butaperazine
   - butethal
   - carphenazine
   - chloral hydrate
   - chlordiazepoxide
   - chlorprocaaine
   - chlorpromazine
   - clonazepam
   - clorazepate
   - cyclobarbital
   - cyclopentobarbital
   - deserpidine
   - desipramine
   - diazepam
   - diethazine
   - ethopropazine
   - etidocaine
   - fluphenazine
   - flurazepam
   - halazepam
   - heptabarbital
   - hexethal
   - hexobarbital
   - imipramine
   - heptobarbital
   - lorazepam
   - meprobamate
   - mepivacaine
   - mesoridazine
   - metharbital
   - methdilazine
   - methitural
   - methohexital
   - methotropine
   - oxazepam
   - pentobarbital
   - perphenazine
   - phencyclidine
   - phenobarbital
   - piperacetazine
   - prazepam
   - pritracaine
   - primidone
   - probarbital
   - procaine
   - prochlordperazine
   - promazine
   - promethazine
   - propiomazine
   - propoxycaïne
   - protriptyline
   - rauwolfia serpentina
   - rescinnamine
   - reserpine
   - secobarbital
   - syrosingopine
   - talbutal
   - temazepam
   - thiamylal
   - thieplperazine
   - thiopental
   - thioproprazine
   - thioridazine
   - triazolam
   - trifluoperazine
   - triflupromazine
   - trimepazine
   - vinbarbital

E. Exposure to manganese
F. Idiopathic
G. Parkinsonism (shaking palsy)
H. Trauma

2. Cerebellar tumors
   A. Astrocytomas
   B. Hemangioblastomas
   C. Medulloblastomas

3. With homonymous hemianopia, indicates parietal or occipital lobe involvement


**PENDULAR NYSTAGMUS**

Pendular nystagmus comprises oscillations that are approximately equal in rate in two directions; they may be horizontal or vertical.

1. Albinism in which the macula does not develop
2. Aniridia (see p. 364-365)
3. Bilateral chorioretinal lesions involving the macula in early infancy (congenital toxoplasmosis)
4. Brainstem or cerebellar dysfunction
   *5. Congenital-cause unknown, may be inherited as autosomal dominant recessive or X-linked recessive trait; not infrequently associated with astigmatism and convergent strabismus*
   *6. Congenital cataracts*
7. Congenital glaucoma
8. Corneal scars
9. Demyelinating disease
10. High myopia of early life
11. Internuclear ophthalmoplegia
12. Laurence-Moon-Bardet-Biedl syndrome (retinitis pigmentosa-polydactyly-adiposogenital syndrome)
13. Leber congenital amaurosis
14. Monocular or binocular visual deprivation
15. Optic nerve hypoplasia, coloboma
   *16. Total color blindness (monochromatism)*
17. Work in poor illuminations (e.g., mining) (rare)

HORIZONTAL JERK NYSTAGMUS (HORIZONTAL OSCILLATORY MOVEMENT OF EYES WITH A FAST AND SLOW PHASE)

1. Albinism
2. Amblyopia (manifest latent nystagmus)
3. Cerebellar disease, acute or chronic; fast component to side of lesion
4. Chediak-Higashi syndrome (anomalous leukocytic inclusions with constitutional stigmata)
5. Congenital achromatopsia
6. Congenital cataracts
7. Congenital stationary night-blindness
8. Congenital X-linked, dominant, recessive
9. Leber congenital amaurosis
10. Lesions of labyrinth (e.g., Meniere syndrome) or when one labyrinth has been removed
11. Neoplastic angioendotheliomatosis
12. Optic nerve hypoplasia, coloboma
13. Vestibular nuclei involvement as in persons with multiple sclerosis


VERTICAL NYSTAGMUS (SPONTANEOUS VERTICAL OSCILLATIONS OF EYES)

1. Upbeat nystagmus-nystagmus in which the fast component is upward and usually most marked when the gaze is directed upward; usually due to a lesion in the posterior fossa
   A. Brainstem lesion, such as that of the vestibular nuclei
   B. Cerebellar disease-acute or chronic, especially in the vermis
   C. Cerebellar degeneration
   D. Drugs-barbiturates and Dilantin (phenytoin)
E. Encephalitis
F. Labyrinth disease—rare; has no lateralizing value
G. Multiple sclerosis
H. Idiopathic

2. Downbeat nystagmus—nystagmus in which the fast component is downward and usually most marked when the gaze is directed downward; probably due to a lesion in the lower end of the brain stem or cerebellum
   A. Alcoholic cerebellar disease
   B. Aneurysm of the supraclinoid part of left carotid siphon
   C. Arnold-Chiari malformation—herniation of cerebellar tonsils and part of medulla through foramen magnum
   D. Cerebellar atrophy/degeneration
   E. Carbamazepine
   F. Deformities of cervical spine
   G. Diabetes mellitus
   H. Encephalopathy
   I. Ependymoma of posterior part of the fourth ventricle
   J. Idiopathic
   K. Insufficiency of basilar artery
   L. Klippel-Feil anomaly—upward displacement of odontoid process into foramen magnum
   M. Meningioma extending into pontine cistern
   N. Morphine poisoning
   O. Multiple sclerosis (disseminated sclerosis)
   P. Neurogenic muscular atrophy
   Q. Platybasia (cerebellomedullary malformation syndrome)


**ROTARY NYSTAGMUS (ROTARY OSCILLATORY MOVEMENT OF EYES)**

1. Benign paroxysmal positional nystagmus—fast component toward lower ear
  *2. Cerebellar disease—acute or chronic
  3. Cerebrotendinous xanthomatosis
  4. Encephalitis
  5. Lesion of vestibular nuclei in floor of fourth ventricle associated with multiple sclerosis, syringobulbia, or thrombosis of posteroinferior cerebellar artery or its branches
*6. Superior oblique myokymia - benign, intermittent, uniocular
7. Vestibular involvement (e.g., labyrinthitis, Meniere syndrome)


**SEE-SAW NYSTAGMUS**

One eye moves up as other eye moves down; in addition, there is torsion of eyes - eye moving up intorts, and eye moving down extorts. This nystagmus probably is due to lesions located in mesodiencephalic region, hypothalamus, and thalamus; it may be associated with bitemporal hemianopsia and reduced vertical optokinetic nystagmus.

1. Brainstem vascular disease
*2. Chiasmal glioma
*3. Chromophobe adenoma of the pituitary gland involving the optic chiasm and third ventricle
4. Congenital
*5. Craniopharyngioma, involving the optic chiasm and hypothalamus
6. Head injury with fracture of frontal
7. Idiopathic
8. Multiple sclerosis
9. Oligodendroglioma involving the pons and third ventricle
10. Postoperative after strabismus surgery
11. Retinitis pigmentosa
12. Septooptico dysplasia
13. Suprasellar epidermoid tumor involving optic chiasm and hypothalamus
14. Syringomyelia and syringobulbia
15. Toxoplasmosis of the brainstem


**RETRACTION NYSTAGMUS**

Spasmodic retraction of eyes can occur when an attempt is made to move them in any direction; it is caused by lesions of midbrain, especially lesions in the vicinity of aqueduct of Sylvius).

1. Arteriovenous aneurysm
2. Brucellosis (Bang disease)
3. Cysticercus cyst
4. Ependymoma
5. Koerber-Salus-Elschnig syndrome (sylvian aqueduct syndrome)
*6. Parinaud syndrome (paralysis of vertical movement)
7. Vascular lesions


**MONOCULAR NYSTAGMUS**

1. Horizontal
   - A. Lesions of optic nerve, chiasm, midbrain, or brainstem
   - B. Nervous system disease, such as multiple sclerosis, epidemic meningitis, and congenital syphilis
   - C. Seizures
   - D. Superior oblique myokymia-benign, intermittent, uniocular
   - E. Spasmus nutans-most common cause in children
   - F. Tumors of brainstem
   - G. Unilateral amblyopia
   - H. Unilateral astigmatism or high refractive error
   - I. Unilateral opacity of the ocular media

2. Vertical
   - A. Multiple sclerosis
   - B. Myokymia of lower eyelid
   - C. Sleep
   - D. Spasmus nutans
   - E. Unilateral amblyopia


**PERIODIC ALTERNATING NYSTAGMUS**

Periodic alternating nystagmus is central vestibular nystagmus with rhythmic jerk type of nystagmus that undergoes phasic or cyclic changes in amplitude and direction.

1. Arnold-Chiari malformation
2. Cerebral trauma or fractured skull
3. Chiasmal lesion, such as craniopharyngioma
4. Chronic otitis media  
5. Congenital  
6. Diabetes mellitus  
7. Encephalitis  
8. Friedreich hereditary ataxia  
9. Meningioma of tentorium cerebelli, cerebellar glioma, and cholesteatoma of the cerebellopontine angle  
10. Head trauma  
11. Mesencephalic brainstem and cerebellar disease  
12. Multiple sclerosis (disseminated sclerosis)  
13. Syringobulbia (Passow syndrome)  
14. Syphilitic optic atrophy  
15. Tumor of the corpus callosum  
16. Vertebrobasilar artery insufficiency  
17. Vestibular nuclei lesions  
18. von Recklinghausen syndrome (neurofibromatosis)


**POSITIONAL NYSTAGMUS**

Positional nystagmus is nystagmus that appears or changes in form or intensity after certain positional changes of the head indicate vestibular stimulation.

1. After general anesthesia  
2. After head injury  
3. Drugs, including the following:
   - acetophenazine  
   - alcohol  
   - allobarbital  
   - alprazolam  
   - amiodarone  
   - amitriptyline  
   - amobarbital  
   - amodiaquine  
   - aprobarbital  
   - aspirin  
   - auranofin  
   - aurothioglucose  
   - aurothioglycanide  
   - Bacille Calmette-Guerin (BCG) vaccine  
   - baclofen  
   - barbital  
   - bleomycin(?)  
   - bromide  
   - bromisovalum  
   - broxyquinoline  
   - bupivacaine  
   - butabarbital  
   - butalbital  
   - butallylonal  
   - butaperazine  
   - butethal  
   - calcitriol  
   - carbamazepine  
   - carboxaminexine  
   - carbon monoxide  
   - carbroxamine  
   - carphenazine  
   - carisoprodol  
   - carphenazime  
   - cefadroxil  
   - cefamandole  
   - cefazolin  
   - cefoperazone  
   - ceforanide  
   - cefotaxime  
   - cefotetan  
   - cefoxitin  
   - cefsulodin  
   - ceftazidime  
   - ceftizoxime
4. Inner ear pathologic changes, including hemorrhage, inflammation, thrombosis, emboli, circulatory and secretory conditions
5. "Normal" persons
6. Other causes include neuritis, meningitis, tumors, vascular anomalies, degeneration, atrophy, syphilis, arteriosclerosis, hypertonia, vasomotor disturbance, allergic and toxic conditions, cranial trauma, hemorrhage, emboli, or thrombosis


**OPTOKINETIC NYSTAGMUS**

Normal physiologic nystagmus is obtained by watching moving targets; slow components in direction targets are moving, and fast component move in the opposite direction. Abnormal optokinetic nystagmus can be seen in the following:

1. Aberrant regeneration of third nerve-absent vertical optokinetic nystagmus, normal horizontal optokinetic nystagmus
2. Internuclear palsies-horizontal targets bring out dissociation of ocular response movements
3. Lesions of optic tract, geniculate body, temporal and occipital lobes show no asymmetry of horizontal optokinetic responses
4. Lesions of parietal lobe give asymmetric horizontal optokinetic responses
5. Occipital lobe lesions with homonymous hemianopia and asymmetrical horizontal optokinetic responses suggests a mass lesion extending into parietal lobe rather than a vascular lesion
6. Parinaud syndrome-vertical optokinetic nystagmus with targets moving downward enhances the retraction nystagmus seen on attempted upgaze.
7. Parkinsonism (shaking palsy)-vertical optokinetic nystagmus may be reduced
8. See-saw nystagmus-vertical optokinetic nystagmus may be reduced either upward or downward (see p. 144)
9. Test for malingering in "blind" eye or eyes with normal optokinetic responses


**SYNDROMES AND DISEASES ASSOCIATED WITH NYSTAGMUS**

1. African eye-worm disease
2. Albers-Schonberg syndrome (osteosclerosis fragilis generalisata)
3. Albinism, ocular
4. Alexander disease
5. Anterior spinal artery syndrome
6. Apert syndrome (acrocephalosyndactylism syndrome)
7. Arnold-Chiari syndrome (platybasia syndrome)
8. Arylsulfatase A deficiency syndrome
9. Babinski-Nageotte syndrome (medullary tegmental syndrome)
10. Bacterial endocarditis
11. Bassen-Kornzweig syndrome (abetalipoproteinemia)
12. Behçet syndrome (dermatostomatoophthalmic syndrome)
13. Behr disease (optic atrophy-ataxia syndrome)
14. Bielschowsky-Lutz-Cogan syndrome (internuclear ophthalmoplegia)
15. Bloch-Sulzberger disease (incontinentia pigmenti)*
16. Blocked nystagmus syndrome (nystagmus blockage syndrome)
17. Bonnet-Dechaume-Blanc syndrome (neuroretinoangiomatosis syndrome)
18. Botulism
20. Caisson syndrome (bends)
21. Canavan disease (spongy degeneration of the white matter)
22. Cerebral palsy
23. Cestan-Chenais syndrome (combination of Babinski-Nageotte and Avellis syndrome)
24. Charcot-Marie- Tooth disease (progressive peroneal muscular atrophy)
25. CHARGE syndrome (colomba, heart disease, atresia, choanae, retarded growth and retarded development or central nervous system anomalies, genital hypoplasia, and ear anomalies or deafness syndrome)
26. Chediak-Higashi syndrome (anomalous leukocytic inclusions with constitutional stigmata)
27. Cherry-red spot myoclonus syndrome
28. Chromosome 6p12
29. Chromosome 18, partial deletion (long-arm) syndrome
30. Cockayne syndrome (dwarfism with retinal atrophy and deafness)
31. Cogan syndrome (II) (oculomotor apraxia syndrome)
*32. Cone dysfunction syndrome (achromatopsia)
33. Costen syndrome (temporomandibular joint syndrome)
34. Craniocervical syndrome (whiplash syndrome)
35. Craniofaryngioma
36. Craniostenosis
37. Creutzfeldt-Jakob syndrome (spastic pseudosclerosis)
38. Crouzon disease (craniofacial dysostosis)
39. Curtius syndrome (ectodermal dysplasia with ocular malformations)
40. Cushing syndrome (II) (angle tumor syndrome)
41. Cytomegalic inclusion disease, congenital
42. Dawson disease (subacute sclerosing panencephalitis)
43. de Lange syndrome (congenital muscular hypertrophy-cerebral syndrome)
44. Diencephalic syndrome (autonomic epilepsy syndrome) (Russell syndrome)
45. Disseminated lupus erythematosus (Kaposi-Libman-Sacks syndrome)
*46. Disseminated sclerosis (multiple sclerosis)
47. Down disease (mongolism, trisomy 21)
48. Drummond syndrome (idiopathic hypercalcemia)
49. Eclampsia and preeclampsia
50. Electrical injury
51. Encephalitis, acute
52. Epidermal nevus syndrome (ichthyosis hystrix)
53. Epiphysial dysplasia, microcephaly, and nystagmus-autosomal recessive
54. Extreme hydrocephalus syndrome
55. Fanconi-Turler syndrome (familial ataxic diplegia)
56. Fetal hydantoin syndrome
57. Forsius-Eriksson syndrome (Aland disease)
58. François dyscephalic syndrome
59. Gangliosidosis (generalized gangliosidosis, infantile)
60. General fibrosis syndrome
61. Goltz syndrome (focal dermal hypoplasia syndrome)
62. Gorlin-Chaudhry-Moss syndrome
63. Guillain-Barré syndrome (acute infectious neuritis)
64. Hallervorden-Spatz syndrome (pigmentary degeneration of globus pallidus)
65. Hallgren syndrome (retinitis pigmentosa-deafness-ataxia syndrome)
66. Hand-Schüller-Christian syndrome (histiocytosis X)
67. Hanhart syndrome (recessive keratosis palmoplantaris)
68. Hartnup syndrome (niacin deficiency)
69. Hennebert syndrome (luetic-otic-nystagmus syndrome)
70. Hermansky-Pudlak syndrome (oculocutaneous albinism and hemorrhagic diathesis)
71. Hurler syndrome (mucopolysaccharidoses I-H)
72. Hypervitaminosis D
73. Hypomelanosis of Ito syndrome (incontinentia pigmenti achromians)
74. Hypothyroidism (cretinism)
75. Hysteria
76. Infantile globoid cell leukodystrophy (Krabbe disease)
77. Infantile neuroaxonal dystrophy
78. Infectious mononucleosis
79. Japanese River fever (typhus)
80. Jeune disease (asphyxiating thoracic dystrophy)
81. Kernicterus-high levels of bilirubin in the blood
82. Klippel-Feil syndrome (congenital brevicollis)
83. Koerber-Salus-Elschnig syndrome (sylvian aqueduct syndrome)
84. Kohn-Romano syndrome (blepharophimosis, ptosis, epicanthus inversus, telecanthus)
85. Kugelberg-Welander syndrome (progressive proximal muscle atrophy)
86. Laurence-Moon-Bardet-Biedl syndrome (retinitis pigmentosa-polydactyly-adiposogenital syndrome)*
87. Leber congenital amaurosis syndrome (retinal aplasia)
88. Leigh disease (subacute necrotizing encephalomyelopathy)
89. Lenoble-Aubineau syndrome (nystagmus-myoclonia syndrome)
90. Lermoyez syndrome (form of Meniere disease)
91. Linear nevus sebaceous of Jadassohn
92. Lockjaw (tetanus)
93. Louis-Bar syndrome (ataxia-telangiectasia syndrome)
94. Lowe disease (oculocerebrorenal syndrome)
95. Malignant hyperthermia syndrome
96. Maple syrup urine disease
97. Marfan syndrome (acrachnadactyly-dystrophia mesodermalis congenita)
98. Marinesco-Sjögren syndrome (congenital spirocerebellar ataxia-congenital cataract-oligophrenia syndrome)
99. Meniere syndrome (vertigo, tinnitus, nystagmus)
100. Meningococcemia (meningitis)
101. Mietens syndrome
102. Morning glory syndrome (optic nerve dysplasia, encephalocele)
103. Moyamoya disease (multiple progressive intracranial arterial occlusion)
104. Naegeli syndrome (melanophoric nevus syndrome)
105. Nystagmus, congenital
106. O'Donnell-Pappas syndrome (dominant foveal hypoplasia and presenile cataracts)*
optic nerve hypoplasia, coloboma
107. Papillon-Lerevre syndrome (hyperkeratosis palmoplantaris with periodontosis)
108. Parkinson disease
109. Passow syndrome (status dysraphicus syndrome)
110. Pelizaeus-Merzbacher disease (aplasia axialis extracorticalis congenita) - x-linked
111. Photomyoclonus, diabetes mellitus, deafness, neuropathy, and cerebellar
dysfunction-autosomal dominant
112. Poliomyelitis
113. Posthypoxic encephalopathy syndrome
114. Pyle disease (familial metaphyseal dysplasia)
115. Quincke disease (angioedema)
116. Rubella, congenital
117. Reimann syndrome (hyperviscosity syndrome)
118. Relapsing polychondritis
119. Scaphocephaly syndrome
120. Schilder disease (encephalitis periaxialis diffusa)
121. Seckel syndrome (bird-headed dwarf syndrome)*
122. Septooptic dysplasia (de Morsier syndrome)
123. Smith-Lemli-Opitz syndrome (cerebrohepatorenal syndrome)
124. Sorsby syndrome (hereditary macular coloboma syndrome)
125. Spastic paraplegia - x-linked
126. Split hand with congenital nystagmus, fundal changes, cataracts-autosomal dominant
127. Stannus cerebellar syndrome (riboflavin deficiency)
128. Subclavian steal syndrome
129. Tay-Sachs disease (familial amaurotic idiocy)
130. Traumatic encephalopathy syndrome (punch-drunk syndrome)
131. Tremor, nystagmus, and duodenal ulcer-autosomal dominant
132. Tuomaala-Haapanen syndrome (similar to pseudohypoparathyroidism)
133. Vermis syndrome
134. Vertebral basilar artery syndrome
135. von Economo syndrome (encephalitis lethargica)
136. von Reuss syndrome (galactosemic syndrome)
137. Wagner syndrome (hyaloideoretinal degeneration)
138. Wallenberg syndrome (lateral bulbary syndrome)
139. Werner syndrome (progeria of adults)
140. Wernicke syndrome (superior hemorrhagic polioencephalopathic syndrome)
141. Wildervanck syndrome (cervicooculoacoustic syndrome)
142. Wilson disease (hepatolenticular degeneration)
143. Wolf syndrome (monosomy partial syndrome)
144. Zellweger syndrome (cerebrohepatorenal syndrome of Zellweger)


**OCULOLOGYRIC CRISIS (SPASMODIC AND INVOLUNTARY DEVIATION OF EYES, USUALLY UPWARD, LASTING FROM A FEW MINUTES TO SEVERAL HOURS)**

1. Cerebellar disease
2. Drugs, including the following:

- acetophenazine
- alprazolam
- alseroxylon
- amantadine
- amitriptyline
- amodiaquine
- butaperazine
- carbamazepine
- carphenazine
- chlordiazepoxide
- chloroquine
- chlorpromazine
- chlorprothixene
- cisplatin
- clonazepam
- clorazepate
- clorpromazine
- clorprothixene
- cisplatin
- clonazepam
- clorazepate
- deserpidine
- desipramine
- diazepam
- diethazine
- doxepin
- droperidol
- ethopropazine
- fluphenazine
- flurazepam
- halazepam
- haloperidol
- hydroxychloroquine
- imipramine
- influenza virus vaccine
- levodopa
- lithium carbonate
- lorazepam
- loxapine
- mesoridazine
- methdilazine
- methotrimeprazine
- metoclopramide
- metronidazole
- midazolam
- nitrazepam
- nortriptyline
- oxazepam
- pemoline
- pentazocine
- perazine
- pericyazine
- perphenazine
- phencyclidine
- pimozide
- piperacetazine
- prazepam
- prochlorperazine
- promazine
- promethazine
- propiomazine
- protriptyline
- rauwolfia serpentina
- rescinnamine
- reserpine
- syrosingopine
- temazepam
- thiethylperazine
- thiopropazate
- thioproperazine
- thioridazine
- thiothixene
- triazolam
- trifluoperazine
- trifluperidol
- triflupromazine
- trimeprazine

3. Late manifestations of encephalitis
4. Lesions of fourth ventricle and cerebellum, especially lesions of the flocculus
5. Multiple sclerosis (disseminated sclerosis)
6. Parkinsonism syndrome (shaking palsy)
7. Syphilis (acquired lues)
8. Trauma


**OCULAR BOBBING**

Both globes move synchronously in vertical plane by spontaneously and intermittently dipping downward through an arc of a few millimeters and then return to primary position; reverse ocular bobbing also has been described. Ocular bobbing differs from vertical nystagmus by virtue of absence of a fast and a slow component in movements; it is due to advanced pontine disease.
1. Acute organophosphate poisoning (Diazinon)
2. Associated with palatal myoclonus*
3. Encephalitis
4. Fibrocartilaginous embolism to the anterior spinal artery
5. Hypertensive pontine hemorrhage
6. Leigh encephalopathy (gangliosidosis GM type 3)
7. Locked-in syndrome
8. Phenothiazine and benzodiazepine poisoning (combined; reverse)
9. Ruptured giant distal posterior inferior cerebellar artery aneurysm
10. Thrombosis of basilar, middle cerebral, or vertebral arteries with posterior fossa infarction


**PARALYSIS OF THIRD NERVE (OCULOMOTOR NERVE)**

This type of paralysis includes ptosis, an inability to rotate the eye upward or inward, a dilated unreactive pupil (iridoplegia), and paralysis of accommodation (cycloplegia).

**Extracted table paralysis of third nerve**

1. Intracerebral
   A. Lesion of red nucleus (Benedikt syndrome) - homolateral oculomotor paralysis with contralateral intention tremor
   B. Myasthenia Gravis and Mesencephalic Cavernous Angioma
   C. Nuclear types-pareses of a single or a few extraocular muscles supplied by the oculomotor nerve in one or both eyes; there may not be pupillary disturbances (mydriasis, sluggish pupillary reaction) and paresis of accommodation; in tumors within or near the midbrain (pinealomas), there is a combination of isolated muscle pareses with vertical gaze palsy, possibly a disturbance of convergence, and nystagmus retractorius (Parinaud syndrome, sylvian aqueduct syndrome, pineal syndrome); includes Axenfeld-Schurenberg syndrome (cyclic oculomotor paralysis), Bruns syndrome (postural change syndrome), Claude syndrome (inferior nucleus ruber syndrome), congenital vertical retraction syndrome, and Nothnagel syndrome (ophthalmoplegia-cerebellar ataxia syndrome)
D. Occlusion of basilar artery due to emboli especially but also to hemorrhage or aneurysm
E. Recurrent third nerve palsy secondary to vascular spasm of migraine
F. Syndrome of cerebral peduncle (Weber syndrome)-homolateral oculomotor paralysis and cross-hemiplegia
G. Tumors

2. Intracranial
A. Amebic dysentery
B. Aneurysm rupture at base of brain-third nerve paralysis, pain around the face (fifth nerve), and headache
C. Botulism
D. Chickenpox
E. Craniopharyngioma
F. Dengue fever
G. Devic syndrome (optical myelitis)
H. Diphtheria
I. Encephalitis, acute
J. Hepatic failure
K. Hepatitis
L. Influenza
M. Lockjaw (tetanus)
N. Lymphoma
O. Malaria
P. Measles immunization
Q. Meningococcal meningitis
R. Multiple sclerosis (disseminated sclerosis)
S. Ophthalmic migraine
T. Periarteritis nodosa
U. Poliomyelitis
V. Polyneuritis because of toxins such as alcohol, lead, arsenic, and carbon monoxide; dinitrophenol or carbon disulfide poisoning; or diabetes mellitus, herpes zoster, or mumps
W. Rabies
X. Relapsing polychondritis
Y. Smallpox vaccination
Z. Subdural hematoma
AA. Syphilis (acquired lues)
BB. Temporal arteritis syndrome (Hutchinson-Horton-Magrath-Brown syndrome)
CC. Tuberculosis

3. Lesions affecting exit from cranial cavity
A. Cavernous sinus syndrome-paralysis of third, fourth, and sixth nerves with proptosis
   (1) Aneurysm (arteriovenous fistula syndrome)
   (2) Carotid-cavernous fistula
   (3) Cavernous sinus thrombosis
(4) Extension from lateral sinus thrombosis
(5) Extension of nasopharyngeal tumor
(6) Pituitary adenoma - lateral extension
(7) Tolosa-Hunt syndrome (painful ophthalmoplegia)

B. Superior orbital fissure syndrome—same as for cavernous sinus syndrome except exophthalmos is less likely to occur and optic nerve involvement and miotic pupil are more likely
   (1) Aneurysm of internal carotid artery syndrome (foramen lacerum syndrome)
   (2) Occlusion of superior ophthalmic vein
   (3) Skull fractures or hemorrhage
   (4) Sphenoid sinus suppuration (sphenocavernous syndrome)
   (5) Temporal syndrome (Gradenigo syndrome)
   (6) Tumors, such as sphenoid ridge meningioma (Rochon-Duvigneaud syndrome), nasopharyngeal tumor, metastatic carcinoma, rhabdomyosarcoma, chordoma, and sarcoma

C. Orbital apex—involve ment of third, fourth, sixth, and first division of fifth cranial nerves and optic nerve proptosis is common

4. Other
   A. Alber-Schonberg syndrome (marble bone disease, osteopetrosis)
   B. Associated with aspirin poisoning
   C. Congenital
   D. Hodgkin disease
   E. Lupus erythematosus (Kaposi-Libman-Sacks syndrome)
   F. Myasthenia gravis (masquerade)
   G. Passow syndrome (status dysraphicus syndrome)
   H. Porphyria cutanea tarda
   I. Sarcoid (Schaumann syndrome)


Childhood Causes of Third Nerve (Oculomotor) Palsy

1. Trauma
2. Neoplasm
3. Undetermined
4. Ophthalmoplegic migraine
5. Postoperative cause
6. Meningitis/encephalitis
7. Subdural hematoma
8. Viral or post-upper respiratory tract infection
9. Varicella-zoster virus
10. Aneurysm
11. Orbital cellulitis
12. Sinus disease
13. Mesencephalic cyst
14. Cyclic oculomotor nerve palsy
15. Poison


**PARALYSIS OF FOURTH NERVE (TROCHLEAR NERVE)**

This type of paralysis produces palsy of superior oblique muscle resulting in limitation of downward movement of eye when it is in adducted position; it is frequently associated with third cranial nerve palsy.

1. Intracerebral
   A. Thrombosis of nutrient vessels, including median penetrating branch of basilar artery to fourth nucleus
   B. Hemorrhage in the roof of the midbrain
   C. Aneurysm, including direct involvement by posterior cerebral and superior cerebellar arteries
   D. Tumors (rare if isolated fourth palsy)
      (1) Primary
         a. Gliomas, such as astrocytomas, ependymomas, and medulloblastomas
         b. Other primary tumors, including meningiomas, pinealomas, craniopharyngiomas, and hemangiomas
      (2) Unilateral trochlear nerve palsy
         E. Metastatic lesions, such as those from the nasopharynx, rhabdomyosarcomas, and neuroblastomas
         F. Neonatal hypoxia
         G. Nuclear type-trochlear paresis combined with a homolateral oculomotor paresis, occasionally in association with vertical gaze palsies, convergence spasm or convergence palsy, and pupillary disturbances seen in tumors of the roof of the midbrain or pinealomas (pineal syndrome)
H. Claude syndrome (inferior nucleus ruber syndrome)
I. Passow syndrome (syringomyelia)
J. Inflammatory lesions, such as meningoencephalitis, cerebellitis, and abscess
K. Pseudotumor cerebri

2. Intracranial
   A. Aneurysms, such as that of the posterior communicating artery or foramen
      lacerum syndrome (aneurysm of internal carotid artery syndrome)
   B. Hematomas, traumatic
   C. Hydrocephalus
   D. Meningitis, encephalitis, polyneuritis-diabetes mellitus, herpes zoster, multiple
      sclerosis, myasthenia gravis, chickenpox, diphtheria, hydrophobia, Gradenigo
      syndrome, influenza, malaria, poliomyelitis
   E. Trauma
   F. Tumors, including cerebellopontine angle tumor and pituitary adenoma

3. Lesions affecting exit from cranial cavity
   A. Cavernous sinus syndrome (Foix syndrome)
   B. Superior orbital fissure syndrome (Rochon-Duvigneaud syndrome)
   C. Orbital apex syndrome (Rollet syndrome)

4. Orbital lesions
   A. Fracture of superior orbital rim
   B. Sinusitis
   C. Operations on the frontal sinus in which there is trochlear displacement
   D. Trochlear disturbance, such as in Paget disease or hypertrophic arthritis
   E. Adherence syndrome-adhesions between the superior rectus and superior
      oblique muscles
   F. Abnormal insertion of superior oblique muscle or abnormal fascial attachments
   G. Rochon-Duvigneaud syndrome (superior orbital fissure syndrome)
   H. Idiopathic

**Extracted Table Paralysis of fourth nerve**

Feinberg AS, Newman NJ. Schwannoma in patients with isolated unilateral trochlear

Holmes JM, et al. Pediatric third, fourth, and sixth nerve palsies: a population-based

Keane JR. Fourth nerve palsy: historical review and study of inpatients. *Neurology* 1993;
43:2439-2433.

Peterman SH, Newman NJ. Pituitary macroadenoma manifesting as an isolated fourth

Roy FH. *Ocular syndromes and systemic diseases*, 3rd ed. Philadelphia: Lippincott
Williams & Wilkins, 2002.

**Childhood Causes of Fourth Nerve (Trochlear) Palsy**

1. Trauma  
2. Neoplasm  
3. Undetermined  
4. Postoperative cause  
5. Meningitis  
6. Hydrocephalus  
7. Ophthalmoplegic migraine  
8. Viral infection  
9. Aneurysm  
10. Other


**PSEUDOABDUCENS PALSY**

1. Accommodative spasm  
2. Blowout fracture (medial rectus entrapment)  
3. Cross fixation (congenital esotropia)  
4. Duane syndrome (retraction syndrome)  
5. Fibrosis of medial rectus  
6. Horizontal gaze palsy (bilateral)-with or without contraction  
7. Lack of effort involved in abducting a habitually adducted eye patch on other eye differentiates from abducens palsy  
8. Myasthenia gravis  
9. Myositis  
10. Orbital cellulitis (abscess)  
11. Overambitious (large) resection of medial rectus  
12. Thyroid myopathy (Graves disease, hyperthyroidism)  
13. Unwillingness to cooperate-doll’s head phenomenon (sudden passive turning of head) differentiates from abducens palsy


Extracted Table Paralysis of sixth nerve
Extracted Table Paralysis of third, fourth, and sixth cranial nerves

PARALYSIS OF SIXTH NERVE (ABDUCENS PALSY)
This type of paralysis produces palsy of the lateral rectus muscle with esotropia increasing when the eye is moved laterally. The course of the sixth nerve makes it more vulnerable to injury than other cranial nerves.

1. Intracerebral
   A. Foville syndrome (Foville peduncular syndrome)
   B. Gaucher disease (cerebroside lipidosis)
   C. Hydrocephalus
   D. Inflammatory lesions, such as meningoencephalitis, cerebellitis, and abscess
   E. Lateral ventricular cyst
   F. Leukemia
   G. Millard-Gubler syndrome (abducens-facial hemiplegia alternans)
   H. Mycoplasma pneumoniae
   I. Nuclear aplasia - autosomal dominant
   J. Platybasia (cerebellomedullary malformation syndrome)
   K. Spontaneous subdural hematoma
   L. Thrombosis or aneurysm of nutrient vessels to sixth nucleus-basilar artery
   M. Tumors - intracranial, pontine glioma, or metastatic tumor from breast, thyroid glands, or nasopharynx
      (1) Primary
         a. Gliomas, such as astrocytomas, ependymomas, and medulloblastomas
         b. Other primary tumors, including meningiomas, pinealomas, craniopharyngiomas, and hemangiomas
      (2) Metastatic lesions, such as those from the nasopharynx, rhabdomyosarcomas, and neuroblastomas
   N. Wernicke encephalopathy-thiamine deficiency in alcoholics with sixth nerve palsy, paresis of horizontal conjugate gaze, nystagmus, ataxia, and Korsakoff psychosis

2. Intracranial
   A. Carotid artery aneurysm (foramen lacerum syndrome)
   B. Cerebellopontine angle tumor, such as acoustic neuroma, producing unilateral deafness, facial paralysis, diplopia, and papilledema
   C. Chickenpox
   D. Coccidioidomycosis
   E. Congenital absence of sixth nerve
   F. Cushing syndrome (II) (angle tumor syndrome)
   G. Dandy-Walker syndrome (atresia of the foramen Magendie)
H. Diphtheria
I. Gradenigo syndrome - osteitis of petrous tip of pyramid following homolateral mastoid or middle ear infection; facial pain (fifth nerve involvement)
J. Greig syndrome (ocular hypertelorism syndrome)
K. Hydrophobia (rabies)
L. Hydrocephalus (decreased intracranial pressure)
M. Increased intracranial pressure
N. Malaria
O. Massive pituitary adenoma
P. Measles
Q. Meningitis
R. Møbius syndrome (congenital paralysis of sixth and seventh nerves)
S. Neuritis because of diseases such as diabetes mellitus, herpes zoster, poliomyelitis, lead or arsenic poisoning, multiple sclerosis, syphilis, brucellosis
T. Ophthalmoplegic migraine syndrome
U. Osteosarcoma
V. Passow syndrome (status dysraphicus syndrome)
W. Pseudotumor cerebri (Symonds syndrome)
X. Raymond syndrome (pontine syndrome)
Y. Relapsing polychondritis
Z. Skeletal dysplasia (mental retardation, abducens palsy)-x-linked
AA. Skull fractures-usually crush injury
BB. Spontaneous dissection of the internal carotid artery
CC. Subdural hematoma
DD. Trichinellosis
EE. Tumor extension as chordoma
FF. Vascular lesions, because of congenital aneurysm, arteriovenous fistulas, diabetes, hypertension
GG. Water-soluble contrast myelography

3. Lesions affecting exit of sixth nerve from cranial cavity
   A. Cavernous sinus syndrome (Foix syndrome)
   B. Le Fort I maxillary osteotomy
   C. Optic nerve sheath fenestration
   D. Orbital apex lesion
   E. Percutaneous thermal ablation of trigeminal nerve rootlet
   F. Sphenocavernous syndrome
   G. Sphenopalatine fossa lesion-loss of tearing and paresis of second division of fifth nerve, most frequently because of malignant tumor
   H. Superior orbital fissure syndrome
   I. Tolosa-Hunt syndrome (painful ophthalmoplegia)
   J. Transient in newborns

4. Other
   A. Cluster headache
   B. Cretinism (hypothyroid goiter)
   C. Duane syndrome (retraction syndrome)
   D. Engelmann syndrome (hereditary multiple diaphyseal sclerosis)
E. Following lumbar puncture, lumbar anesthesia, or Pantopaque injection for myelography
F. Kahler disease (multiple myeloma)
G. Lupus erythematosus (Kaposi-Libman-Sacks syndrome)
H. Myasthenia gravis
I. Optic nerve sheath fenestration (rare)
J. Preeclampsia
K. Sarcoidosis
L. Secondary to immunization or viral illness
M. Toxic substances, such as arsenic, carbon tetrachloride, dichloroacetylene, Dilantin, gold salts, isoniazid, nitrofurane, thalidomide, trichloroethylene, furaltadone (Altafur), lithium


**Childhood Causes of Sixth Nerve (Abducans) Palsy**

1. Trauma
2. Neoplasm
3. Undetermined
4. Viral/benign
5. Gradenigo syndrome
6. Meningitis/encephalitis
7. Pseudotumor cerebri
8. Leukemia
9. Hydrocephalus
10. Arteriovenous malformation, brain
11. Multiple sclerosis
12. Miscellaneous


**Childhood Causes of Third, Fourth, and Sixth Nerve Palsy**

1. Trauma
2. Neoplasm
3. Undetermined
4. Postoperative cause
5. Meningitis
6. Hydrocephalus
7. Ophthalmoplegic migraine
8. Viral infection
9. Aneurysm
10. Other


**ACUTE OPHTHALMOPLEGIA (ACUTE ONSET OF EXTRAOCULAR MUSCLE PALSY)**

1. Infranuclear
   A. Aneurysm of internal carotid artery or circle of Willis
   B. Trauma
      (1) Orbital fracture
      (2) Orbital hematoma
   C. Orbital cellulitis secondary to acute paranasal sinusitis including mucormycosis in a diabetic
   D. Ophthalmoplegic migraine
   E. Myasthenia gravis
   F. Orbital pseudotumor
   G. Orbital tumors
      (1) Lymphoma
      (2) Metastatic
      (3) Rhabdomyosarcoma
2. Nuclear
   A. Acute and subacute infections
      (1) Infectious encephalitis
         a. Viral encephalitis
            (i) Anterior poliomyelitis
            (ii) Encephalitis lethargica and other epidemic viral encephalitides
(iii) Fisher syndrome (ophthalmoplegia, ataxia, areflexia)
(iv) Rabies
(v) Vaccinal encephalitis
(vi) Varicella, variola, measles, mumps, influenza, infectious mononucleosis
(vii) Zoster

b. Organismal encephalitic infections
   (i) Typhoid
   (ii) Scarlet fever
   (iii) Whooping cough
   (iv) Gas gangrene
   (v) Septicemia
   (vi) Pneumonia
   (vii) Typhus
   (viii) Malaria

c. Acute central nervous system diseases
   (i) Acute demyelinating diseases-acute disseminated encephalomyelitis, acute multiple sclerosis
   (ii) Neuritic infections
       (a) Polyradiculoneuritis
       (b) Epidemic paralyzing vertigo
       (c) Acute infectious (rheumatic) polynueritis
       (d) Interstitial neuritis-meningitis, cranial sinusitis, petrositis, nasal sinusitis, orbital periostitis, orbital abscess
   (iii) Widespread infections
       (a) Meningovascular syphilis
       (b) Mucormycosis (diabetes, immunosuppressed, AIDS)
       (c) Tuberculosis
       (d) Torula and cryptococcosis
   (iv) Toxic conditions
       (a) Diphtheria
       (b) Tetanus
       (c) Botulism
   (v) Allergic conditions
       (a) Sarcoidosis syndrome (Schaumann syndrome)
       (b) Recurrent multiple cranial nerve palsies

B. Metabolic diseases
   (1) Deficiency diseases
       a. Thiamine deficiency (Wernicke- Korsakoff syndrome)
       b. Nicotinic acid deficiency-pellagra
       c. Ascorbic acid deficiency-scurvy
   (2) Diabetes
   (3) Anemias
       a. Primary anemia-leukemia
b. Secondary anemia (loss of blood)

(4) Exophthalmic ophthalmoplegia
(5) Porphyria

C. Poisoning such as lead, carbon monoxide, snake poisons, wasp stings, ergot, sulfuric acid, phosphorus, triorthocresylphosphate, and dichloroacetylene

D. Drugs, including the following:

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<th>Drug B</th>
<th>Drug C</th>
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E. Neoplasms and cysts
F. Trauma affecting the midbrain, base of the skull, and orbit
G. Vascular lesions as arteriosclerosis, hemorrhage and thrombosis in the midbrain, subarachnoid, hemorrhage, aneurysms, congenitally dilated arteries, giant-cell arteritis
H. Idiopathic-etiologic basis undetermined


**CHRONIC OPHTHALMOPLEGIA (SLOW ONSET OF EXTRAOCULAR MUSCLE PALSY)**

1. Degenerative conditions
   A. Amyotrophic lateral sclerosis-progressive bulbar palsy
   B. Chronic progressive external ophthalmoplegia
   C. Hereditary ataxias-Friedreich ataxia, Sanger-Brown ataxia
   D. Progressive supranuclear palsy
   E. Syringomyelia (syringobulbia)
   F. Thyroid myopathy (Graves disease)

2. Infective conditions
   A. Diffuse sclerosis
   B. Disseminated sclerosis (multiple sclerosis)
   C. Syphilis


**BILATERAL COMPLETE OPHTHALMOPLEGIA (BILATERAL PALSY OF OCULAR MUSCLES, PTOSIS, WITH PUPIL AND ACCOMMODATION INVOLVEMENT)**

1. Arteriosclerotic hemorrhage and occlusion
2. Cerebellopontine angle tumors (Cushing syndrome II)
3. Encephalitis, acute
4. Fisher syndrome (ophthalmoplegia-ataxia areflexia syndrome)
5. Giant-cell arteritis (Hutchinson-Horton-Magath-Brown syndrome)
6. Kiloh-Nevin syndrome (ocular myeomyopathy)
7. Midbrain tumors
8. Multiple sclerosis (rare)
9. Mucormycosis
10. Ohasha syndrome (ophthalmoplegia, hypotonia, ataxia hypacusis, athetosis)
11. Orbital abscess
12. Parinaud syndrome (conjunctiva-adenitis syndrome)
13. Retrobulbar block complication
14. Rochon-Duvigneaud syndrome (superior orbital fissure syndrome)
15. Rollet syndrome (orbital apex-sphenoidal syndrome)
16. Syphilis (acquired lues)
17. Trauma
18. Wernicke encephalopathies (thiamine deficiency)
19. Whipple disease (intestinal lipodystrophy)


**EXTERNAL OPHTHALMOPLEGIA (PARALYSIS OF OCULAR MUSCLES INCLUDING PTOSIS WITH SPARING OF PUPIL AND ACCOMMODATION)**

1. Abiotrophy-specific for one particular tissue, bilateral, symmetric
2. Amyloidosis (Lubarsch-Pick syndrome)
3. Aneurysm of internal carotid artery (foramen lacerum syndrome)
4. Bassen-Kornzweig syndrome (familial hypolipoproteinemia)
5. Bee sting
6. Chronic progressive external ophthalmoplegia
7. Congenital and familial
8. Diabetes mellitus (Willis disease)
9. Diphtheria
10. Epidemic encephalitis
11. Friedreich ataxia
12. Garcin syndrome (Schmincke tumor unilateral cranial paralysis)
13. Graves disease (hyperthyroidism)
14. Jacod syndrome (petrosphenoidal space syndrome)
15. Kearns-Sayre syndrome (ophthalmoplegic retinal degeneration syndrome)
16. Mumps
17. Myasthenia gravis (Erb-Goldflam syndrome)
18. Myotonic dystrophy (Curschmann-Steinert syndrome)
19. Myositis
20. Nevus sebaceous of Jadassohn
21. Nothnagel syndrome (ophthalmoplegia-cerebellar ataxia syndrome)
22. Oculopharyngeal syndrome (progressive muscular dystrophy with ptosis and dysphagia)
23. Olivopontocerebellar atrophy III (with retinal degeneration)-dominant
24. Ophthalmoplegia, progressive external, and scoliosis (horizontal gaze paralysis, familial)-recessive
25. Pernicious anemia
26. Polyradiculoneuritis (Guillain-Barré and Fisher syndromes)
27. Progressive facial hemiatrophy (Parry-Romberg syndrome)
28. Pseudotumor (orbital)
29. Refsum syndrome (heredopathia atactica polyneuritiformis syndrome)
30. Scleroderma (progressive systemic sclerosis)
31. Shy-Drager syndrome (orthostatic hypotension syndrome)
32. Shy-Gonatas syndrome (accumulation of lipids in muscles simulates gargoylism)
33. Tick paralysis (Lyme disease, Rocky Mountain spotted fever)
34. Vincristine—may have fifth and seventh nerve and peripheral neuropathies
35. Wernicke encephalopathies (beriberi, thiamine deficiency)


**INTERNUCLEAR OPHTHALMOPLEGIA**

This condition comprises paralysis of the medial rectus muscles on attempted conjugate lateral gaze without other evidence of third nerve paralysis due to involvement of medial longitudinal fasciculus. Jerk nystagmus of abducting eye and vertical nystagmus, usually on upward gaze, may be present.

1. Bilateral
   A. Arnold-Chiari malformation (cerebellomedullary malformation syndrome)
   B. "Crack" cocaine
   C. Fabry disease (glycosphingolipid lipidosis)
   D. Inflammation, such as upper respiratory infection
   E. Midbrain infarction
   *F. Multiple sclerosis (disseminated sclerosis)
   G. Myasthenia gravis (Erb-Goldflam syndrome)
   H. Occlusive vascular disease
   I. Oculocerebellar tegmental syndrome
   J. Pontine hematoma
   K. Syphilis (acquired lues)
   L. Temporal arteritis
   M. Vertebral basilar artery syndrome (whiplash injury)
   N. Webino syndrome (wall-eyed exotropia bilateral internuclear ophthalmoplegia)
   O. Wernicke encephalopathy

2. Unilateral
   A. Bielschowsky-Lutz-Cogan syndrome (internuclear ophthalmoplegia)
   B. Cryptococcosis (torulosis)
C. Multiple sclerosis (disseminated sclerosis)
D. Myasthenia gravis (Erb-Goldflam syndrome)
E. Neuro-Behçet Disease
F. Tumors of the brainstem
*G. Vascular lesion-infarct of small branch of basilar artery


**PAINFUL OPHTHALMOPLEGIA (PALSY OF OCULAR MUSCLES WITH PAIN)**

1. Adenocarcinoma metastatic to the orbit
2. Atypical facial neuralgia
3. Cavernous sinus syndrome (Foix syndrome)
4. Collier sphenoidal palsy
5. Diabetic ophthalmoplegia
6. Intracavernous carotid aneurysm
7. Myositis (orbital)
8. Nasopharyngeal tumor
9. Ophthalmoplegic migraine
10. Orbital abscess (mucormycosis-diabetes, immunosuppressed, AIDS)
11. Orbital apex sphenoidal syndrome (Rollet syndrome)
12. Orbital periostitis
13. Postherpetic neuralgia
14. Pseudotumor of orbit
15. Superior orbital fissure syndrome (Rochon-Duvigneaud syndrome, including superior orbital fissuritis)
16. Temporal arteritis
17. Tic douloureux of the first trigeminal division
18. Tolosa-Hunt syndrome (inflammatory lesion of cavernous sinus)


**Extracted Table internuclear ophthalmoplegia**

**Extracted Table Painful Ophthalmoplegia**

**TRANSIENT OPHTHALMOPLEGIA (EXTRAOCULAR MUSCLE PARALYSIS OF SHORT DURATION)**

1. Cranial irradiation and intrathecal chemotherapy
2. Cyclic esotropia
3. Cyclic oculomotor palsy
4. Following internal carotid artery ligation for treatment of intracavernous giant aneurysm
5. Lethargic encephalitis
6. Multiple sclerosis (disseminated sclerosis-usually the lateral rectus)
7. Myasthenia gravis (ocular, early)
8. Oculomotor nuclear complex infarction
9. Ophthalmoplegia migraine
10. Post lumbar puncture abducens palsy
11. Syphilis (acquired lues)
12. Tabes dorsalis
13. Treatment of arteriovenous fistulas with Debrun balloon technique
14. Wilson disease (hepatolenticular degeneration)


**PAINFUL OCULAR MOVEMENTS (PAIN WITH MOVEMENT OF THE EYES)**

1. Bone-break fever (dengue fever) (rare)
2. Influenza
3. Myositis
   A. "Collagen diseases"
   B. Infectious myositis
   C. Trichinosis
4. Orbital cellulitis
5. Orbital periostitis
6. Retrobulbar neuritis


**POOR CONVERGENCE (INABILITY OF BOTH EYES TO FIXATE SIMULTANEOUSLY ON A NEAR OBJECT)**

1. Functional
   A. Convergence insufficiency
   B. Exophoria
   C. Exotropia
   D. Hysteria
   E. Poor attention span

2. Organic
   A. Brain lesion, to include bilateral occipital lobe lesions, superior colliculi, and anterior internuclear ophthalmoplegia, such as in hemorrhage, trauma, or tumors
   B. Dorsal midbrain syndrome
   C. Encephalitis
   D. Exophthalmic goiter-Möbius sign
   E. Exophthalmos
   F. Multiple sclerosis
   G. Myotonic dystrophy
   H. Narcolepsy
   I. Poor visual acuity in one or both eyes
   J. Postencephalitis
   K. Syphilis and tabes
   L. Third nerve paralysis (see p. 153)
   M. Whiplash injury

3. Drugs, including the following:

<table>
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<tr>
<th>Alcohol</th>
<th>Cyclophosphamide</th>
<th>Morphine</th>
</tr>
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<td>Allobarbital</td>
<td>Dextroamphetamine</td>
<td>Opium</td>
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<tr>
<td>Amobarbital</td>
<td>Dimethyl tubocurarine</td>
<td>Penicillamine</td>
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<tr>
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<td>Iodide</td>
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<tr>
<td>Cyclophosphamide</td>
<td>Metocurine iodide</td>
<td></td>
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</tbody>
</table>
SPASM OF CONVERGENCE

Spasm of convergence occurs with spasm of accommodation and miosis (i.e., spasm of the near reflex).

1. Encephalitis-accompanied with nystagmus
2. Hysteria-may be confused with lateral rectus palsy
3. Labyrinthine fistulas
4. Kenny syndrome
5. Oculogyric crisis in myasthenia gravis
6. Paralysis of horizontal gaze with compensatory spasm of near reflex
7. Parinaud syndrome (divergence paralysis)
8. Tabes dorsalis
9. Trauma
10. Wernicke syndrome (avitaminosis B\textsubscript{1})


DIVERGENCE PARALYSIS

Divergence paralysis is defined as having a supranuclear cause with sudden onset of comitant esotropia and uncrossed diplopia at distance, fusion at near (usually 1-2 m) normal ductions and versions, and gross impairment of fusional amplitudes of divergence.

*1. Brainstem lesions
   A. Cerebellar cyst
   B. Hemangioma
   C. Tumors, such as cerebellar and acoustic neuromas and pontine glioma
2. Cerebral hemorrhage
3. Diazepam
4. Diphtheria
5. Dorsal midbrain syndrome
6. Epidemic encephalitis
7. Functional
8. Head injuries
9. Increased intracranial pressure
10. Influenza
11. Lead poisoning  
12. Multiple sclerosis (disseminated sclerosis)  
13. Poliomyelitis  
14. Syphilis  
15. Unknown  
16. Vascular disease  
   A. Diabetes mellitus  
   B. Hypertension  
   C. Occlusion of subclavian artery with reversal of flow in vertebral artery  
   D. Vertebral basilar insufficiency


**OCULOCARDIAC REFLEX**
Bradycardia, nausea, and faintness occur and are dependent on trigeminal sensory stimulation evoked by pressure on or within the eyeball or from sensory impulses by stretching of ocular muscles.

1. Acute glaucoma  
2. Anophthalmic socket  
3. During ophthalmoscopy examination of premature infants  
4. Exaggerated in epidemic encephalitis  
5. Intermittent exophthalmos due to congenital venous malformations of the orbit  
6. Intraocular injections  
7. Orbital hematoma  
   *8. Pressure on globe  
9. Retinal detachment operation  
10. Severe injury to eye or orbit  
   *11. Traction on extraocular muscles including levator palpebrae superioris


RETRACTION OF THE GLOBE (ON HORIZONTAL CONJUGATE GAZE)

*1. Duane syndrome (retraction syndrome)-cocontraction of horizontal rectus muscles, lateral rectus, and both vertical muscles, or medial and inferior rectus muscles or fibrotic lateral rectus
   A. Acrorenoocular syndrome
   B. Goldenhar syndrome
   C. Hanhart syndrome
   D. Isolated
   E. Okihiro syndrome
   F. Wildervanck syndrome (Klippel-Feil anomaly with Duane syndrome)

2. Fibrosis secondary to strabismus surgery

3. Medial wall fracture with incarceration of orbit contents-retraction of globe with attempted abduction

4. Orbital mass
   A. Dermoid cyst
   B. Hemangioma
   C. Lymphangioma
   D. Osteofibroma

5. Retraction of convergent nonfixating eye associated with loss of conjugate lateral gaze and occurrence of the near reflex on attempted lateral gaze

6. Thyroid myopathy


FORCED DUCTION TEST
The eyeball is moved away from the muscle being tested; this is accomplished by grasping with a forceps the conjunctiva and episclera close to the limbus.

1. Supraduction - infraduction
   A. Resistance
      (1) Abnormal fascial or muscle attachments
      (2) Congenital fibrosis syndrome
      (3) Double elevator palsy
      (4) Orbital floor fracture
      (5) Orbital mass
      (6) Thyroid myopathy of inferior rectus muscle
   B. Unrestricted
      (1) Elevator paresis
(2) Paresis of inferior superior rectus muscle

2. Supraduction in adduction
   A. Brown superior oblique tendon sheath syndrome-resistance (see p. 132)
   B. Paresis of inferior oblique muscle-unrestricted

3. Adduction
   A. Resistance
      (1) Chronic third nerve palsy with contracture lateral rectus
      (2) Congenital fibrosis syndrome
      (3) Duane retraction syndrome because of fibrosis of lateral rectus muscle
      (4) Orbital mass
      (5) Tight lateral rectus following excessive resection operation
      (6) Thyroid myopathy
   B. Unrestricted
      (1) Extensive medial rectus recession
      (2) Duane retraction syndrome because of central or peripheral cocontraction of medial and lateral rectus on attempted adduction

4. Abduction
   A. Resistance
      (1) Abnormal fascial or muscle attachments including strabismus fixus
      (2) Blowout fracture
      (3) Chronic sixth nerve palsy with contracture medial rectus
      (4) Myositis
      (5) Orbital mass
      (6) Thyroid myopathy
   B. Unrestricted
      (1) Extensive lateral rectus recession
      (2) Paralysis of lateral rectus muscle


DOUBLE ELEVATOR PALSY (APPARENT PARALYSIS OF GLOBE ELEVATORS [I.E., SUPERIOR RECTUS AND INFERIOR OBLIQUE MUSCLES])

1. Inferior rectus muscle restriction
   A. Anomalous insertion of inferior rectus muscle
   B. Blow-out fracture
   C. Congenital orbital fibrosis syndrome
   D. Thyroid ophthalmopathy

2. Monocular elevation paresis secondary to central nervous system lesions
   A. Monocular elevation paresis
   B. Skew deviation

3. Neurogenic or myogenic superior rectus muscle weakness
   A. Myasthenia gravis (Erb-Goldflam syndrome)
   B. Postoperative Berke-Motais surgery
C. Superior oblique tendon sheath syndrome (see p. 132)
D. Third cranial nerve palsy (see p. 153)
E. Trauma


**OCULAR NEUROMYOTONIA (PAROXYSMAL MONOCULAR DEVIATIONS ASCRIBABLE TO INVOLUNTARY CONTRACTION OF MUSCLES INNERVATED BY THIRD, FOURTH, OR SIXTH CRANIAL NERVES)**

1. Aneurysmal compression of third nerve
2. Chondrosarcoma
3. Cystic craniopharyngioma
4. Following radiation therapy
5. Rhabdomyosarcoma


**EXTRAOCULAR MUSCLE ENLARGEMENT ON ORBITAL COMPUTERIZED AND TOMOGRAPHIC SCAN**

1. Diffuse
   A. Acromegaly
   B. Amyloidosis
   *C. Graves disease as thyroid ophthalmopathy
   *D. Infection
   E. Parasitic infiltration
   F. Trauma
   G. Tumors, including pseudotumor
   H. Vascular abnormalities as arteriovenous fistula
   I. Collagen vascular disease
2. Focal
   A. Cysticercosis
   B. Hemorrhagic cyst
   C. Primary or metastatic carcinoma
   D. Trichinella


CELLULAR RESPONSES

1. Basophilic reaction—significant only when seen in large numbers; immunoglobulin E (Ig-E) mediated allergic conditions; key role in asthma, atopy, and hypersensitivity responses; present in vernal (eosinophils) and giant papillary conjunctivitis (GPC) conjunctivitis.

2. Eosinophilic reaction-GPC (giant papillary conjunctivitis); Charcot-Leyden crystals are more prominent than intact eosinophils in chronic allergy; parasitic conjunctivitis.

   *A. Vernal conjunctivitis—characteristic with fragmentation of eosinophil
B. Hay fever conjunctivitis—rarely fragmentation of eosinophil
C. Allergic conjunctivitis from various drugs, cosmetics, and other antigens
D. Atropine sensitivity—not present when eserine or pilocarpine is used

3. Mononuclear reaction
   A. Viral disease-100% without secondary infection; usually lymphocytic
      *(1) Epidemic keratoconjunctivitis—adenovirus type 8
      *(2) Pharyngoconjunctival fever—adenovirus type 3
      *(3) Herpetic keratoconjunctivitis
      *(4) Acute follicular conjunctivitis of Beal

B. Chronic ocular infections

4. Neutrophilic reaction—early stage of severe viral conjunctivitis
   A. All bacteria but two—Neisseria catarrhalis and Haemophilus duplex (Morax-Axenfeld diplobacillus)
   B. Viruses of the family Chlamydiaceae (trachoma inclusion conjunctivitis (TRIC) agent)
      *(1) Trachoma
      *(2) Inclusion conjunctivitis
      *(3) Lymphogranuloma venereum
   C. Fungal disease
      *(1) Streptothrix conjunctivitis secondary to canaliculitis
      *(2) Nocardial corneal ulcers
      *(3) Monilial corneal ulcers
   D. Unknown cause
      *(1) Erythema multiforme (Stevens-Johnson syndrome)
      *(2) Conjunctivitis of Reiter disease

E. Vernal conjunctivitis—eosinophilic and neutrophilic reaction

F. Epidemic keratoconjunctivitis and herpetic keratoconjunctivitis have a shift from mononuclear to polymorphonuclear reaction when a membrane is formed

5. Plasma-cell reaction—trachoma—especially with spontaneous rupturing of follicles; chlamydial conjunctivitis

6. Epithelial changes
   A. Keratinization of conjunctival epithelial cells
      *(1) Alkali bum
      *(2) Vitamin A deficiency
      *(3) Exposure
      *(4) Cicatrization (such as pemphigoid and Stevens-Johnson syndrome)
      *(5) Keratoconjunctivitis sicca—partially keratinized epithelial cells, specific
      *(6) Epithelial plaque
      *(7) Superior limbic keratoconjunctivitis (SLK)
   B. Large, multipointed epithelial cells
      *(1) Characteristic of viral infection
      *(2) Most often found in herpetic keratitis
   C. Intracellular granules
      *(1) Pseudoinclusion bodies—extension of nuclear material into cytoplasm
      *(2) Intracellular-free green pigment in cytoplasm—present in persons with dark complexion
3. Intracellular-free blue granules—present in cytoplasm in about 12% of normal individuals
4. Sex chromatin—present in nuclei of females only

7. Cellular inclusions
   A. Trachoma and inclusion conjunctivitis have identical inclusions—basophilic, cytoplasmic (Halberstaedter-Prowazek)
   *B. Molluscum contagiosum—eosinophilic, cytoplasmic (Henderson/Patterson)
   C. Lymphogranuloma venereum—eosinophilic
   *D. Herpes simplex and herpes zoster—eosinophilic, internuclear (Lipschütz)
   E. Measles—multinucleated giant cells with eosinophilic internuclear inclusion bodies and cytoplasmic, eosinophilic masses
   *F. Chickenpox—eosinophilic, internuclear
   G. Smallpox—eosinophilic, cytoplasmic


**PURULENT CONJUNCTIVITIS**

Purulent conjunctivitis is characterized as violent acute conjunctival inflammation, great swelling of lids, copious secretion of pus, and a marked tendency to corneal involvement and even possible loss of the eye.

1. Gram-positive group
   A. Bacillus of Doderlein (Lactobacillus sp.)
   B. Listeria monocytogenes
   *C. Pneumococcus
   D. Staphylococcus
   *E. Streptococcus

2. Gram-negative group
   A. Aerobacter aerogenes
   B. Enterobacteriaceae
   C. Escherichia coli
   *D. Haemophilus influenzae biotype III
   E. Klebsiella pneumoniae
   F. Moraxella lacunata
   *G. Neisseria gonorrhoeae
   H. Neisseria meningitidis
   I. Proteus species
J. *Pseudomonas* species
K. *Serratia marcescens*

3. Vaccinia virus

4. Fungus
   A. *Actinomyces* species
   B. *Candida* species
   *C. Nocardia* species

5. Wiskott - Aldrich syndrome- x-linked


**ACUTE MUCOPURULENT CONJUNCTIVITIS**

This type of conjunctivitis is epidemic pink eye, marked hyperemia and a mucopurulent discharge, which tends toward spontaneous recovery.

1. Gram-positive group
   *A. Pneumococcus*
   B. Staphylococcus-eyelid lesions and punctate staining of the lower cornea may occur

2. Gram-negative group
   *A. *Haemophilus aegyptius* (Koch-Weeks bacillus)*
   B. *H. influenzae*

3. Associated with exanthems and viral infections
   A. German measles (Greig syndrome)
   B. Measles (rubeola)
   C. Mumps
   *D. Reiter syndrome (conjunctivourethrosynovial syndrome)*
   E. Scarlet fever

4. Fungus
   A. *Candida albicans*
   B. Leptothrix

5. Lyell disease-toxic epidermal necrolysis or scalded-skin syndrome

6. Relapsing polychondritis

7. Sjögren syndrome (secretoinhibitor syndrome)

8. Etiology obscure in many cases


CHRONIC MUCOPURULENT CONJUNCTIVITIS (MUCOPURULENT DISCHARGE, MODERATE HYPEREMIA WITH A CHRONIC COURSE)

1. Infective element-lids or lacrimal apparatus
   A. Monilia species
   B. Morax-Axenfeld diplobacillus (angular conjunctivitis)
   *C. Pneumococcus
   D. Pubic lice
   E. Staphylococcus
   F. Streptothrix foersteri
2. Allergic-cosmetic
3. Irritative
   A. Associated infections or irritation of lids, lacrimal apparatus, nose, or skin
   B. Deficiency of lacrimal secretions
   C. Direct irritants-foreign body, mascara, dust, wind, smog, insecticides, chlorinated water, and many others
   D. Exposure-ectropion, facial paralysis, exophthalmos, and others
   E. Eyestrain
   F. Metabolic conditions-gout, alcoholism, or prolonged digestive disturbances
   G. Overtreatment by drugs-antibiotics, miotics, mydriatics


MEMBRANOUS CONJUNCTIVITIS

Exudate permeates epithelium to such an extent that removal of membrane is difficult and a raw bleeding surface results. Membranous conjunctivitis can lead to symblepharon, ankyloblepharon, and entropion with trichiasis.

1. Chemical irritants
   A. Acids, such as acetic or lactic
   *B. Alkalis, such as ammonia or lime
C. Metallic salts, such as silver nitrate or copper sulfate

2. *Corynebacterium diphtheriae*

3. Ligneous conjunctivitis-chronic, cause unknown

4. Pneumococcus

5. Streptococcus


**PESEOMEMBRANOUS CONJUNCTIVITIS**

In pseudomembranous conjunctivitis, the fibrin network is easily peeled off, leaving the conjunctiva intact; it forms on the conjunctiva.

1. Bacteria
   
   A. *C. diphtheriae*
   
   *B. Gonococcus*
   
   *C. Meningococcus*
   
   D. Pneumococcus
   
   E. Staphylococcus
   
   *F. Streptococcus*
   

2. Viral
   
   *A. Epidemic keratoconjunctivitis (type adenovirus)*
   
   *B. Herpes simplex*
   
   C. Herpes zoster
   
   D. Reiter syndrome (conjunctivourethrosynovial syndrome)
   
   E. Vaccina

3. Fungal- *C. albicans*

*4. Allergic- vernal conjunctivitis*

5. Toxic
   
   *A. Stevens-Johnson syndrome can be caused by drugs, including the following:

   - acetaminophen
   - acetanilid
   - acetazolamide
   - acetohexamide
   - acetonphenazine
   - allobarbital
   - allopurinol

   - amidone
   - aminosalicylate(?)
   - aminosalicylic acid(?)
   - amithiozone
   - amobarbital
   - amodiaquine
   - amoxicillin

   - ampicillin
   - antipyrine
   - aprobarbital
   - aspirin
   - auranofin
   - aurothioglucose
   - aurothioglycanide
barbital  clindamycin  methdilazine
belladonna  cloxacillin  methicilllin
bendroflumethiazide  cyclobarbital  methitural
benzathine penicillin G  cyclopentobarbital  methohexital
benzthiazide  cyclothiazide  methotrimethoprim
bromide  danazol  methsuximide
bromisovalum  demeclocycline  methyclothiazide
demeclocycline  dichlorphenamid  methylphenidate
butabarbital  dicloxacillin  metolazone
butalbital  diethazine  minocycline
dutallylonal  diphenylhydantoin  minoxidil
butaperazine  doxycycline  moxalactam
butethal  enalapril  nafcinil
butamazine  erythromycin  naproxen
captopril  ethopropazine  oxacillin
carbamazepine  ethosuximide  oxyphenbutazone
carbenicillin  ethotoin  oxytetracycline
carbromal  ethoxzolamide  paramethadione
carisoprodol  fenoprofen  penicillin
carphenazine  fluphenazine  pentobarbital
cefaclor  furosemide  perazine
cefadroxil  gentamicin  pericyazine
cefamandole  glyburide  perphenazine
cefazolin  heptabarbital  phenacetin
cefonicid  gold Au 198  phenobarbital
cefoperazone  gold sodium thiomalate  phenoxymethyl
ceforanide  gold sodium thiosulfate  penicillin
cefotaxime  heptabarbital  phensuximide
cefotetan  hetacillin  phenylbutazone
cefoxitin  hexethal  phenytoin
cefsulodin  hexobarbital  piperacetazine
ceftazidime  icubrofen  piroxicam
ceftizoxime  hydrochlorothiazide  probarbital
ceftiraxone  hydroflumethiazide  prochloperazine
cefuroxime  hydroxychloroquine  promazine
cephalexin  ibuprofen  promethazine
ccephalosporin  indapamide  propracaine
ccephalothin  indomethacin  propiomazine
cephaolin  isoniazid  ""
B. Benign mucous membrane pemphigoid can be caused by drugs, including the following:

- carbamazepine
- carbimazole
- diphenylhydantoin
- ethosuximide
- griseofulvin
- hydralazine

C. Lyell disease (toxic epidermal necrolysis or scalded-skin syndrome) can be caused by drugs, including the following:

- acetaminophen
- acetylsalicylic acid
- ampicillin
- amoxicillin
- aurothioglucose
- aurothioglucianide
- barbital
- bendroflumethiazide
- benzathine penicillin G
ethoxzolamide
fludrocortisone
fluprednisolone
gold Au 198
gold sodium thiomolate
heptabarbital
hetacillin
hexethal
hexobarbital
hydrabamine
hydrocortisone
ibuprofen
indapamide
indomethacin
isoniazid
kanamycin
mechloremethamine
melphalan
mephenytoin
mephobarbital
meprednisone
methacycline
metharbital
methazolamide
methicillin
methitural
methotrexate
methyclothiazide
methylprednisolone
metolazone
minocycline
nafcillin
nitrofurantoin
oxacillin
oxyphenbutazone
oxytetracycline
paramethadione
paramethasone
penicillamine
penicillin
phenylbutazone
phenobarbital
phenoxymethyl
potassium penicillin G
potassium penicillin V
potassium phenethicillin
prednisolone
primidone
probartal
procarbazine
quinethazone
secobarbital
smallpox vaccine
sodium salicylate
streptomycin

D. Pemphigus vulgaris
E. Hereditary epidermolysis bullosa

6. Chemical irritants
   A. Acids, such as acetic or lactic
   *B. Alkalis, such as ammonia or lime
   C. Metallic salts, such as silver nitrate or copper sulfate
   D. Vegetable and animal irritants

7. Acute graft-versus-host disease
8. Foot-and-mouth disease
9. Koch-Weeks bacillus
10. Ligneous conjunctivitis-chronic, cause unknown
11. Lipoid proteinosis (Urbach-Wiethe disease)
   *12. Superior limbic keratoconjunctivitis
13. Traumatic or operative healing of wounds
14. Wegner granulomatosis


**OPHTHALMIA NEONATORUM (CONJUNCTIVITIS OCCURRING IN NEWBORNS)**

* 1. Chemical conjunctivitis, such as from silver nitrate instillation
* 2. Chlamydial trachomatis
3. Bacteria
   A. Gram positive
      (1) *C. diphtheriae*
      (2) *Staphylococcus aureus*
      (3) *Staphylococcus epidermidis*
      (4) Streptococcus group D
      (5) *Streptococcus pneumoniae*
      (6) *Streptococcus viridans*
   B. Gram negative
      (1) Coliform bacillus, such as *E. coli*
      (2) *Enterobacter cloacae*
      (3) *Haemophilus influenzae*
      (4) *Haemophilus parainfluenzae*
      (5) *K. pneumoniae*
      (6) Meningococcus
      (7) Mima polymorpha - gram negative
      (8) *N. gonorrhoeae and N. catarrhalis*
      (9) *Neisseria* organisms
      (10) Pneumonococcus
      (11) *Proteus mirabilis*
      (12) *P. aeruginosa*
      (13) *Pseudomonas pyocyanea*
      (14) *S. marcescens*

4. Virus
   A. Herpes simplex
   B. *Streptococcus viridans*
   C. Coxsackie A
   D. TRIC virus

5. Other
   A. *Acinetobacter* species
B. *Branhamella catarrhalis*
C. *C. albicans*
D. *Citrobacter feundi*
E. *Clostridium perfringens*
F. Inclusion blennorrhea
G. *Listeriosis (L. monocytogenes)*
H. *Moraxella* species
I. *Mycoplasma* organisms
J. *Peptococcus prevotii*
K. *Propionibacterium* species
L. *Trichomonas vaginalis*


**ACUTE FOLLICULAR CONJUNCTIVITIS LYMPHOID FOLLICLES (COBBLESTONING) OF conjunctiva with RAPID ONSET**

*1. Inclusion conjunctivitis-adult inclusion conjunctivitis (AIC) (begins 2 days after exposure to organism, may be bilateral, no systemic symptoms, and a unilateral or bilateral preauricular node is often present.
*2. Adenovirus conjunctivitis-EKC has been reported worldwide from virus serotypes (the most common are 8, 11, and 19); pharyngoconjunctival fever (PCF) is usually caused by serotypes 3, 4, and 7.
  A. Pharyngoconjunctival fever-usually because of type adenovirus; common in swimming-pool epidemics in the summer and fall
  B. Epidemic keratoconjunctivitis because of adenovirus type (rarely occurs in children)
*3. Primary herpetic keratoconjunctivitis-conjunctival reaction may be follicular or pseudomembranous
4. Newcastle disease (fowlpox) conjunctivitis-usually seen in poultry handlers, veterinarians (caused by a paramyxovirus: single-stranded RNA virus that causes respiratory infections)
5. Influenza virus A
6. Herpes zoster
*7. Cat-scratch fever (Parinaud oculoglandular syndrome)-fever caused by two types of rickettsia: *Rochalimaea henselae* and *Afipia felis*
8. Echovirus keratoconjunctivitis
9. Trachoma (sometimes)
10. Bacterial *Streptococcus, Moraxella, and Treponema* organisms
11. Mesantoin use
12. *Chlamydia epizootic* (feline pneumonitis)
13. Ophthalmomyiasis
14. Acute hemorrhagic conjunctivitis
15. Neonatal inclusion conjunctivitis
16. Unknown types—a case that resists etiologic classification is encountered occasionally; it is probable that other viruses occasionally produce acute follicular conjunctivitis
17. Associated with regional adenitis
   A. Angelucci syndrome (critical allergic conjunctivitis syndrome)
   B. Anoxic overwear syndrome
   C. Benjamin-Allen syndrome (brachial arch syndrome)
   D. Floppy eyelid syndrome
   *E. Giant papillary conjunctivitis syndrome
   *F. Inclusion conjunctivitis in adults—acute mucopurulent follicular inflammation, persisting as long as several months, sometimes with scarring
   G. Syndrome of Beal—transient unilateral disease, usually resolving in weeks


**CHRONIC FOLLICULAR CONJUNCTIVITIS (LYMPHOID FOLLICLES COBBLES TONING) OF CONJUNCTIVA WITH LONG-TERM COURSE**

1. Chronic follicular conjunctivitis—Axenfeld's type (orphan's) frequently found in institutionalized children; almost asymptomatic; long duration (to months or longer); no keratitis; cause unknown

*2. Chronic follicular conjunctivitis, toxic type
   A. Bacterial origin, such as that due to a diplobacillus or other microorganism
   B. Drugs, including the following:

<table>
<thead>
<tr>
<th>acyclovir</th>
<th>diatrizoate meglumine</th>
<th>F3T</th>
</tr>
</thead>
<tbody>
<tr>
<td>adenine arabinoside</td>
<td>and sodium</td>
<td>framycetin</td>
</tr>
<tr>
<td>amphotericin B</td>
<td>diisopropyl</td>
<td>gentamicin</td>
</tr>
<tr>
<td>apraclonidine</td>
<td>fluorophosphate</td>
<td>homatropine</td>
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<tr>
<td>atropine</td>
<td>dipivefrin</td>
<td>hyaluronidase</td>
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<tr>
<td>carbachol</td>
<td>DPE</td>
<td>idoxuridine</td>
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<tr>
<td>clonidine</td>
<td>echothiophate</td>
<td>isoflurophate</td>
</tr>
<tr>
<td>demecarium</td>
<td>eserine</td>
<td>ketorolac tromethamine</td>
</tr>
</tbody>
</table>
methscopolamine  pilocarpine  sulfisoxazole
neomycin  scopolamine  trifluorothymidine
neostigmine  sulfacetamide  trifluridine
physostigmine  sulfamethizole  vidarabine

3. Chronic follicular conjunctivitis with epithelial keratitis; differentiated from Axenfeld type by shorter duration (to months) and by epithelial keratitis involving upper third of cornea; epidemic in schools; can be transmitted by mascara pencil; cause unknown
4. Ectodermal syndrome (Rothmund syndrome)
5. Folliculosis-associated general lymphoid hypertrophy
*6. Molluscum contagiosum conjunctivitis
7. Neurocutaneous syndrome (ectodermal dysgenesis)
8. Parinaud syndrome-chronic fever and regional lymphadenopathy, frequently cat-scratch fever
*9. Postoperative penetrating keratoplasty or cataract surgery sutures
10. Sebaceous carcinoma with papillary conjunctivitis
11. Trachoma-stages to 3
12. Use of hard and soft contact lens
*14. Use of ocular prostheses
15. With generalized lymphadenopathy


CICATRICIAL CONJUNCTIVITIS (SCARRING OF CONJUNCTIVA)

*1. General: a postinfectious type of membranous conjunctivitis such as C. diphtheriae, streptococcal conjunctivitis, autoimmune or presumably autoimmune sarcoidosis, scleroderma, Stevens-Addison, pemphigoid, lichen planus, atopic blepharoconjunctivitis, miscellaneous causes and linear IgA dermatosis.
2. Upper lid
   A. Trachoma
3. Lower lid
   *A. Acne rosacea (ocular rosacea)
   B. Chemical (especially alkali)
   C. *Chlamydia* organisms (psittacosis-lymphogranuloma group)
   D. Chronic cicatricial conjunctivitis - occurs in the elderly; has a chronic course; may have concurrent skin and mucous membrane lesions
   *E. Congenital syphilis
   F. Dermatitis herpetiformis
   G. Epidemic keratoconjunctivitis
   H. Epidermolysis acuta toxica (Lyell syndrome)
   I. Epidermolysis bullosa
   *J. Erythema multiforme (Stevens-Johnson disease)
   K. Erythroderma ichthyosiforme
   L. Exfoliative dermatitis
   M. Fuchs-Lyell syndrome
   N. Hydroa vacciniforme
   O. Impetigo
   P. Lamellar ichthyoses
   *Q. Ocular pemphigoid
   R. Paraneoplastic lichen planus
   S. Radium burns
   T. Reiter syndrome (conjunctivourethrosynovial syndrome)
   U. Sjögren syndrome (secretoinhibitor syndrome)
   V. Staphylococcal granuloma
   W. Syphilis (acquired lues)
   X. Systemic scleroderma (progressive systemic sclerosis)
   Y. Vaccinia

*4. Drugs
   A. Demecarium bromide
   *B. Echothiophate iodide
   C. Idoxuridine
   D. Penicillamine
   E. Pilocarpine
   F. Practolol
   G. Thiabendazole
   H. Timolol
   *I. Topical ocular epinephrine


**ANGULAR CONJUNCTIVITIS (INFLAMMATION AT ANGLE OF EYE, USUALLY LATERAL)**

1. *C. albicans*
2. *M. lacunata* (Morax-Axenfeld diplobacillus)
3. Stannus cerebellar syndrome (riboflavin deficiency)
4. *S. aureus*


**CONJUNCTIVAL DISORDERS ASSOCIATED WITH DERMATOLOGIC DISORDERS**

1. Dermatoses
   A. Acanthosis nigricans
   B. Acne rosacea
   C. Acrodermatitis chronica atrophicans
   D. Acrodermatitis enteropathica
   *E. Atopic eczema dermatitis
   F. Diffuse cutaneous mastocytosis
   G. Erythroderma exfoliativa (Wilson-Brocq disease)
   H. Ichthyosis
   I. Keratosis follicularis
   J. Keratosis follicularis spinulosa decalvans
   K. Lichen planus
   L. Pityriasis rubra pilaris; lichen acuminatus
   M. Porokeratosis
   N. Psoriasis vulgaris
   O. Seborrhea
   P. Xeroderma pigmentosum
2. Mucocutaneous eruptions
   *A. Behçet disease (dermatostomatooophthalmic syndrome)
   B. Benign mucous membrane pemphigoid
   C. Dermatitis herpetiformis (Duhring-Brocq disease)
   *D. Erythema multiforme (Stevens-Johnson disease)
   E. Epidermolysis bullosa
   F. Hydroa vacciniforme (recurrent summer eruption)
   G. Pemphigus-vulgaris, vegetans, foliaceus
   H. Pyostomatitis vegetans
   I. Reiter disease (polyarthritis enterica)


**CONJUNCTIVAL DISORDERS ASSOCIATED WITH GENITAL DISORDERS**

1. Bacteria
   A. *Bacteroides* species
   B. *Calymmatobacterium granulomatis* (granuloma inguinale)
   C. *E. coli*
   D. *Haemophilus ducreyi*
   E. *Haemophilus vaginalis*
   F. *Mimeae* species
   G. *Mycobacterium leprae*
   H. *M. tuberculosis*
   *I. N. gonorrhoeae*
   J. *Proteus* species
   *K. P. aeruginosa*
   L. *Staphylococcus* species
   M. *Streptococcus* species

2. Fungi
   A. *Candida* species
   B. Other

3. Viruses
   A. Cytomegalovirus
   *B. Herpes virus hominis 2
   *C. *Molluscum contagiosum* virus
   D. Rubella
   E. Varicella zoster
   F. Verruca virus

4. Spirochetes
   A. *T. pallidum*

5. Chlamydiae
   A. *Chlamydia lymphogranuloma*
   *B. Chlamydia oculogenitalis*
C. Unclassified Chlamydia from Reiter disease

6. Parasites
   A. Beetles
   B. Fly larvae
   C. Moths
   D. Phthirus pubis
   E. T. vaginalis


CONGESTION OF CONJUNCTIVA (NONINFECTIOUS HYPEREMIA OF THE CONJUNCTIVA)

* 1. Acute lupus erythematosus (Kaposi-Libman-Sacks syndrome)
2. Alcoholism
*3. Allergic conjunctivitis, such as contact with cosmetics or plastic
4. Avitaminosis
5. Carcinoid syndrome
*6. Carotid-cavernous fistula or arteriovenous aneurysm
7. Cavernous sinus thrombosis (Foix syndrome)
8. Conjunctivitis caused by air pollution (smog, dust, or smoke)
*9. Drugs causing conjunctival hyperemia, including the following:

acetoxyamide  aurothioglucose  cefazolin
acetylcholine  aurothioglycanide  cefonicid
acyclovir  barbital  cefoperazone
adrenal cortex injection  beclomethasone  ceforanide
adsorbed  belladonna  cefotaxime
alcohol  bendroflumethiazide  cefotetan
aldosterone  benzalkonium  cefoxitin
allobarbital  benzthiazide  cefsulodin
allopurinol  betamethasone  ceftazidime
alprazolam  bischloroethylnitrosourea  ceftizoxime
alseryxolol  (BCND)  ceftriaxone
alseryxolol  brimonidine tartrate  cefuroxime
antazoline  buptalbital  cephaloglycin
aminohydrozone  buptalbital  cephalodine
amobarbital compounds  butalbital  cephalothin
and pertussis vaccine  butalbital  cephapirin
antazoline  butalbital  cephapirin
apractoloidine  carbachol  cephadine
aprobarbital  carmustine  chloral hydrate
aspirin  cefaclor  chloramphenicol
atropine  cefadroxil  chlordiazepoxide
auranofin  cefamandole  chlordiazepoxil
| chloroform                      | emetine                      | lithium carbonate              |
| chloroprocaine                 | ephedrine                    | lorazepam                      |
| chlorothiazide                 | erythromycin                 | maprotiline                    |
| chlorpropamide                 | ether                        | meperidine                     |
| chlortetracycline              | ethotoin                     | mephenytoin                    |
| chlorthalidone                 | ethyl biscoumacetate         | mepobartal                     |
| chrysarobin                    | etidocaine                   | mepivacaine                    |
| cimetidine                     | F3T                          | mercuric oxide                 |
| cisplatin                      | fenoprofen                   | metaraminol                   |
| clindamycin                    | fludrocortisone              | methacholine                   |
| clindamycin                    | fluorescein                  | metharbital                    |
| clofibrate                     | fluorometholone              | methimazole                    |
| clonazepam                     | fluprednisolone              | methitural                     |
| clorazepate                    | flurazepam                   | methocarbamol                  |
| colchicine                     | flurbiprofen                 | methohexital                   |
| colloidal silver               | gentamicin                   | methoxamine                    |
| cortisone                      | glyburide                    | methoxsalen                   |
| cyclobarbital                  | gold Au                      | methylchlorothiazide           |
| cyclopentobarbital             | gold sodium thiomalate       | methylcyclohexyl               |
| cycloserine                    | gold sodium thiosulfate      | methylflumethiazide            |
| cyclosporine                   | griseofulvin                 | methylprednisolone             |
| cyclothiazide                  | halazepam                    | methyldopa                     |
| cytarabine                     | heparin                      | methylprednisolone             |
| deferoxamine                   | heptabarbital                | methylthiouracil               |
| deserpidine                    | hexethal                     | metolazone                     |
| desoxyxycorticosterone         | hexobarbital                 | metoprolol                     |
| dexamethasone                  | homatropine                  | metronidazole                  |
| dextran                        | hydralazine                  | mianserin                     |
| dextrothyroxine                | hydrochlorothiazide          | midazolam                     |
| diacetylmorphine               | hydrocortisone               | minoxidil                      |
| diatrizoate meglumine          | hydroflumethiazide           | morphine                       |
| and diazepam                   | ibuprofen                    | moxalactam                     |
| dicumarol                      | idoxuridine (IDU)            | mumps virus vaccine            |
| diethylycarbamazine            | indapamide                   | (live)                         |
| difumarate                     | indomethacin                 | naproksen                      |
| diltiazem                      | interferon                   | nifedipine                     |
| dimercaprol                    | iodide and iodine            | nitrazepam                     |
| diphenadione                   | solutions                    | nitromersol                    |
| diphtheria and tetanus         | iothalamate meglumine        | norepinephrine                 |
| toxoids                        | and iothalamic acid          | olopatadine HCI                |
| disodium cladronate            | ketoprofen                   | opium                          |
| disodium etidronate            | levothryroxine               | oxazepam                       |
| disodium pamidronate           | lidocaine                    |                              |
| emedastine                     | lincomycin                   |                              |
| disopyramide                   | liothyronine                 |                              |
| doxorubicin                    | liotrix                      |                              |
oxprenolol
oxyphenbutazone
oxyphenonium
paramethasone
pentazocine
pentobarbital
phenacetin
phenindione
phenobarbital
phenoxybenzamine
phenprocoumon
phénybutazone
phenylephrine
phenylmercuric acetate
phenylmercuric nitrate
pilocarpine
piroxicam
polythiazide
practolol
prazepam
prazosin
prednisolone
prilostone
prilocaine
primidone
probabral
procaine
propoxycaine
propranolol
propylthiouracil
quinethazone
radioactive iodides
ranitidine
rauwolfia serpentina
rescinnamine
rifampin
rubella and mumps virus
rubella virus vaccine
(slive)
scopolamine
secobarbital
sildenafil citrate
silver nitrate
silver protein
sodium chloride
sodium salicylate
streptomycin
sulfacetamide
sulfachlorpyridazine
sulfacytine
sulfadiazine
sulfadimethoxine
sulfamerazine
sulfameter
sulfamethazine
sulfamethoxazole
sulfamethoxypyridazine
sulfanilamide
sulfaphenazole
sulfapyridine
sulfathiazole
sulfisoxazole
syrosingopine
talbutal
temazepam
tetanus immune globulin
tetanus toxoid
tetracycline
thiabendazole
thiacetazone
thiamylal
thiopental
thiopental
thiogloablurb
thiobendazole
thiacetazone
thiamylal
thiopental
thiopental
thyroglubulin
thyroid
tolazamide
tolazoline
tolbutamide
triamcinolone
triazolam
trichloromethiazide
trichloroethylene
trifluridine
trioxsalen vaccine (live)
vancocycin
verapamil
vidarabine
vinbarbital
vitamin A

10. Gout (hyperuricemia)
11. Hormone deficiency (estrogenic)
12. Hypothyroidism
13. Irritative follicular conjunctivitis (see p. 193)

*A. Chemical conjunctivitis because of drugs, including the following:

acenocoumarol
acematinophen
aceltanilid
acetohexamide
allobarbital
allopurinol
alprazolam
amobarbital
anisindione
antazoline
antipyrine
apobarbital
aspirin
barbital
bendroflumethiazide
benzalkonium
benzathine
benztiazide
butabarbital
butalbital
butallylinal
butethal
carbamazepine
carbimazole
cefaclor
B. Topical drugs that are hypotonic, hypertonic, or in which the pH is above or below 6.9 or a drug degradation causing chemical irritation
C. Toxic conjunctivitis because of drugs such as miotics or cycloplegics
D. Vegetable irritants (e.g., caster bean)

*14. Malignant lymphoma
15. Ophthalmic vein thrombosis
16. Photosensitive conjunctivitis
*17. Polycythemia vera (Vaquez disease)
18. Sjögren syndrome (secretoinhibitor syndrome)
19. Vascular changes
   A. Facial paralysis (see p. 66-68)
   *B. Hereditary hemorrhagic telangiectasis (Rendu-Osler-Weber disease)
   C. Petechial hemorrhage of conjunctiva (see p. 206)


**CILIARY FLUSH**

Ciliary flush involves circumcorneal congestion and congestion of ciliary vessels immediately surrounding the cornea; individual vessels are not seen; color is violaceous; redness fades toward the fornices; and vessels do not move with conjunctiva.

1. Corneal disease, such as with inflammations and erosions
2. Glaucoma, especially acute glaucoma
3. Iridocyclitis
4. Iris irritation, such as with corneal foreign bodies
5. Iritis

CONJUNCTIVAL ANEURYSMS, VARICOSITIES, TORTUOUSITIES, AND TELANGIECTASIS

1. Local causes
   *A. Acne rosacea
   B. Chronic congestive glaucoma
   C. Delayed mustard gas keratitis
   D. Idiopathic anomaly
   E. Irradiation of the eye
   F. Long-standing ocular inflammation
   *G. Metastatic primary tumor
   *H. Pterygium
   *I. Underlying choroidal or ciliary body melanomas

2. Systemic causes
   A. Acquired immunodeficiency syndrome (AIDS)
   B. Arteriosclerosis
   C. Associated with familial amyloidotic polyneuropathy, type 1
   D. Ataxic telangiectasia (Louis-Bar syndrome)
   *E. Degos syndrome (malignant atrophic papulosis)
   F. Diabetes
   G. Dysproteinemia as in Waldenström macroglobulinemia, cryoglobulinemia, and multiple myeloma
   H. Endangiitis obliterans
   I. Fabry disease (diffuse angiokeratosis)
   J. Hereditary hemorrhagic telangiectasia (Rendu-Osler-Weber disease)
   K. Hypertension
   L. Klippel-Trenaunay-Weber syndrome (angioosteohypertrophy syndrome)
   M. Normal individuals
   *N. Pulmonary insufficiency
   O. Reimann syndrome (hyperviscosity syndrome)
   P. Renal failure
   Q. Rheumatic fever or rheumatic heart disease
   R. Scleroderma (progressive systemic sclerosis)
   S. Sturge-Weber syndrome (meningocutaneous syndrome)
   *T. Syphilis (acquired lues)
   U. Varicose veins-generalized


CONJUNCTIVAL SLUDGING AND SEGMENTATION

1. Local
   A. Aging
   B. Hypothermia
   C. Sympathetic irritation
   D. Vasodilator drugs that are applied locally

2. Systemic or hyperviscosity with increase in serum proteins
   A. Cryoglobulinemia
   B. Hyperglobulinemia
   C. Hypertension
   D. Macroglobulinemia (Waldenström syndrome)
   *E. Multiple myeloma (Kahler disease)
   *F. Sickle cell disease (Herrick syndrome)


CONJUNCTIVAL EDEMA (CHEMOSIS)

1. Acquired blockage of orbital lymphatics following orbital surgery (lateral orbitotomy) or because of erysipelas or lymphogranuloma venereum
   *2. Chronic hereditary lymphedema (Nonne-Milroy-Meige disease)
   *3. Drugs, including the following:

   acetohexamide  antimony lithium  benzalkonium
   acetophenazine  thiomalate  benzathine penicillin G
   acetyldigitoxin  antimony potassium tartrate  betamethasone
   actinomycin C  tartrate  bleomycin
   adrenal cortex injection  antimony sodium  brinzolamide
   albuterol  tartrate  bupivacaine
   aldosterone  antimony sodium  butabarbital
   allobarbital  thioglycollate  butacaine
   amantadine  antipyrine  butalbital
   aminopterin  aprobarbital  butallylone
   aminosalicylate  aspirin  butaperazine
   aminosaliclyc acid  auranofin  butethal
   amobarbital  aurolithoiglucose  cactinomycin
   amodiaquine  aurolithoiglycanide  captopril
   amphotericin B  barbital  carbachol
   and compounds  benoxinate  carbamazepine
carbenicillin
carphenazine
cefaclor
cefadroxil
cefamandole
cefazolin
cefonidic
cefoperazone
ceforanide
cefotaxime
cefotetan
cefoxitin
cefsulodin
ceftazidime
ceftizoxime
ceftriaxone
cefuroxime
cephaloglycin
cephaloridine
cephalothin
cephapirin
cephaprin
cephadine
chloral hydrate
chlorambucil
chloramphenicol
chlorhexidine
chlorisondamine
chloroprocaine
cloroquine
chlorpromazine
chlorpropamide
chlorotetrapylicine
chrysarobin
cisplatin
clofibrate
clomipramine
cloxacinil
cocaine
colistin
colloidal silver
cortisone
cyclobarbital
cyclopentobarbital
cyproheptadine
dactinomycin
danazol
dapsone
daunorubicin
demecarrium
demethylcycline
desipramine
deslanoside
desoxycorticosterone
dexamethasone
dextrothyroxine
diacetylmorphine
diacezizole meglumine
and
dibucaine
dicloxacillin
diethazine
diethylcarbamazine
digoxin
dionin
dorzolamide
doxepin
doxorubicin
doxycycline
dromostanolone
dyclonine
diothiolophate
dETIME
enalapril
epinephrine
ergonovine
ergotamine
erythromycin
ethopropazine
etidocaine
etretinate
F3T
floxacuridine
fluocortisone
fluorouracil
fluoxymestosterone
fluphenazine
fluprednisolone
flumethasone
fluoride
flurbiprofen
folic acid
foam
fordalin
folic acid
formalin
forxidine
furosemide
fusidic acid
fusidate
galaxy
galactose
gallamine pentolin
glyburide
gold Au 198
gold sodium thiomolate
griseofulvin
heptabarbital
hetacillin
hexachlorophene
hexamethonium
hexethal
hexobarbital
hydabamine penicillin
hydroazine
hydrocortisone
hydroxychloroquine
ibuprofen
idoxuridine
IDU
imipramine
iodide and iodine solutions
iron dextran
isofluorophate
isosorbide
isotretiino
ketoprofen
lanatoside C
levodopa
levothyruxine
lidocaine
lithium carbonate
mannitol
maprotiline
mecamylamine
medrysone
melphalan
mephabartal
mepivacaine
meprednisone
mercuric oxide
mesoridazine
metharbital
methdilazine
methitural
methoxhexital
methotrimetrazine
methylboma
methylergonovine
methylpentynol
methylprednisolone
methysergide  pipobroman  stibocaptate
metoclopramide  poliovirus vaccine  stibophen
metrizamide  polynoxylin B  streptomycin
metronidazole  potassium penicillin G  succinylcholine
mianserin  potassium phen ethicillin  sulindac
mild silver protein  potassium phen ethicillin  suramin
minocycline  practolol  talbutal
mitomycin  prazosin  testolactone
moxalactam  prednisolone  testosterone
nafcillin  prednisone  tetracaine
naproxen  prilocaine  tetracycline
neomycin  pridomide  tetraethylammonium
nitromersol  probarbital  thiamylal
nortriptyline  procaine  thiethylperazine
oral contraceptives  procaine penicillin G  thiamyl
ouabain  prochlorperazine  thioceptorzate
oxacillin  promazine  thioridazine
oxprenolol  promethazine  thiopropazate
oxyphenbutazone  propiomazine  thiopropazine
oxytetracycline  propoxyca ine  thiotepa
paramethasone  protriptyline  thyroid
pentobarbital  quinacrine  tolazamide
pentolinium  radioactive iodides  tolbutamide
perazine  rifampin  triamcinolone
pericyazine  rubella virus vaccine  trifluoperazine
perphenazine  (live)  trifluromazine
phenacaine  sanguinarine  trimetrazine
phenazine  secobarbital  trimethaphan
phenobarbital  silicone  trimethidinium
phenylbutazone  silver nitrate  urokinase
phenylephrine  silver protein  vidarabine
pilocarpine  sodium  vinbarbital
piperacetazine  sodium  vinblastine
piperazine  antimonylgluconate
piperocaine  sodium salicylate

4. Glandular fever
5. Hypersensitivity-local topical allergies
*6. Increased bulk of orbital contents-orbital tumors, cysts, or endocrine exophthalmos
7. Local inflammatory conditions
  *A. Cerebral cavity-acute meningitis
  *B. Eye-viral conjunctivitis, corneal ulcer, fulminating iritis, or panophthalmitis
  C. Lac rimal passages-dacryocystitis
  D. Lids-styes, vaccinia, acute meibomitis, insect bites, or vaccinal pocks
  E. Nasal cavity-sinusitis
  *F. Orbit-cellulitis, periostitis, dacrooadenitis, tenonitis
8. Myxedema-infiltration with mucopolysaccharides
9. Reduced plasma protein level-nephrotic state
10. Systemic lupus erythematosus
11. Vasomotor instability-angioneurotic edema or premenstrual phase of water retention
12. Venous congestion-local obstruction of orbital apex, carotid-cavernous fistula, thrombosis of cavernous sinus, or right-sided heart failure
13. Whipple disease


**CONJUNCTIVAL XEROSIS (DRYNESS OF CONJUNCTIVA)**

1. Absence of blinking
2. Drugs, including the following:
   - acebutolol (?)
   - amiodarone
   - atenolol (?)
   - betaxolol
   - busulfan
   - chlorambucil
   - clonidine (?)
   - cyclophosphamide
   - doxepin
   - ibuprofen
   - ketoprofen
   - labetalol (?)
   - levobunolol
   - methyldopa
   - metoprolol (?)
   - nadolol (?)
   - naproxen (?)
   - oxprenolol (?)
   - perhexiline
   - pindolol (?)
   - practolol
   - primidone
   - propanolol (?)
   - quinidine
   - sulindac
   - thiabendazole
   - timolol
   - vinblastine (?)

3. Following cicatricial conjunctivitis (see p. 194)
4. Illness or coma
   *5. Lack of closure of lids in sleep
5. Result of exposure of conjunctiva to air
   A. Deficient closure of lids, such as with paralysis of orbicularis, as part of facial palsy, spasms of the levator, or ectropion
   B. Excessive proptosis, such as in exophthalmic goiter or orbital tumor
   *7. Vitamin A deficiency
   A. Dietary deficiencies, including malnutrition, cystic fibrosis, anorexia nervosa, and bulimia
   B. Digestive tract disorders
      (1) Colitis and enteritis
      *(2) In pancreas-chronic pancreatitis
      (3) In stomach-achlorhydria, chronic gastritis or diarrhea, peptic ulcer
C. Hookworm disease
* D. Liver disease, such as chronic cirrhosis
E. Malaria
F. Pregnancy
G. Pulmonary tuberculosis
H. Skin disorders, such as pityriasis rubra pilaris
*I. Thyroid gland disorder, such as hyperthyroidism
J. Uyemura syndrome (fundus albipunctatus with hemeralopia and xerosis)

8. Decrease tear production
   A. Congenital alacrima
   B. Keratoconjunctivitis sicca
   C. Riley-Day syndrome (familial dysautonomia)
   D. Sjögren syndrome
   E. Surgical excision of the lacrimal and accessory lacrimal glands
   F. X-irradiation of the lacrimal gland

9. Following x-irradiation of the conjunctiva


**BITOT SPOTS**

Bitot spots are small gray or white, sharply outlined areas, cheeselike or foamy, occurring on either side of the limbus but especially in the temporal area.

1. Associated with coloboma of lid
2. Associated with corectopia, nystagmus, and absent foveal reflexes
3. Associated with Rieger anomaly
4. Congenital anomaly
5. Corneal snowflake dystrophy
6. Exposure
7. Idiopathic
*8. Keratosis follicularis (Darier -White disease) associated with retinitis pigmentosa
9. Pellagra or other poor nutritional states
10. Vitamin A deficiency


**SUBCONJUNCTIVAL HEMORRHAGE (BLOOD UNDER CONJUNCTIVA)**

1. Acute febrile systemic infections

   A. Bacteria, such as those responsible for meningococcal septicemia, subacute bacterial endocarditis, scarlet fever, diphtheria, typhoid fever, or cholera
   B. Parasites, such as plasmodia (malaria)
   C. Rickettsia, such as those causing typhus fever
   D. Unknown infective agents, such as those causing glandular fever
   E. Viruses, such as those responsible for influenza, smallpox, measles, yellow fever, sandfly fever, or Kaposi sarcoma

2. Associated with use of drugs, including the following:

   acetylcholine  
   acid bismuth sodium  
   tartrate  
   adrenal cortex injection  
   aldosterone  
   allopurinol  
   alseroxylon  
   aspirin  
   benoxinate  
   betamethasone  
   bismuth carbonate  
   bismuth oxychloride  
   bismuth salicylate  
   bismuth sodium tartrate  
   bismuth thioglycollate  
   bismuth sodium thiglycollamate  
   bismuth subcarbonate  
   bismuth subsalicylate  
   bupivacaine  
   butacaine  
   chloroprocaine  
   cobalt (?)  
   cocaine  
   combination products of estrogens and progestogens  
   cortisone  
   deserpidine  
   desoxycorticosterone  
   dexamethasone  
   dibucaine  
   dyclonine  
   epinephrine  
   ethambutol  
   etidocaine  
   fludrocortisone  
   fluorometholone  
   fluorouracil  
   glycerin  
   heparin  
   hexachlorophene  
   hydrocortisone  
   indomethacin (?)  
   iodide and iodine solutions and compounds (?)  
   isosorbide  
   ketoprofen  
   leuprolide acetate  
   lincomycin  
   mannitol  
   medroxyprogesterone  
   medrysone  
   mepivacaine  
   meprednisone  
   methaqualone  
   methylphenidate  
   methylprednisolone  
   mithramycin  
   mitotane  
   oxyphenbutazone  
   paramethasone  
   penicillamine  
   phenacaine  
   phenylbutazone  
   phenytin  
   piperocaine  
   plicamycin  
   pralidoxime  
   prilocaine  
   procaine  
   proparacaine  
   propoxycaire  
   radioactive iodides (?)  
   rauwolfia serpentina  
   rescinnamine  
   reserpine  
   sodium chloride  
   sodium salicylate  
   sulfacetamide  
   sulfachlorpyridazine  
   sulfacytine  
   sulfadiazine  
   sulfadimethoxine  
   sulfamerazine
sulfameter  sulfapyridine  tetracaine
sulfamethazine  sulfasalazine  triamcinolone
sulfamethizole  sulfathiazole  trichloroethylene
sulfamethoxazole  sulfisoxazole  urea
sulfamethoxypyridazine  sulindac  urokinase (?)
sulfanilamide  syrosingopine  vitamin A
sulfaphenazole  tamoxifen
sulfapyridine  sulfasalazine  sulfathiazole  sulfisoxazole  sulindac  syrosingopine  tamoxifen

3. Blood dyscrasias
   A. Associated with thrombocytopenia
      *(1) Anemias, especially, aplastic anemia
      *(2) Drugs, including the following:
      absorbed acebutolol  aurothioglycanide  cefonicid
      acebutolol  azatadine  cefoperazone
      acenocoumarin  azathioprine  ceforanide
      acenocoumarol  bacillus Calmette Guérin (BCG) vaccine  cefotaxime
      acetaminophen  barbital  cefotetan
      acetanilid  BCNU  cefoxitin
      acetzolamide  bendroflumethiazide  cefotetan
      acetohexamide  benzathine penicillin G  cefuroxime
      acetonide  benzthiazide  cephalexin
      acyclovir  bishydroxycoumarin  cephalexin
      allopurinol  bleomycin  cephaloglycin
      allopurinol  brompheniramine  cephaloridine
      alprazolam  butalbital  cephalothin
      amifostine  butalbital  cephalothin
      amitriptyline  butalbital  cepharadine
      amobarbital  butylbital  chlorambucil
      amodiaquine  butylbital  chlorambucil
      amphetamine  cactinomycin  chlorambucil
      amphotericin B  calcifediol  chlorambucil
      ampicillin  calcitriol  chlorambucil
      amisidine  captopril  chloroquine
      antimony lithium  carbamazepine  chloroquinone
      thiomalate  carbamazepine  chloroquinone
      antimony potassium  carbimazole  chlorothiazide
      tartrate  carbinoxamine  chlorthiazide
      antimony sodium  carisoprodol  chlorphentermine
      thioglycollate  carmustine  chlorpromazine
      antipyrene  carphenezine  chlorpromazine
      aprobarbital  cefaclor  chlorpromazine
      atenolol  cefadroxil  chlorpromazine
      auranofin  cefamandole  chlorprothixene
      aurothioglycine  cefazolin  chlortetracycline
      aurothioglycine  clotiapine  chlortetracycline
      aurothioglycanide  clofibrate  chlortetraycline
      azatadine  clofibrate  chloralose
clofibrate
clonazepam
clofibrate
cloxacillin
colchicine
cyclophosphamide
clorazepate
cyclopentobarbital
cycloferon
cycloserine
cyclophosphamide
closoxazole
cyclothiazide
cyclothiazide
cyclothiazide
cyclothiazide
cloproheptadine
cytarabine
dacarbazine
dactinomycin
dapsone
daunorubicin
deferoxamine
demeclocycline
desipramine
dexamethasone
dexchlorpheniramine
dezocine
diazepam
diazoxide
dichlorphenamide
dicloxacillin
dicumarol
diethazine
dihydrotachysterol
diltiazem
dimercaprol
dimethindene
dimethyl imidazole

carboxamide (DIC)
dimethyl sulfoxide
dimethyl sulfoxide
(DMSO)
diphenadione
diphenhydramine
diphenylhydantoin
diphenylpyrinaline
diphtheria and tetanus toxoids and pertussis vaccine
divalproex sodium
doxorubicin
doxycycline
doxylamine
dromostanolone
droperidol
enalapril
erythromycin
ethacrynic acid
ethopropazine
ethosuximide
ethotoin
ethoxzolamide
ethyl bisoumacetate
fenfluramine
fenoprofen
flecainide
flouxuridine
fluorouracil
fluoxymesterone
fluphenazine
flurazepam
furosemide
ganciclovir
gentamicin
glutethimide
glyburide
heparin
heptabarbitral
hetacillin
hexethal
hexobarbital
hydractin
hydrallazine
hydrallazine
hydrochlorothiazide
hydroflumethiazide
hydroxychloorquine
hydroxyurea
ibuprofen
imipramine
indapamide
indomethacin
interferon alpha, beta, or gamma
iopamidol
isocarboxazid
isoniazid
ketoprofen
labelalol
levodopa
lincomycin
lithium carbonate
lomustine
lorazepam
loxpine
maprotiline
measles virus vaccine
mechloretamine
mefenamic acid
melphalan
mephenytoin
mepobarbital
meprobamate
mercaptopurine
mesoridazine
methaclyline
methaqualone
metharbital
methazolamide
methdilazine
methicillin
methimazole
methitural
methohexital
methotrexate
methotrimetrazpine
methsuximide
methyclothiazide
methyldopa
methylen blue
methylene blue
methylphenidate
methythouracil
methyprylon
metolazone
metrizamide
metronidazole
mexiletine  |  mexiletine  |  mexiletine  |  mexiletine  |  mexiletine  |  mexiletine  |  mexiletine  |  mexiletine
mianserin  |  mianserin  |  mianserin  |  mianserin  |  mianserin  |  mianserin  |  mianserin  |  mianserin
minocycline |  minocycline |  minocycline |  minocycline |  minocycline |  minocycline |  minocycline |  minocycline
mitomycin  |  mitomycin  |  mitomycin  |  mitomycin  |  mitomycin  |  mitomycin  |  mitomycin  |  mitomycin
moxalactam  |  moxalactam  |  moxalactam  |  moxalactam  |  moxalactam  |  moxalactam  |  moxalactam  |  moxalactam
mumps virus vaccine (live)
 nadolol  |  nadolol  |  nadolol  |  nadolol  |  nadolol  |  nadolol  |  nadolol  |  nadolol
nafcillin  |  nafcillin  |  nafcillin  |  nafcillin  |  nafcillin  |  nafcillin  |  nafcillin  |  nafcillin
nalidixic acid |  nalidixic acid |  nalidixic acid |  nalidixic acid |  nalidixic acid |  nalidixic acid |  nalidixic acid |  nalidixic acid
naproxen  |  naproxen  |  naproxen  |  naproxen  |  naproxen  |  naproxen  |  naproxen  |  naproxen
nialamide  |  nialamide  |  nialamide  |  nialamide  |  nialamide  |  nialamide  |  nialamide  |  nialamide
nifedipine  |  nifedipine  |  nifedipine  |  nifedipine  |  nifedipine  |  nifedipine  |  nifedipine  |  nifedipine
nitrazepam  |  nitrazepam  |  nitrazepam  |  nitrazepam  |  nitrazepam  |  nitrazepam  |  nitrazepam  |  nitrazepam
nitrofurantoin  |  nitrofurantoin  |  nitrofurantoin  |  nitrofurantoin  |  nitrofurantoin  |  nitrofurantoin  |  nitrofurantoin  |  nitrofurantoin
nitroglycerin  |  nitroglycerin  |  nitroglycerin  |  nitroglycerin  |  nitroglycerin  |  nitroglycerin  |  nitroglycerin  |  nitroglycerin
nortriptyline  |  nortriptyline  |  nortriptyline  |  nortriptyline  |  nortriptyline  |  nortriptyline  |  nortriptyline  |  nortriptyline
oral contraceptives  |  oral contraceptives  |  oral contraceptives  |  oral contraceptives  |  oral contraceptives  |  oral contraceptives  |  oral contraceptives  |  oral contraceptives
orphenadrine  |  orphenadrine  |  orphenadrine  |  orphenadrine  |  orphenadrine  |  orphenadrine  |  orphenadrine  |  orphenadrine
oxacillin  |  oxacillin  |  oxacillin  |  oxacillin  |  oxacillin  |  oxacillin  |  oxacillin  |  oxacillin
oxazepam  |  oxazepam  |  oxazepam  |  oxazepam  |  oxazepam  |  oxazepam  |  oxazepam  |  oxazepam
oxybutazone  |  oxybutazone  |  oxybutazone  |  oxybutazone  |  oxybutazone  |  oxybutazone  |  oxybutazone  |  oxybutazone
oxytetracycline  |  oxytetracycline  |  oxytetracycline  |  oxytetracycline  |  oxytetracycline  |  oxytetracycline  |  oxytetracycline  |  oxytetracycline
paramethadione  |  paramethadione  |  paramethadione  |  paramethadione  |  paramethadione  |  paramethadione  |  paramethadione  |  paramethadione
penicillamine  |  penicillamine  |  penicillamine  |  penicillamine  |  penicillamine  |  penicillamine  |  penicillamine  |  penicillamine
penicillin  |  penicillin  |  penicillin  |  penicillin  |  penicillin  |  penicillin  |  penicillin  |  penicillin
pentobarbital  |  pentobarbital  |  pentobarbital  |  pentobarbital  |  pentobarbital  |  pentobarbital  |  pentobarbital  |  pentobarbital
perazine  |  perazine  |  perazine  |  perazine  |  perazine  |  perazine  |  perazine  |  perazine
pericyazine  |  pericyazine  |  pericyazine  |  pericyazine  |  pericyazine  |  pericyazine  |  pericyazine  |  pericyazine
perphenazine  |  perphenazine  |  perphenazine  |  perphenazine  |  perphenazine  |  perphenazine  |  perphenazine  |  perphenazine
phenacetin  |  phenacetin  |  phenacetin  |  phenacetin  |  phenacetin  |  phenacetin  |  phenacetin  |  phenacetin
phenelzine  |  phenelzine  |  phenelzine  |  phenelzine  |  phenelzine  |  phenelzine  |  phenelzine  |  phenelzine
phenformin  |  phenformin  |  phenformin  |  phenformin  |  phenformin  |  phenformin  |  phenformin  |  phenformin
phenindione  |  phenindione  |  phenindione  |  phenindione  |  phenindione  |  phenindione  |  phenindione  |  phenindione
 phenoxymethyl
pheniramine  |  pheniramine  |  pheniramine  |  pheniramine  |  pheniramine  |  pheniramine  |  pheniramine  |  pheniramine
phenobarbital  |  phenobarbital  |  phenobarbital  |  phenobarbital  |  phenobarbital  |  phenobarbital  |  phenobarbital  |  phenobarbital
phenoxyisopropyl
phenoxymethyl
phenoxymethylpenicillin  |  phenoxymethylpenicillin  |  phenoxymethylpenicillin  |  phenoxymethylpenicillin  |  phenoxymethylpenicillin  |  phenoxymethylpenicillin  |  phenoxymethylpenicillin  |  phenoxymethylpenicillin
phenprocoumon  |  phenprocoumon  |  phenprocoumon  |  phenprocoumon  |  phenprocoumon  |  phenprocoumon  |  phenprocoumon  |  phenprocoumon
phensuximide  |  phensuximide  |  phensuximide  |  phensuximide  |  phensuximide  |  phensuximide  |  phensuximide  |  phensuximide
phenylbutazone  |  phenylbutazone  |  phenylbutazone  |  phenylbutazone  |  phenylbutazone  |  phenylbutazone  |  phenylbutazone  |  phenylbutazone
phenytoin  |  phenytoin  |  phenytoin  |  phenytoin  |  phenytoin  |  phenytoin  |  phenytoin  |  phenytoin
pindolol  |  pindolol  |  pindolol  |  pindolol  |  pindolol  |  pindolol  |  pindolol  |  pindolol
piperacetazine  |  piperacetazine  |  piperacetazine  |  piperacetazine  |  piperacetazine  |  piperacetazine  |  piperacetazine  |  piperacetazine	pipibroman  |  pipibroman  |  pipibroman  |  pipibroman  |  pipibroman  |  pipibroman  |  pipibroman  |  pipibroman	polio virus vaccine  |  polio virus vaccine  |  polio virus vaccine  |  polio virus vaccine  |  polio virus vaccine  |  polio virus vaccine  |  polio virus vaccine  |  polio virus vaccine
polythiazide  |  polythiazide  |  polythiazide  |  polythiazide  |  polythiazide  |  polythiazide  |  polythiazide  |  polythiazide
potassium penicillin G  |  potassium penicillin G  |  potassium penicillin G  |  potassium penicillin G  |  potassium penicillin G  |  potassium penicillin G  |  potassium penicillin G  |  potassium penicillin G
potassium penicillin V  |  potassium penicillin V  |  potassium penicillin V  |  potassium penicillin V  |  potassium penicillin V  |  potassium penicillin V  |  potassium penicillin V  |  potassium penicillin V
potassium phenethicillin  |  potassium phenethicillin  |  potassium phenethicillin  |  potassium phenethicillin  |  potassium phenethicillin  |  potassium phenethicillin  |  potassium phenethicillin  |  potassium phenethicillin
potassium  |  potassium  |  potassium  |  potassium  |  potassium  |  potassium  |  potassium  |  potassium

(3) Leukemia
(4) Septicemias
Splenic disorders, such as Banti or Gaucher disease, Felty syndrome, and hemolytic icterus
*(6) Systemic lupus erythematosus (Kaposi-Libman-Sacks syndrome)
B. Ehlers-Danlos syndrome (fibrodysplasia elastica generalisata)
C. Hemochromatosis
D. Schomberg disease
E. Scurvy (avitaminosis C)
F. Secondary, such as that because of nephritic, cardiac, or hepatic disease
G. Thrombocytopenia purpura
4. Fragility of vessel walls because of systemic vascular disease
   A. Age
   B. Arteriosclerosis
   C. Diabetes
   D. Hypertension
   E. Nephritis
5. Gravity inversion
6. Injury to orbital or adjacent structures, such as sinus, basal skull fracture, subarachnoid hemorrhage
7. Local acute inflammation, including, acute pneumococcal conjunctivitis, leptospirosis ictero-hemorrhagica, epidemic typhus, and scrub typhus
8. Local trauma, including surgical trauma
9. Remote injury associated with fractured bones and fat emboli following angiography or open heart operation causing "splinter" subconjunctival hemorrhage
10. Spontaneous during menstruation
11. Spontaneous rupture of telangiectasis, varicosities, aneurysm, or angiomatous tumor (see p. 201)
* 12. Sudden severe venous congestion of head, including that because of coughing, vomiting, epileptic fit, strangulation, or an orbital tumor (neuroblastoma)
13. Without apparent cause-most common


TUMORS OF THE CONJUNCTIVA

1. Epithelial tumors
   A. Keratoacanthoma
B. Dyskeratosis
   (1) Epithelial plaques-leukoplakia, hereditary benign intraepithelial dyskeratosis
   (2) Intraepithelial epithelioma (Bowen disease) (61N)
C. Metastatic uveal melanoma
*D. Papilloma-including virus types 11, 16, and 18
E. Epithelioma
F. Adenoma
   (1) Papillary cystadenoma lymphomatosum (Warthin tumor)
   (2) Oncocytoma (oxyphil-cell adenoma)
   (3) Pleomorphic adenoma of Krause glands

2. Mesoblastic tumors
   A. Inflammatory hyperplasias
      (1) Granuloma
      (2) Plasmoma
   B. Connective tissue tumors
      (1) Fibroma
      (2) Lipoma
      (3) Myxoma

3. The reticuloses
   *A. Lymphoma
   B. Lymphosarcoma
   *C. Mycosis fungoides

4. Vascular tumors-angiomas
   A. Polymorphous hemangioma, telangiectatic granuloma, granuloma pyogenicum
   B. Lymphangioma
   C. Angiosarcoma monomorphous angioma, Kaposi (hemorrhagic) sarcoma

5. Pigmented tumors
   *A. Nevus
   B. Malignant melanoma
   C. Intraepithelial melanoma-precancerous melanosis

6. Peripheral nerve tumors
   A. Neurofibroma
      *(1) Neurilemmoma (neurinoma, schwannoma)
      (2) Malignant schwannoma (neurogenic sarcoma; neurofibrosarcoma)
      (3) Plexiform neurofibromatosis
   B. Tuberous sclerosis (Bourneville disease)
      *C. Intracranial nerve loops

7. Amyloidosis
8. Metastatic renal cell carcinoma
9. Trematode-induced granulomas
10. Hypertrophic discoid lupus erythematosus


**CONJUNCTIVAL CYSTS**

1. Congenital corneoscleral cyst (rare)
2. Epibulbar dermoids with cystic form
3. Epithelial cyst
   A. Apposition of folds of conjunctival mucosa (common)
   B. Downgrowth of epithelium-chronic inflammatory conditions, such as that following inflammation of pterygium
   C. Glandular retention-involvement of Krause glands in chronic inflammatory conditions, including trachoma and pemphigus
   D. Pigmented cyst appearing after prolonged topical use of cocaine or epinephrine
4. Limbal wounds with iris prolapse
5. Lymphatic cyst
6. Muscle inclusion cyst/complication of strabismus surgery
7. Parasitic cyst such as filarial cyst
8. Traumatic cyst (epithelial implantation)


**LIMBAL MASS**

1. Allergic reaction
   *A. Phlyctenules
   B. Vernal limbal lesions
2. Amyloid - perilimbal
3. Associated with skin disease
   *A. Acne rosacea (ocular rosacea)
   B. Hereditary benign intraepithelial dyskeratosis
   *C. Hodgkin disease
   D. Limbal squamous carcinoma in xeroderma pigmentosa
   E. Pityriasis rubra pilaris
   F. Psoriasis (psoriasis vulgaris)
   *G. Reticulum cell sarcoma-raised, pink, smooth lesions
4. Benign nodular fascitis
5. Dermoids
6. Ectopic lacrimal gland tissue
7. Epithelial hyperplasia
8. Fibrous histiocytoma
9. Fibroxanthoma
10. Granular cell tumor
11. Granulomas
12. Hemangioma
13. Intraepithelial epitheliomas (Bowen disease)
14. Lymphomas
15. Malignant melanomas
16. Mononucleosis (infectious)
17. Nevi
18. Papillomas
19. Pterygia
20. Sarcomas
21. Salmon patch associated with relapsing polychondritis
22. Squamous cell carcinoma
23. Subconjunctival nodules associated with Crohn disease
24. Synthetic fiber granuloma


**LARGE, FLAT, FLESHY LESIONS OF PALPEBRAL CONJUNCTIVA**

1. Accidental or surgical injuries
2. Carthy disease (pyorhinoblepharostomatitis vegetans)
   *3. Chalazion
   4. Embryonal rhabdomyosarcoma of children
   5. Granuloma pyogenicum
3. Ligneous conjunctivitis
4. Lymphogranuloma venereum
5. Meibomian cell carcinoma
6. Myopic infection
7. Papillary hyperplasia of vernal conjunctivitis
   *11. Syphilis
8. Tuberculosis
9. Tularemia


**CHRONIC OR RECURRENT ULCERS OF THE CONJUNCTIVA**

1. Behçet disease
2. Crohn disease
3. Drugs, including the following:
   - allopurinol
   - amphotericin B
   - aspirin
   - ferrocholinate
   - ferrous fumarate
   - ferrous gluconate
   - ferrous succinate
   - ferrous sulfate
   - floxuridine
   - fluorouracil
   - gentamicin
   - iron dextran
   - iron sorbitex
   - iron sulfate
   - phenytoin
   - polysaccharide- iron complex
   - sodium salicylate
4. Fungi
   *5. Herpes simplex*
6. Mucous membrane pemphigoid
7. Pseudomonas ulcer in patients with AIDS
8. Soft chancre
9. Syphilis (acquired lues)
10. Tuberculosis
11. Wegener granulomatosis


**PHLYCTENULAR KERATOCONJUNCTIVITIS**

This condition involves a localized conjunctival, limbal, or corneal nodule measuring about 1 to 3 mm.

* 1. Delayed hypersensitivity to bacterial protein, particularly tuberculoprotein and staphylococci; lymphopathia venereum and coccidioidomycosis may also be allergens
2. Malnutrition
3. Secondary infection of the conjunctiva, especially from S. aureus, pneumococcus, and Koch-Weeks bacillus
4. Systemic infection
   A. Bang disease (brucellosis)
   B. Candidiasis
   C. Neurodermatitis
   D. Mikulicz-Radecki syndrome (dacryosialoadenopathy)
   E. Trachoma
   F. Sjögren syndrome (secretoinhibitor syndrome)


**PIGMENTATION OF THE CONJUNCTIVA (SEE PIGMENT SPOTS OF SCLERA AND EPISCLERA)**
1. Blood pigments
   *A. After subconjunctival hemorrhage - red or later fine brown spots
   B. Yellow tinge of malaria, blackwater fever, or yellow fever
   C. Pigmentary limbal ring associated with senile, traumatic, or diseased conditions
2. Bile pigments (yellow)-obstructive or hemorrhagic jaundice
3. Melanin pigmentation
   A. Acanthosis nigricans
   *B. Addison disease (adrenal cortical insufficiency)
   C. Alcaptonuric ochronosis
   D. Chlorpromazine (Thorazine)
   E. Endogenous ochronosis
   F. Keratomalacia
   G. Trachoma
   *H. Use of epinephrine or epinephrine bitartrate, borate, and hydrochloride
   I. Vernal conjunctivitis
   J. Vitiligo (leukoderma)-increased conjunctival pigmentation
   K. Xeroderma pigmentosum
4. Drugs, including the following:
   acid bismuth sodium, bismuth subcarbonate, iron dextran
   tartrate, bismuth subsalicylate, iron sorbitol
   Alcian blue, captopril, ketoprofen
   amidarone, chloroquine, methacycline
   amodiaquine, chlortetracycline, methylene blue
   amphotericin B, chrysarobin, minocycline
   antimony lithium, clofazimine, minoxidil
   thiomalate, colloidal silver, oxytetracycline
   antimony potassium, demeclocycline, penicillamine
   tartrate, diethazine, polysaccharide iron complex
   antimony sodium, doxycycline
   tartrate, enalapril, quinacrine
   antimony sodium, ethopropazine, rifabutin
   thioglycollate, ferrocholinolate, rifampin silver nitrate
   antipyrine, ferrous fumarate, silver protein
   bismuth carbonate(?), ferrous gluconate, rose bengal
   bismuth oxychloride(?), ferrous succinate, sodium
   bismuth salicylate(?), ferrous sulfate, antimonylgluconate
   bismuth sodium tartrate, fluorescein, stibocaptate
   bismuth sodium, gold AU 198, stibophen
   thioglycollate (?), gold sodium thioglycollate, tetracycline
   bismuth sodium, gold sodium thiosulfate, trypan blue
   triglycollamate(?), hydroxychloroquine, vitamin A
5. Foreign substances such as silver (argyrosis), iron (siderosis), copper (chalcosis), arsenic (arsenic melanosis), gold (chrysisasis), aluminum, quinones, aniline dyes, and eye cosmetics containing carbon black
*6. Benign melanosis-overactivity of melanocytes
   A. Epithelial-congenital or acquired, for example, following radiation or use of chemicals (arsenic); in Addison disease; because of chronic conjunctivitis (trachoma, vernal conjunctivitis, onchocerciasis, keratomalacia)
   B. Subepithelial-congenital or in association with melanosis oculi or nevus of Ota

7. Neoplasms
   *A. Nevus-most common in children, localized stationary, elevated, cystic, mayor may not have pigmentation
   *B. Malignant melanoma arising from preexisting nevus, apparently normal conjunctiva, or from an area of acquired pigmentation (intraepithelial melanoma); occurs primarily in middle age; diffuse, flat, pigmentation; progressive; no cysts
   C. Secondary melanotic tumors
   D. Incidentally pigmented tumors, such as a melanocarcinoma
   E. Secondary metastatic tumors from lung or breast

8. Ocular causes, including the following:
   A. Apocrine adenocarcinomas
   B. Foreign bodies
   C. Hematic cysts
   D. Moll gland cystadenomas
   E. Staphylomas
   F. Subconjunctival hematomas


**DISCOLORATION OF CONJUNCTIVA**

1. Red
   A. Subconjunctival hemorrhage

2. Yellow
   A. Bilirubinemia-obstructive or hemorrhagic jaundice
   B. Picric acid
   C. Leptospirosis
   D. Brucellosis (Barg disease or Mediterranean fever)
   E. Aromatic nitro and amino compounds
   F. Conjunctival fat-occurs primarily in older and black patients
G. Blood pigment tinge of malaria, blackwater fever, and yellow fever

3. Gray (black)
   *A. Argyrosis (silver)
   B. Drugs, including the following:
      atabrine
      nitrochlorobenzene
      phenols, specifically phenylic acid and carbon disulfide
   C. Chrysiasis (gold)-grayish green effect
   D. Arsenicals-ash white
   E. Mascara

4. Brown
   A. Subconjunctival hemorrhage-fine brown spots
   B. Pigmentary limbal ring associated with senile, traumatic, or diseased conditions
      *C. Benign melanosis-overactivity of melanocytes
         (1) Epithelial-congenital or acquired, following radiation or use of chemicals (arsenic); in Addison disease (adrenal cortical insufficiency); because of chronic conjunctivitis (trachoma, vernal conjunctivitis, onchocerciasis, keratomalacia)
         (2) Subepithelial-congenital or in association with melanosis oculi or nevus of
   D. Neoplasms
      *(1) Nevus-most common in children, localized, stationary, elevated, cystic, mayor may not have pigmentation
      *(2) Malignant melanoma arising from preexisting nevus, apparently normal conjunctiva, or from an area of acquired pigmentation (intraepithelial melanoma); occurs primarily in middle age; diffuse, flat, pigmentation; progressive; no cysts
   E. Drugs, including the following:
      aminoquinoline combinations
      (benzoquinone, paraquinone, hydroquinone)
      aniline dyes
      bromides
      chromic acid and chromates phenol derivatives
      phenothiazine
      sympathomimetics (adrenalin)

5. Blue pigmentation
   A. Ink tattoo from pens
   B. Manganese dust


**SYMBLEPHARON**

Symblepharon involves fusing of the eyelid to an opposing surface, such as the tarsal and bulbar conjunctiva.

1. Physical trauma with denuded epithelium, including purulent, membranous, bullous, or ulcerative conjunctivitis and trauma
   2. Chemical burns—especially lime or caustic burns
   3. Inflammation—especially from drug reactions, including:

<table>
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<th>Drug</th>
<th>Effect</th>
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<tr>
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<td>vidarabine</td>
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<tr>
<td>silver nitrate</td>
<td>vinbarbital (?)</td>
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4. Long-standing acute inflammation
   A. Pemphigus (Cazenave disease)
   B. Stevens-Johnson disease (dermatostomatitis)

5. Congenital
6. Associated with cyanoacrylate tissue adhesive
7. Epidemic keratoconjunctivitis

CONJUNCTIVAL CONCRETIONS

Conjunctival concretions are small yellow spots that are most common in tarsal conjunctiva.
1. Chronic inflammatory conditions, including atopic keratoconjunctivitis, vernal conjunctivitis, and posttrachomatous degenerations
2. Elderly
3. Calcium deposits in patients with chronic renal failure treated with maintenance hemodialysis


LESIONS OF CARUNCLE

1. Apocrine hydrocystoma
*2. Basal cell carcinoma
3. Capillary hemangioma
4. Chronic inflammation
5. Dermoid
6. Ectopic lacrimal gland
7. Epithelial inclusion cyst
8. Foreign-body granuloma
9. Granular cell myeloblastoma
10. Histiocytic lymphoma
11. Lipogranuloma
12. Lymphangiectasis
13. Malignant melanoma
14. Nevus
15. Normal caruncle
*16. Oncocytoma
17. Papilloma
18. Pilar cyst
19. Plasmacytoma
20. Pyogenic granuloma
21. Reactive lymphoid hyperplasia
22. Sebaceous gland hyperplasia
*23. Sebaceous gland adenoma
24. Seborrheic keratosis
25. Squamous cell carcinoma


MICROPHTHALMIA (SMALL GLOBE)

1. Microphthalmia associated with the following:
   A. Cataract-dominant inheritance
   B. Coloboma-dominant and sex-linked inheritance
   C. Congenital spastic diplegia-x-linked
   D. Ectopic pupils-dominant inheritance
   E. Glaucoma-recessive inheritance
   F. Harelip and cleft palate-autosomal recessive
   G. High hypermetropia-recessive inheritance
   H. Malformation of hands and feet-autosomal recessive
   I. Polydactyly-autosomal recessive
   J. Retinitis pigmentosa and glaucoma-dominant inheritance

2. Colobomatous microphthalmia
   A. X-linked
      (1) Aicardi syndrome
      (2) Bloch-Sulzberger syndrome (incontinentia pigmenti)
      (3) Goltz syndrome (focal dermal hypoplasia)
      (4) Lenz microphthalmia syndrome
   B. Autosomal recessive
      (1) Cohen syndrome
      (2) Ellis-van Creveld syndrome
      (3) Hepatic fibrosis, polycystic kidneys, colobomas, and encephalopathy
      (4) Humeroradial synostosis
      (5) Kartagener syndrome
      (6) Laurence-Moon-Biedl syndrome
      (7) Marinesco-Sjögren syndrome
      (8) Meckel syndrome
      (9) Micro syndrome
(10) Sjögren-Larsson syndrome
(11) Warburg syndrome

C. Autosomal dominant
(1) Basal cell nevus syndrome
(2) Congenital contractual arachnodactyly
(3) Crouzon syndrome
(4) Stickler syndrome
(5) Treacher Collins syndrome
(6) Tuberous sclerosis
(7) Zellweger syndrome

D. Chromosomal abnormalities
(1) Deletions 4p, 4r, 11q, 13q, 18q, 18r, XO
(2) Duplications 3q, 4p, 4q, 7q, 9p, 9q, 13q, 22q
(3) Ring B syndrome
(4) Triploidy
(5) Trisomy 8, 9, 13, 17, 18, XXX, XYY

E. Unknown cause
(1) Amniogenic band syndrome (Streeter dysplasia)
(2) Cat’s-eye syndrome (Schmid-Fraccaro syndrome)
(3) CHARGE (colomba, heart disease, atresia choanae, retarded growth and retarded growth development or central nervous system anomalies, genital hypoplasia, and ear anomalies, or deafness) syndrome
(4) Dyscraniopygophalangea (Ullrich syndrome)
(5) Facial-clefting syndromes
(6) Frontonasal dysplasia (median cleft face syndrome)
(7) Goldenhar syndrome (oculoauriculo vertebral syndrome)
(8) Hemifacial microsomia syndrome
(9) Linear sebaceous nevus syndrome
(10) Rubinstein - Taybi syndrome

3. Noncolobomatous microphthalmia
   A. X-linked
(1) Anderson-Warburg syndrome
(2) Forsius-Eriksson syndrome (Aland disease)
(3) Lowe syndrome (oculocerebrorenal syndrome)

B. Autosomal recessive
(1) Cerebrooculofaci oskeletal syndrome
(2) Comadi syndrome
(3) Cross syndrome
(4) Diamond-Blackfan syndrome
(5) Fanconi
(6) Obesity-cerebral-ocular-skeletal anomalies syndrome

C. Autosomal dominant
(1) Blatt syndrome
(2) Gansslen syndrome
(3) Hypomelanosis of Ito syndrome
(4) Leri syndrome
(5) Myotonic dystrophy
(6) Rieger syndrome

D. Chromosomal abnormalities
(1) Duplication 10q
(2) Trisomy 21Q syndrome
(3) Chromosome deletion Xp22.1

E. Unknown cause
(1) Arachnoidal cyst
(2) Gorlin-Chaudhry-Moss syndrome
(3) Hallerman-Streiff syndrome
(4) Hutchinson-Gilford syndrome (progeria)
(5) Krause syndrome (encephalooophthalmic)
(6) Meyer-Schwickerath and Weyers syndrome
(7) Pierre Robin syndrome
(8) Retinal disinsertion syndrome
(9) Sabin-Feldman syndrome
(10) Weyers syndrome

F. Infectious etiology
(1) Congenital rubella (Gregg syndrome)
(2) Congenital spherocytic anemia
(3) Congenital toxoplasmosis
(4) Cytomegalovirus
(5) Epstein-Barr syndrome
(6) Herpes virus
(7) Mumps
(8) Varicella

G. Intoxicants
(1) Fetal alcohol effects
(2) Maternal phenylketonuria fetal effects

4. Idiopathic
5. Nanophthalmos


**BUPHTHALMOS (LARGE GLOBE)**

Buphthalmos usually is associated with corneal abnormalities such as opacities and rupture of Descemet membrane; the transition from cornea to sclera is unclear, and a thin, bluish sclera may be present.
1. Associated with anterior chamber cleavage syndrome (Reese-Ellsworth syndrome)
2. Autosomal recessive inheritance
3. Cerebrohepatorenal syndrome (Smith-Lemli-Opitz syndrome)
4. Chondrodystrophy calcificans congenita (Conradi syndrome)
5. Congenital glaucoma
6. Congenital rubella syndrome (Gregg syndrome)
7. Cryptophthalmia syndrome (cryptophthalmos-syndactyly)
8. Hurler syndrome
9. Krabbe syndrome
10. Lowe syndrome (oculocerebrorenal syndrome)
11. Milroy disease (chronic hereditary edema; Noone-Milroy-Meige disease)
12. Neurofibromatosis (von Recklinghausen disease)
13. Oculodentodigital dysplasia
14. Rieger syndrome (hypodontia and iris dysgenesis)
15. Sporadic occurrence
16. Sturge-Weber syndrome (encephalotrigeminal syndrome)


**PSEUDOENDOPHTHALMITIS (CONDITIONS THAT SIMULATE ENDOPHTHALMITIS)**

*1. Chemical reactions from irritating chemicals (irrigating solutions or medications) introduced into the anterior chamber
2. Foreign material in the anterior chamber
3. Metastatic carcinoma
4. Retained lenticular material
5. Severe postoperative iridocyclitis
6. Toxic anterior segment syndrome (TASS)*


**ENDOPHTHALMITIS (INTRAOCULAR INFECTION)**
1. **Bacterial agents**
   
   A. **Gram positive**
      
      (1) Bacillus subtilis, megaterium, anthracis, cereus
      (2) Clostridium peifringens (B. welchii)
      (3) Clostridium tetani
      (4) Coryneform bacterium
      (5) Diplococcus pneumoniae (Pneumococcus)
      (6) Diphtheroids
      (7) Listeria monocytogenes
      *(8) Propionibacterium acnes
      *(9) Staphylococcus aureus, albus, and epidermidis
      (10) Streptococcus viridans, S. hemolytic, S. pneumoniae, and Pyogenes salivarius
   
   B. **Gram negative**
      
      (1) Aerobacter aerogenes
      (2) Enterobacter cloacae
      (3) Escherichia coli
      (4) Fusobacterium organisms
      (5) Klebsiella pneumoniae (Friedlander bacillus)
      (6) Meningococci
      (7) Morganella species
      (8) Mycobacterium organisms
      (9) Neisserias catarrhalis
      (10) Ochrobactrum anthropi
      (11) Pasteurella multocida and tularensis
      (12) Proteus vulgaris (B. proteus) and mirabilis
      *(13) Pseudomonas aeruginosa (B. pyocyaneus)
      (14) Serratia marcescens
      (15) Yersinia enterocolitica or Y. pseudotuberculosis

2. **Fungal agents**
   
   A. Acanthamoeba
   B. *Actinomyces* species, including *Nocardiosis*
   C. Aspergillus species
   D. *Blastomyces dermatitidis*
   E. *Candida* species
   F. *Cephalosporium* species, hyphas
   G. *Coccidioides immitis*
   H. *Cryptococcus neoformans*
   I. *Hormodendrum*
   J. *Hyalopus bogolepofi*
   K. *Hyalosporus*
   L. *Mucormycosis* species
   M. *Neurospora sitaphila*
   N. *Sporothrix schenckii*
   O. *Sporotrichum schenckii*
   P. *Volutella* species
3. Viral agents
   A. Behçet syndrome (dermatostomatoophthalmic syndrome)
   B. Cytomegalovirus
   C. Myxovirus (influenza)
   D. Nocardia asteroides
   E. Vaccinia
   F. Variola

4. Nematode agents
   A. Taenia solium
   B. Toxocara canis and T. cati

5. Other agents
   A. Mycosis fungoides
   B. Exophiala jeanselmei (yeast)


**INTRAOCULAR CARTILAGE**

1. Angiomatosis of the retina
2. Chromosome deletion
3. Chronic inflammation
4. Facial nevus of Jadassohn (linear sebaceous nevus syndrome)
5. Incidental findings in microphthalmic eye, microphthalmos with cyst, microphthalmic eye from a cyclopic orbit, in eyes with coloboma of the choroid and retina or ciliary body
6. Incontinentia pigmenti (Bloch-Sulzberger disease)
7. Persistent hyperplastic primary vitreous
8. Retinal dysplasia
9. Teratoid medulloepithelioma (dictyomas)
10. Trisomy (13-Patau syndrome) (globe less than 10 mm in diameter)


**INTRAOCULAR CALCIFICATIONS**

1. Choroidal osteoma
2. Facial nevus of Jadassohn (linear sebaceous nevus syndrome)
3. Intraocular calcifications
   A. Congenital deformity
   B. Gitelman syndrome
   C. Recurrent iritis and keratitis
   D. Retinal detachment
   E. Trauma (perforating, nonperforating, or surgical)
4. Intraocular sarcoma
   *5. Retinoblastoma
6. Retinopathy of prematurity (end stage)
7. Sites of intraocular calcification
   A. Calcific emboli of retinal and ciliary arteries
   B. Cyclitic membrane
   C. Lens
   D. Peripapillary choroid
   E. Posterior pole to ora serrata in region of choroid and pigment epithelium
   F. Retina
   G. Vitreous


**INTRAOCULAR ADIPOSE TISSUE**

1. Congenital malformations
   A. Dermoid or dermolipoma extending from the cornea or limbus into the globe
   B. Malformed optic nerve
C. Persistent hyperplastic vitreous (PHPV) and other related ocular malformations, such as microphthalmia, persistent hyaloid vessels, cataract, and abnormal differentiation of the angle of the anterior chamber
2. Embolic phenomenon secondary to crush wounds of the thorax and abdomen or fracture of long bones of the extremities
3. Formation of fatty tissue within the marrow spaces of metaplastic bone
4. Missile passing through orbit carrying orbital fat into the eye


**SOFT GLOBE (DECREASED INTRAOCULAR PRESSURE)**

* 1. Fistula from intraocular source, including penetrating intraocular trauma or surgery and ruptured wall of the globe
* 2. Laser or cryotherapy ciliodestructive procedure
  3. *Phthisis bulbi*
4. Choroidal detachment
5. Injury to the cervical sympathetic nerve
6. Serous detachment of the retina
7. Myotonic dystrophy (Curschmann-Steinert syndrome)
8. Systemic disturbances
   A. Cardiac edema
   B. Diabetic coma
   C. Extreme or rapid dehydration because of malnutrition, cholera, or diarrhea
   D. Decrease in ocular blood pressure due to hypotension, ligation of the carotid artery, carotid occlusion, or pulseless disease (Takayasu syndrome)
   E. Giant cell arteritis (temporal arteritis syndrome)
   F. Leprosy (Hansen disease)
   G. Parkinson disease (shaking palsy)
   H. Postencephalitic syndrome following severe cerebral trauma, barbiturate poisoning, in deep anesthesia, following leukotomy, or on the paralyzed side in cases of cerebral hemiplegia
   I. Severe abdominal disturbances, such as intestinal perforation or obstruction
   J. Profound anemias
   K. Uremic coma
9. Drugs, including the following:
   acebutolol      adrenal cortex injection      amobarbital
   acebutolol      albuterol                  amyl nitrite
   aceclidine      alcohol                    antazoline
   acetazolamide   aldosterone               aprobabarbital
   acetylcholine   allobarbital              aspirin
   acetyldigitoxin alseroxylon               atenolol
propranolol  succinylcholine  trichloroethylene
protriptyline  syrosingopine  trifluoperidol
pyrilamine  talbutal  trimethaphan
triplelenamine  tetraethylammonium  trimethidinium
quinethazone  tetrahydrocannabinol  trolnitate
rauwolfia serpentina  tetrahydrozoline  tubocurarine
rescinnamine  thiamylal  urea
reserpine  thiopental  urokinase
seco barbital  timolol  vinbarbital
sodium salicylate  tolazoline  vitamin A
spironolactone  trichlormethiazide

10. Detachment of the ciliary body, planned or inadvertent
11. Hyperosmotic agents, such as mannitol or urea
12. Iritis or iridocyclitis
13. After central retinal vein occlusion
14. Myopia—low scleral rigidity may give false low readings with Schiötz tonometer, but normal readings with applanation intraocular pressure
15. Herpes zoster
16. Following irradiation by roentgenograms or beta rays
17. Congenital lesions, including microphthalmos, aniridia, and coloboma
18. Concussion trauma
19. Necrosis of anterior segment of the eye
20. Idiopathic, including normal variation


PHTHISIS BULBI (DEGENERATIVE SHRINKAGE OF EYEBALL WITH HYPOTONY)

1. Cilioodestructive procedures such as cyclocryotherapy or laser
2. Endophthalmitis
3. Following cataract surgery, especially with rubella syndrome (German measles)
4. Panophthalmitis
5. Severe ocular injury with loss of tissue
6. Severe uveitis
7. Sympathetic ophthalmia
8. Tumor, such as retinoblastoma or malignant melanoma


**CLINICAL ANOPHTHALMOS (APPARENT ABSENCE OF GLOBE)**
1. Anencephaly
2. Gross midline facial defects (median cleft face syndrome)
3. Dyscraniopygophalangea
4. Goldenhar syndrome (oculoauriculovertebral syndrome)
5. Goltz syndrome (focal dermal hypoplasia syndrome)
6. Hallermann-Streiff syndrome (dyscephalic mandibulo-oculofacial syndrome)
7. Hypervitaminosis A
8. Idiopathic
9. Klinefelter syndrome (gynecomastia-aspermatogenesis)
10. Lanzieri syndrome (craniofacial malformations)
11. Leri syndrome (carpal tunnel syndrome)
12. Meckel syndrome (dysencephalia splanchnocyntica syndrome)
13. Oculovertebral dysplasia (Weyers-Thier syndrome)
14. Otocephaly
15. Trisomy 13-15
16. Sex-linked or recessive hereditary
17. Waardenburg anophthalmia syndrome (anophthalmos with limb anomalies)-recessive


**OCULODIGITAL STIMULATION**

The patient presses on the globe through the lids with the index finger or hand; the patient has poor visual acuity.

1. Bilateral congenital cataracts
2. Combined retinal detachment and congenital cataract
3. Congenital glaucoma
4. Congenital rubella syndrome (German measles)
5. Leber amaurosis congenita or other congenital retinal degeneration (Leber tapetoretinal dystrophy syndrome)
6. Norrie disease (fetal iritis syndrome)
7. Total corneal leukoma

ANTERIOR SEGMENT ISCHEMIA

This condition involves hypoxia with involvement of the cornea, iris, anterior chamber, lens, and ciliary body.

1. Damage to normal intact anterior vessels
   A. Pressure
      (1) Scleral buckle
      (2) Suture (Jensen procedure)
   B. Thermal
      (1) Cryotherapy
      (2) Diathermy

2. Disinsertion of normal vessels (Hummelsheim or Knapp procedure)

3. Fuchs syndrome (I) (heterochromic cyclitis syndrome)

4. Hematologic abnormality
   A. Extreme leukocytosis
   B. Extreme thrombocytosis
   C. Hyperglobulinemia
   D. Red blood cell dysfunction including sickle cell trait
      (1) Hemoglobinopathy
      (2) Polycythemia vera (Vaquez-Osler syndrome)

5. Vessel wall abnormality (arteriosclerosis)
   A. Arteriosclerosis
   B. Giant cell arteritis


BLUE SCLERA

Blue sclera is characterized by localized or generalized blue coloration of sclera because of thinness and loss of water content, which allow underlying dark choroid to be seen.

1. Associated with high urine excretion
   A. Folling syndrome (phenylketonuria)
   B. Hypophosphatasia (phosphoethanolaminuria)
   C. Lowe syndrome (oculocerebrorenal syndrome; chondroitin-4-sulfate-uria)
2. Associated with skeletal disorders
   A. Brachmann-de Lange syndrome
   B. Brittle cornea syndrome (blue sclera syndrome)-recessive
   C. Crouzon disease (craniofacial dysostosis)
   D. Hallermann-Streiff syndrome (dysecephalia mandibulooculofacial syndrome)
   *E. Marfan syndrome (dystrophia mesodermalis congenita)
   F. Marshall-Smith syndrome
   G. McCune-Albright syndrome (fibrous dysplasia)
   H. Mucopolysaccharidosis VI (Maroteaux-Lamy syndrome)
   I. Osteogenesis imperfecta (van der Hoeve syndrome)
   J. Paget syndrome (osteitis deformans)
   K. Pierre Robin syndrome (micrognathia-glossoptosis syndrome)
   L. Robert syndrome
   M. Silver-Russell syndrome
   N. Werner syndrome (progeria of adults)
3. Chromosome disorders
   A. Trisomy syndrome
   B. Turner syndrome
4. Ocular
   *A. Congenital glaucoma
   B. Myopia
   *C. Repeated surgeries
D. Scleromalacia (perforans)
E. Staphyloma
F. Trauma

5. Miscellaneous
   A. Ehlers-Danlos syndrome (fibrodysplasia elastica generalisata)
   B. Goltz syndrome (focal dermal hypoplasia syndrome)
   C. Incontinentia pigmenti (Bloch-Sulzberger syndrome)
   D. Lax ligament syndrome
   E. Minocycline-induced
   F. Oculodermal melanocytosis (nevus of Ota)
   G. Pseudoxanthoma elasticum (Grönblad-Strandberg syndrome)
   H. Relapsing polychondritis


**Extracted Table Blue Sclera**

**DILATED EPISCLERAL VESSELS**

1. Carotid-cavernous fistula
2. Cavernous sinus thrombosis (Foix syndrome)
*3. Chronic respiratory diseases
*4. Glaucoma, untreated
5. Increased viscosity of circulating blood
   A. Leukemia (early)
   B. Polycythemia vera (erythema, Vaquez-Osler syndrome)
6. Occlusion of orbital veins of the apex of the orbit
   A. Endocrine exophthalmos of rapid development
   B. Inflammatory lesions
   C. Orbital thrombophlebitis
   D. Tumor (rare)
7. Ophthalmic vein thrombosis
8. Tricuspid Incompetence
9. Uveal neoplasm with localized engorgement


**EPISCLERITIS**

Episcleritis is a benign, self-limited, nodular or diffuse disease that usually resolves spontaneously within weeks but has a tendency to recur. Inflammation of episcleral tissues causes discomfort rather than pain; it does not affect visual acuity. Even recurrent attacks do not produce scleritis. Complications are minimal and include areas of scleral transparency and localized keratitis.

*1. Idiopathic (single, short episode that does not recur)*

2. Associated with the following diseases (recurrent attacks)
   A. Addison syndrome (adrenal cortical insufficiency)
   B. Arthritis
      (1) Involving small and medium-sized vessels
         a. Necrotizing granulomatous arthritis; Wegener granulomatosis (Wegener syndrome)
         b. Polyarteritis nodosa (Kussmaul disease)
      (2) Involving small, medium, and large vessels
         a. Arteritis in collagen vascular diseases
            (i) Progressive systemic sclerosis (PSS; scleroderma)
            (ii) Rheumatoid arthritis
            (iii) Rheumatic fever
   C. Cogan syndrome
   D. Crohn disease (granulomatous ileocolitis)
   E. Goodpasture syndrome (pulmonary hemosiderosis)
   F. Heerfordt disease (uveoparotid fever)
   G. Inflammatory pseudotumor
   H. Initial manifestation of uveal melanoma (ciliary body)
   I. Myeloproliferative diseases
      (1) Hodgkin disease
      (2) T-cell leukemia
   J. Paraneoplastic syndromes
      (1) Dermatomyositis
      (2) Sweet syndrome (cutaneous paraneoplastic syndrome)
   K. Paraproteinemia
      (1) Familial Mediterranean fever
      (2) Necrobiotic xanthogranuloma [increased immunoglobulin G (IgG)/IgA]
   L. Parry-Romberg syndrome (progressive hemifacial atrophy)
   M. Relapsing polychondritis
   N. Skin diseases
      (1) Chronic cutaneous lupus erythematosus (CCLE)
      (2) Erythema elevatum diutinum
      (3) Lichen planus
      (4) PSS; scleroderma
(5) Psoriasis
(6) Reiter syndrome (polyarteritis enterica)
(7) Wiskott-Aldrich syndrome

O. Terrien marginal corneal disease
P. Ulcerative colitis (regional enteritis)
Q. Weber-Christian disease (systemic panniculitis)
R. Pseudoepiscleritis (lesions resembling episcleritis)
   (1) Conjunctivitis
   (2) In-growing lash
   *(3) Inflamed pinguecula
   (4) Punctate keratitis
   (5) Sclerosing keratitis
   (6) Wegener granulomatosis

3. Drugs
   A. Pamidronate disodium

4. Infectious
   A. Brucellosis (Bang disease, undulant fever)
   B. Coccidioidomycosis
   C. Influenza
   D. Leprosy (Hansen disease)
   E. Leptospirosis (Weil disease)
   F. Lyme disease (borreliosis, relapsing fever)
   G. Lymphogranuloma venereum (Nichols-Favre disease)
   H. Nematode (Angiostrongylus cantonensis)
   I. Q fever

5. Trauma
   *A. Episcleral foreign body
   B. Following transscleral fixation of posterior chamber IOL (intraocular lens)
   C. Insect bite granuloma
   D. Malpositioned (Jones) tube


**PIGMENT SPOTS OF SCLERA AND EPISCLERA**

*1. Acquired melanosis
2. Cysts
3. Drugs, including the following:
acetophenazine  butaperazine  carphenazine  chlorpromazine  diethazine  ethopropazine  ferrocholinate  ferrous fumarate  ferrous gluconate  ferrous succinate  ferrous sulfate  fluphenazine  iron dextran  mesoridazine  methdilazine  methotrimeprazine  perazine  pericyzine  perphenazine  piperacetazine  polysaccharide iron complex  prochlorperazine  promazine  propiomazine  thiethylperazine  thiopropazate  thioproperazine  thioridazine  trifluoperazine  triflupromazine  trimeprazine  vitamin D  vitamin D$_2$  vitamin D$_3$

4. Extension of adjacent or underlying malignant melanoma
5. Foreign body
6. Intrascleral nerve loops with uveal pigment (painful to touch)
   *7. Nevus
8. Ochronosis with melanin deposition
   *9. Resolving hemorrhage
10. Staphyloma
11. Transscleral migration of pigment following cryotherapy of intraocular tumor or trauma
12. Uveal melanocytes carried by the scleral emissaria into the episclera (most often in eyes, with dark irides in superior, inferior temporal, and nasal quadrants in descending frequency; conjunctiva freely movable over them)


**SCLERITIS**

Scleritis involves a potentially destructive inflammatory process that may accompany severe systemic disease. Ocular pain occasionally radiates to the temple, jaw, or sinuses. Women are more frequently affected than men. Most cases begin with bilateral involvement. Early perforation of sclera is possible. The anterior portion of the eye is affected most severely. Posterior scleritis may be a diagnostic challenge.

1. Associated with systemic disease
   A. Collagen diseases
      *(1) Dermatomyositis (Wagner-Unverricht syndrome)
      *(2) Felty syndrome
(3) Giant cell (temporal) arteritis  
(4) Juvenile rheumatoid arthritis (Still disease)  
(5) Polyarteritis nodosa (Kussmaul disease)  
(6) PSS; scleroderma  
(7) Relapsing polychondritis  
(8) Reiter syndrome (polyarteritis enteric a)  
(9) Rheumatoid arthritis  
(10) Sjögren syndrome  
(11) Systemic lupus erythematosus (SLE)  
(12) Wegener granulomatosis (Wegener syndrome)

B. Metabolic diseases  
(1) Cretinism (hypothyroidism)  
(2) Gout  
(3) Porphyria cutanea tarda

C. Myeloproliferative diseases  
(1) Hodgkin disease (lymph node disease)  
(2) Mycosis fungoides syndrome (Sézary syndrome)

2. Infectious  
A. Bacterial  
(1) Leprosy  
(2) Lymphogranuloma venereum (Nichols-Favre disease)  
(3) Syphilis (acquired lues)  
(4) Tuberculosis

B. Viral infections  
(1) Herpes simplex  
(2) Herpes zoster  
(3) Influenza  
(4) Mumps

C. Fungal-aspergillosis

D. Helminth infection- acanthamoeba

E. Protozoan-toxoplasmosis

F. Infections  
(1) Associated with skin disease or immunosuppressive status  
(2) Spreading directly from conjunctiva, cornea, uvea, periorbital tissues, nose, or sinuses

3. Miscellaneous  
A. Cogan syndrome  
B. Crohn disease (granulomatous ileocolitis)  
C. Goodpasture syndrome (pulmonary hemosiderosis)  
D. Erythema nodosum  
E. Exogenous infection via penetration through conjunctiva  
F. Hashimoto thyroiditis  
G. Heerfordt disease (uveoparotid fever)  
H. Necrobiotic xanthogranuloma (increased IgG/IgA)  
I. Terrien marginal corneal disease  
J. Uveitis
4. Drugs
   A. Pamidronate disodium

*5. Trauma-following cataract or strabismus surgery


**STAPHYLOMA (STRETCHING AND THINNING OF THE SCLERA WITH INCARCERATION OF UVEAL TISSUE)**

1. Collagen diseases
   A. Felty syndrome
   B. Rheumatoid arthritis (adult)
   C. Wegener syndrome (Wegener granulomatosis)

2. Following trauma
   A. Beta radiation
   B. Deep scleral resection for episcleral malignancies
   C. Pterygium excision and mitomycin therapy
   *D. Scleral buckle removal
   E. Subconjunctival injection of corticosteroids
   F. Ultrasound treatment for glaucoma

3. Infectious
   A. Aspergillosis
   B. Herpes zoster (rare)
   C. Plague (bubonic plague)
   D. Syphilis
   E. Tuberculosis

4. Ocular cause
   A. Buphthalmos associated with increased intraocular pressure
   B. Comeoscleral ectasia
   C. Myopia with increased anteroposterior diameter
D. Scleritis (e.g., secondary to rheumatoid arthritis)
E. Uveitis

5. Miscellaneous
   A. Ehlers-Danlos syndrome (fibrodysplasia elastica generalisata)
   B. Endarteritis
   C. Epidermolysis bullosa
   D. Hyperparathyroidism
   E. Meckel syndrome (dysencephalia syndrome)
   F. Oculodental syndrome (Peter syndrome)
   G. Porphyria cutanea tarda


**EPISCLERAL AND SCLERAL TUMORS**

1. Carcinomas
2. Choroidal melanomas
   *3. Epibulbar tumor
4. Fibromas
5. Hemangiomas
6. Histiocytosis
7. Lymphomas
8. Leiomyoma (transscleral)
9. Melanoblastoma (spread from choroid)
10. Retinoblastoma


8

Cornea

CONTENTS

Crystals of the cornea 242
  Diagnostic table 244
Anesthesia of the cornea 246
Hyperplastic corneal nerves 247
Increased visibility of corneal nerves 248
Pigmentation of the cornea 248
Corneal edema 250
Corneal hydrops 252
Microcornea 252
  Diagnostic table 254
Megalocornea 255
  Diagnostic table 256
Corneal opacification in infancy 258
Band-shaped keratopathy 259
Corneal keloids 260
Punctate keratitis or keratopathy 260
Morphologic classification of punctate corneal lesions 265
Sicca keratitis 266
White rings of the cornea (Coats disease) 267
Dry spots of the cornea 267
Anterior embryotoxon (Arcus) 267
Bowman membrane folds 268
  Diagnostic table 269
Delayed corneal wound healing 270
Anterior corneal mosaic 270
Linear opacity in superficial corneal stroma 270
Superficial vertical corneal striations 271
Dendritic corneal lesions 271
Bullous keratopathy 271
Nummular keratitis 272
Deep keratitis 272
Interstitial keratitis 272
Pannus 274
  Diagnostic table 276
Corneal opacity-diffuse 278
Corneal opacity-localized, diffuse 278
Deep corneal stromal deposits 279
Intracorneal hemorrhage 279
Central posterior stromal corneal deposits 279
Dellen 280
Phlyctenular keratoconjunctivitis 280
Corneal ring lesion 280
Cryoglobulinemia
B. Multiple myeloma
E. Elevated bilirubin with crystalline dystrophy
F. Fine, multicolored glittering crystals following successful transplant that later underwent graft rejection and was treated with steroids
G. Gout (hyperuricemia)
12. Hyperparathyroidism
13. Immunoglobulin G (IgG) K monoclonal gammopathy
*14. Infectious crystalline retinopathy, usually with more indolent streptococcal and staphylococcal species
15. Post keratoplasty (Kaye dots)
16. Renal failure
17. Subconjunctival 5-fluorouracil
18. Uremia
19. Waldenström syndrome (macroglobulinemia syndrome)


**Extracted Table Crystals of the cornea (deposition of crystalline subsyance in the cornea)**

**ANESTHESIA OF THE CORNEA (HYPESTHESIA OR DIMINISHED CORNEAL SENSATION IN TRIGEMINAL DISTRIBUTION)**

1. Cornea  
   *A. Cerebellopontine angle tumors  
   B. Congenital  
   C. Corneal dystrophy-granular, lattice, macular, and crystalline  
   D. Dysautonomia  
   *E. Infections, including herpes zoster, herpes simplex, leprosy, and malaria  
   F. Inflammations, including that occurring after electrocautery of Bowman membrane, stromal edema, vascularized scars, congestive glaucoma, exposure keratitis, radiation damage, and vitamin A deficiency  
   G. Trauma, including constant wearing of contact lenses and postoperatively, including cataract extraction and within corneal transplant, following operation for detached retina-from an encircling band or, less frequently, a circumscribed buckle; from refractive surgery  

2. Maxillary division
A. Congenital
B. Facial trauma
C. Interruption of trigeminal nerve or gasserian ganglion, including
cerebellopontine angle tumor or other space-occupying lesion in the region of the
superior orbital fissure
D. Maxillary antrum carcinoma
E. Neoplasm, foramen rotundum, sphenopterygoid fossa
F. Orbital floor fracture
G. Perineural spread of skin carcinoma
H. Surgery for trigeminal neuralgia

3. Ophthalmic division
   A. Aneurysm, cavernous sinus
   B. Neoplasm, cavernous sinus
   C. Neoplasm, middle fossa
   D. Neoplasm, orbital apex
   E. Neoplasm, superior orbital fissure

4. Syndromes and diseases
   A. Adie syndrome
   B. Anhidrotic ectodermal dysplasia
   C. Barré Lieou syndrome (posterior cervical sympathetic syndrome)
   D. Diabetes mellitus-youth onset, more marked with age
   E. Eaton-Lambert syndrome (myasthenic syndrome)
   F. Familial corneal hypesthesia
   G. Foix syndrome (cavernous sinus syndrome)
   H. Gradenigo syndrome (temporal syndrome)
   I. Hereditary fleck dystrophy of the cornea
   J. Herpes zoster
   K. Hunt syndrome (herpes zoster auricularis)
   L. Hydroa vacciniforme (lower cornea)
   M. Multiple sclerosis
   N. Oculoauriculovertebral dysplasia (Goldenhar-Gorlin syndrome)
   O. Nephropathic cystinosis
   P. Passow syndrome (Bremer status dysraphicus)
   Q. Psoriasis (lower cornea)
   R. Riley-Day syndrome (congenital familial dysautonomia)
   S. Rochon-Duvigneaud syndrome (superior orbital fissure syndrome)
   T. Rollet syndrome (orbital apex-sphenoidal syndrome)
   U. Scholz subacute cerebral sclerosis (arylsulfatase A deficiency syndrome)
   V. Temporal arteritis syndrome (cranial arteritis syndrome)
   W. Tolosa-Hunt syndrome (painful ophthalmoplegia)
   X. Vitamin A deficiency

5. Toxins and drugs, including oleoresin capsicum (pepper spray)
   amiodarone betaxolol carisoprodol
   amitriptyline bromide chloroquine
   amodiaquine carbon dioxide clorazepate
   atenolol (?) carbon disulfide desipramine
<table>
<thead>
<tr>
<th>Diazepam</th>
<th>Methyprylon</th>
<th>Stealer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gentamicin (?)</td>
<td>Metoprolol (?)</td>
<td>Timolol</td>
</tr>
<tr>
<td>Glutethimide</td>
<td>Nadolol (?)</td>
<td>Trichloroethylene</td>
</tr>
<tr>
<td>Hydrogen sulfide</td>
<td>Nortriptyline</td>
<td>Trifluoperazine</td>
</tr>
<tr>
<td>Hydroxychloroquine</td>
<td>Paraldehyde</td>
<td>(Stelazine)</td>
</tr>
<tr>
<td>Imipramine</td>
<td>Phencyclidine</td>
<td>Vinblastine</td>
</tr>
<tr>
<td>Levobunolol</td>
<td>Pindolol (?)</td>
<td>Vincristine</td>
</tr>
<tr>
<td>Meprobamate</td>
<td>Propanolol</td>
<td></td>
</tr>
</tbody>
</table>


**HYPERPLASTIC CORNEAL NERVES**

This condition involves overgrowth of corneal nerves up to 20 times the normal number. This nonspecific change may occur in association with the following conditions:

1. Deep filiform dystrophy of Maeder and Danis
2. Herpes simplex
3. Herpes zoster
4. Multiple endocrine neoplasia - type II B
5. Neurofibromatosis (von Recklinghausen syndrome)
6. Neuroparalytic keratitis
7. Normal eyes at advanced age
8. Ocular pemphigus foliaceus (Cazenave disease)
9. Opaque corneal grafts
10. Phthisis bulbi
11. Posterior polymorphous dystrophy


INCREASED VISIBILITY OF CORNEAL NERVES
1. "Colloidin" skin syndrome (bullous ichthyosiform erythroderma)
2. Congenital
3. Ectodermal dysplasia (Rothmund syndrome)
4. Fuchs dystrophy
5. Ichthyosis
6. Idiopathic
7. Keratoconus
8. Leprosy (Hansen disease)
9. Neurofibromatosis (von Recklinghausen syndrome)
10. Neurofibromatosis associated with pheochromocytoma and thyroid carcinoma (Sipple syndrome)
11. Posterior polymorphous dystrophy
12. Primary amyloidosis
13. Refsum syndrome (phytanic acid storage disease)
14. Siemens disease (keratosis follicularis spinulosa decalvans)


PIGMENTATION OF THE CORNEA
1. Melanin pigmentation
   A. Epithelial melanosis
      (1) Congenital
      (2) Presence of limbal malignant melanoma
      (3) Sequela of trachoma and other inflammations
      (4) Melanocytic migration in heavily pigmented persons
   B. Stromal pigmentation such as that in ochronosis
   C. Endothelial melanosis
      (1) Congenital
      (2) Senile
      (3) Degenerative, including atrophic and inflammatory conditions (such as cornea guttata, herpes simplex, zoster keratitis, myopia, diabetes mellitus, senile cataract, chronic glaucoma, and melanoma)
      *(4) Krukenberg spindle, with or without pigmentary glaucoma, may be present in association with diabetes mellitus
      (5) Trauma—from contusions, wounds, or intraocular operations
      (6) Turks line—fine vertical line in the lower portion of the cornea
2. Hematogenous pigmentation
   *(A. Blood staining of the cornea, most often because of total hyphema associated with elevated intraocular pressure
B. Hemorrhage into cornea-following subconjunctival hemorrhage and intracorneal hemorrhage from newly formed vessels, as in interstitial keratitis or mustard gas keratitis
C. Epithelial deposit associated with spherocytic anemia
D. Hemachromatosis

3. Metallic pigmentation
A. Copper (chalcosis)
   *(1) Kayser-Fleischer ring-limbal ring associated with Wilson disease
   (2) Copper foreign body in cornea or intraocular region
   (3) Occupational exposure or topical therapeutic use of copper-containing substance
   (4) Advanced cirrhosis of the liver, such as that associated with parasitic infestation (schistosomiasis)
B. Silver (argyrosis)-from topical, local, or systemic use; also occupational use
C. Gold (chrysiasis)-from topical, local, or systemic use
D. Iron (siderosis)
   (1) Foreign body in cornea or intraocular area
   (2) Iron lines
      *a. Fleischer ring-associated with keratoconus around base of the cone
      b. Hudson-Stähli line-horizontal line at the junction of the middle and lower one third of the cornea, believed to be related to exposure, trauma of lid closure, and chronic corneal infection
      c. Stocker line-line running parallel with head of the pterygium
      d. Ferry line-associated with filtering blebs, believed to result from minute, repeated, localized trauma caused by eyelid striking the elevated bleb
      e. Circular lesion associated with congenital spherocytosis
      f. Iron lines following refractive corneal surgery, such as radial keratotomy and photorefractive keratectomy, and laser in situ keratomileusis (hyperopic and myopic)
E. Bismuth (bismuthiasis)-from therapeutic use
F. Arsenic melanosis

4. Drugs, discoloration, including the following:
   acetophenazine  auranofin  bismuth subsalicylate
   acid bismuth sodium  aurothioglucose  butaperazine
      tartrate  aurothioglycanide  calcitriol
   alcohol  bismuth carbonate  carphenazine
   amiodarone  bismuth oxychloride  chloroquine
   amodiaquine  bismuth salicylate  chlorpromazine
   antimony potassium tartrate  bismuth sodium tartrate  chlorprothixene
   antimony sodium tartrate  bismuth sodium  chlortetracycline
   antimony sodium thioglycollate  colloidal silver  diethazine
   thioglycollate  bismuth subcarbonate  echothiophate
   bismuth carbonate  bismuth subsalicylate  epinephrine
   bismuth oxychloride  bismuth sodium thioglycollate  echothiophate
   bismuth salicylate  bismuth sodium thioglycollate  epinephrine
   bismuth sodium tartrate  bismuth subcarbonate  epinephrine
ergocalciferol  methotrimetrazine  sodium
ethopropazine  methylene blue  antimonylgluconate
ferrocholinate  mild silver protein  stibocaptate
ferrous fumarate  perazine  stibogluconate
ferrous gluconate  perhexiline  stibophen
ferrous succinate  perichryzaene  tetracycline
ferrous sulfate  perphenazine  thiethylperazine
fluphenazine  phenylmercuric nitrate  thimerosal
gold Au 198  piperaacetazine  thiopropazate
gold sodium thiomalate  polysaccharide-iron  thioridazine
gold sodium thiosulfate  complex  thioridazine
hydroxychloroquine  prazosin  thiopropazate
indomethacin  prochlorperazine  trifluoperazine
iodide  promazine  triflupromazine
iodine solution  promethazine  trimethazine
iron dextran  propiomazine  vitamin A (?)
iron sorbitex  quinacrine  vitamin D
meperidine (?)  quinidine  vitamin D2
mercuric oxide  radioactive solution  vitamin D3
mesoridazine  silver nitrate
methililazine  silver protein

5. Other color changes
   A. White discoloration-scars, fatty degeneration or infiltration, calcified areas
   B. Yellow, discoloration-hyaline or colloid degeneration, and Tangier disease
      (familial deficiency of high-density lipoprotein)
   C. Black discoloration-coal powder, dirt, epinephrine, or ink (tattooing)
   D. Yellow-brown discoloration-Kyrle disease (hyperkeratosis follicularis et
      parafollicularis in cutem penetrans)
   E. Grey-black discoloration-chronic phenol exposure as carbolic acid
   F. Grey-white discoloration-anesthetic cornea
   G. Brown discoloration-aniline (amidobenzole), including benzoquinone and
      hydroquinine


**CORNEAL EDEMA**

1. Drugs, including the following:
   - acetophenazine  
   - acetylcholine  
   - alpha-chymotrypsin  
   - amodiaquine  
   - amphotericin B  
   - bacitracin  
   - benoxinate  
   - benzalkonium chloride  
   - benzathine penicillin G  
   - butacaine  
   - butaperazine  
   - carbachol  
   - carphazine  
   - chloramphenicol  
   - chlorhexidine  
   - chloroquine  
   - chlorpromazine  
   - chlorotetracycline  
   - cocaine  
   - colistin  
   - deslanoside  
   - dibucaine  
   - diethazine  
   - digitoxin  
   - digoxin  
   - dyclonine  
   - epinephrine  
   - erythromycin  
   - ethopropazine  
   - fluphenazine  
   - hydrobamine penicillin  
   - hydrogen peroxide  
   - hydroxychloroquine  
   - idoxuridine  
   - lanatoside C  
   - melphalan  
   - methdilazine  
   - methicillin  
   - methotrimazeprene  
   - neomycin  
   - perazine  
   - pericyazine  
   - perphenazine  
   - phenacaine  
   - phenoxymethylpenicillin  
   - phenylephrine  
   - piperacetazine  
   - piperocaine  
   - polymyxin B  
   - potassium penicillin G  
   - potassium penicillin V  
   - potassium phenethicillin  
   - potassium phenoxymethyl  
   - procaine penicillin G  
   - prochlorperazine  
   - promazine  
   - propantheline  
   - propiomazine  
   - quinacrine  
   - silicone  
   - streptomycin  
   - tetracycline  
   - thiethylperazine  
   - thiopropazate  
   - thiopropazine  
   - thiopropazine  
   - thiostatazide  
   - thiotope  
   - trifluoperazine  
   - triflupromazine  
   - trifluoride  
   - urokinase  
   - vidarabine  
   - vinblastine

2. Endothelial decompensation
   A. Noninflammatory
      *(1) Acute hydrops with keratoconus
      (2) Congenital
         a. Anhidrotic ectodermal dysplasia
         b. Birth trauma, typically a forceps injury
         c. Congenital glaucoma
         d. Congenital hereditary endothelial dystrophy
         e. Posterior polymorphous dystrophy
      (3) Environmental cold in trigeminal nerve palsy
(4) Essential corneal edema
(5) Failed corneal graft
(6) Metabolic such as myxedema and hypercholesteremia
(7) Neuropathic conditions
(8) Postsurgical
  a. Anterior segment ischemia
  b. Anterior synechiae
  c. Direct mechanical damage to endothelium including argon laser
     iridotomy
  d. Epithelial or fibrous downgrowth
  e. Osmotic, such as irrigation of cornea or anterior chamber with distilled water
  f. Plasma gas sterilization
  g. Stripped Descemet membrane
  h. Vitreous touch
*(9) Primary degenerative-Fuchs dystrophy
(10) Traumatic
  a. Anoxia of epithelium, such as from excessive wearing of contact lens (Sattler veil)
  b. Brown-McLean Syndrome
  c. Chemical, such as tear gas, hydrogen peroxide and Hibiclens
  d. Exposure as in exophthalmos
  e. Large epithelial defect
  f. Nonpenetrating including after air bag inflation injury
  g. Penetrating
  h. Radiation injury such as from ultraviolet, roentgenograms, gamma rays
  i. Retained foreign body-anterior chamber
  j. Sympathectomy including jugular vein catheterization and Parry-Robson syndrome
  k. Trigeminal nerve palsy with cold exposure

B. Inflammatory
  (1) Any severe iritis
  (2) Acute graft rejection
  (3) Chandler syndrome (iridocorneal endothelial syndrome)
  *(4) Herpes simplex keratitis or keratouveitis
  *(5) Herpes zoster keratouveitis
  (6) Retinal tacks

3. Increased intraocular pressure
  *A. Acute glaucoma
  B. Chronic glaucoma
    (1) Minimal to moderate pressure elevations in the presence of abnormal endothelium
    (2) Prolonged moderately high elevations in the presence of normal or near-normal endothelium

4. Hypotony


**CORNEAL HYDROPS (RUPTURES OF DESCEMET MEMBRANE WITH CORNEA INTRALAMELLAR DISSECTION AND COLLECTION OF AQUEOUS HUMOR)**

1. Congenital glaucoma
2. Forceps injury
*3. Keratoconus
4. Pellucid marginal degeneration
5. Terrien marginal degeneration
6. Trauma, blunt


**MICROCORNEA (CORNEA WITH A HORIZONTAL DIAMETER OF LESS THAN 10 MM)**

1. Associated ocular findings
   A. Aniridia and subluxated lenses
B. Autosomal-dominant cataract and myopia
C. Autosomal-dominant cataract, nystagmus, and glaucoma
D. Axenfeld syndrome (posterior embryotoxon)
E. Colobomatous microphthalmia
F. Congenital glaucoma
G. Corectopia and macular hypoplasia
H. Hyperopia
I. Meckel syndrome (dysencephalia splanchnocystica syndrome)
J. Nanophthalmos
K. Narrow-angle glaucoma
L. Sclerocornea

2. Aberfeld syndrome (congenital blepharophimosis associated with generalized myopathy)
3. Autosomal recessive or dominant trait
4. Carpenter syndrome (acrocephalopolysyndactyly II)
5. Cataract microcornea syndrome
6. Chromosome partial deletion (long-arm) syndrome
7. Deafness retardation, arched palate syndrome
8. Ehlers-Danlos syndrome (fibrodyplasia elastica generalisata)
9. Gansslen syndrome (familial hemolytic icterus)
10. Hallermann-Streiff syndrome (dyscephalic mandibulooculofacial syndrome)
11. Hemifacial microsomia syndrome (Francois Haustrate syndrome)
12. Hutchinson-Gilford syndrome (progeria)
13. Laurence-Moon-Biedl syndrome (retinitis pigmentosa-polydactylyadiposogenital syndrome)
14. Lenz microphthalmia syndrome
15. Little syndrome (nail patella syndrome)
16. Marchesani syndrome (mesodermal dysmorphodystrophy)
17. Marfan syndrome (arachnodactyly dystrophica mesodermalis congenita)
18. Meckel syndrome (dysencephalia splanchnocystica syndrome)
19. Meyer-Schwickerath-Weyers syndrome (oculodentodigital dysplasia)
20. Microcornea, glaucoma, absent frontal sinuses
21. Micro syndrome
22. Rieger syndrome (hypodontia and iris dysgenesis)
23. Ring chromosome
24. Roberts pseudothalidomide syndrome
25. Rubella syndrome (Gregg syndrome)
26. Sabin-Feldman syndrome
27. Schwartz syndrome (glaucoma associated with retinal detachment)
28. Smith-Magenis syndrome
29. Triploidy
30. Trisomy 13 (D trisomy, Patau syndrome)
31. Trisomy syndrome
32. Waardenburg syndrome (interoculoidid Dermatoauditive dysplasia)


**Extracted Table Microcornea**

**MEGALOCORNEA (CORNEA WITH A HORIZONTAL DIAMETER OF MORE THAN 14 MM)**

1. Aarskog syndrome (faciodigitogenital syndrome)
2. Autosomal dominant or recessive trait
3. Congenital glaucoma (rare)
4. Craniosynostosis
5. Down syndrome
6. Facial hemiatrophy
7. Isolated
8. Marchesani syndrome (brachymorphia with spherophakia)
9. Marfan syndrome (arachnodactyly dystrophy mesodermalis congenita)
10. MMR (megalocornea-mental retardation) syndrome
11. MMMM (megalocornea, macrocephaly, mental and motor retardation) syndrome
12. Mucopolysaccharidoses I-S (Scheie syndrome)
13. Neuhauser syndrome (megalocornea-mental retardation syndrome)
14. Oculocerebrorenal syndrome (Lowe syndrome)
15. Oculodental syndrome (Peter syndrome)
16. Osteogenesis imperfecta (van der Hoeve syndrome)
17. Oxycephaly (dysostosis craniofacial dyostosis)
18. Pierre Robin syndrome (micrognathia-glossoptosis syndrome)
19. Posterior embryotoxon
20. Rieger syndrome (hypodontia and iris syndrome)
21. Rubella syndrome (Gregg syndrome)
22. Sex-linked recessive trait
23. Sturje-Weber syndrome (meningocutaneous syndrome)


Extracted Table Megalocornea

CORNEAL OPAFICATION IN INFANCY (SEE CONDITIONS SIMULATING CONGENITAL GLAUCOMA)

*1. Birth trauma, such as Descemet membrane rupture
2. Chromosomal aberrations
   A. Mongolism (Down syndrome)-trisomy 21
   B. Trisomy 13 (Patau syndrome)
3. Congenital malformations
   A. Amyloidosis (Lubarsch-Pick syndrome)
   B. Anterior chamber cleavage syndromes
      (1) Axenfeld anomaly
      (2) Congenital central anterior synechiae
      (3) Congenital anterior staphyloma
* (4) Peter anomaly
      (5) Rieger anomaly
   C. Bilateral corneal dermis-like choristomas
   D. Congenital glaucoma
   E. De Barsy syndrome
   *F. Dermoid tumors
   *G. Sclerocornea
   H. Xanthomas
4. Corneal dystrophy
   *A. Congenital hereditary endothelial dystrophy
   B. Congenital hereditary stromal dystrophy
   C. Posterior polymorphous dystrophy
5. Idiopathic
6. Inborn errors of metabolism
   *A. Mucopolysaccharidoses (MPS)
      (1) Hurler syndrome (MPS IN)
      (2) Maroteaux-Lamy syndrome (MPS VI)
      (3) Morquio-Brailsford syndrome (MPS IV)
      (4) Scheie syndrome (MPS IS)
   B. Lowe syndrome (oculocerebrorenal syndrome)
   C. von Gierke disease (glycogen disease)
D. Corneal lipoidosis-later
E. Mucolipidosis
   (1) Generalized gangliosidosis (GMI-gangliosidosis I and II)
   (2) ML I (lipomucopolysaccharidosis)
   (3) ML III (pseudo-Hurler polydystrophy)
F. Riley-Day syndrome (congenital familial dysautonomia)

7. Inflammatory processes
   A. Corneal ulceration
   B. Herpes simplex and herpes zoster
   C. Interstitial keratitis
   D. Rubella syndrome (German measles)
   E. Savin syndrome


**BAND-SHAPED KERATOPATHY**

This type of corneal opacification extends horizontally over the cornea, at the level of the Bowman membrane, in the exposed part of the palpebral aperture.

1. Anterior mosaic dystrophy, primary type
   A. Episkopi (sex-linked recessive)
   B. Labrador keratopathy
2. Chemical fume related as mercury vapor or calcium bichromate vapor
3. Cyclosporine as eyedrops
4. De Barsy syndrome
5. Discoid lupus erythematosus
6. Dysproteinemia
7. Gout (hyperuricemia)
8. High levels of visible electromagnetic radiation, such as xenon arc photocoagulation and laser causing acute severe anterior uveitis
9. Hypercalcemia
   A. Excessive vitamin D as with oral intake, Boeck sarcoid with liver involvement, acute osteoporosis, Heerfordt syndrome, and Schaumann syndrome
   B. Hyperparathyroidism
   C. Hypophosphatasia (phosphoethanolaminuria)
   D. Idiopathic hypercalcemia
   E. Milk-alkali syndrome
   F. Paget syndrome (osteitis deformans)
   G. Renal failure, such as that associated with Fanconi syndrome (cystinosis)
10. Ichthyosis vulgaris
*11. Local degenerative diseases, including chronic uveitis, phthisis bulbi, absolute glaucoma, infantile polyarthritis (Still disease), rheumatoid arthritis, interstitial keratitis, Felty syndrome, and juvenile rheumatoid arthritis
12. Long-term miotic therapy
13. Long-term steroid phosphate preparations
14. Progressive facial hemiatrophy (Parry-Romberg syndrome)
15. Rothmund syndrome (ectodermal syndrome)
16. Silicone oil in anterior chamber
17. Traumatic-chronic exposure to irritants, such as mercury fumes, calomel, calcium bichromate vapor, and hair
18. Tuberous sclerosis (Bourneville syndrome)
19. Tumoral calcinosis
20. Viscoat usage
21. Wagner syndrome (hyaloideoretinal degeneration)
22. X-linked recessive ocular dystrophy


**CORNEAL KEOLOIDS**

1. Lowe syndrome (oculocerebrorenal syndrome)
2. Trauma, usually with perforation of the iris


**PUNCTATE KERATITIS OR KERATOPATHY**

1. Alimentary disorders
A. Mouth
*(1) Dry mouth, as in Sjögren syndrome
(2) Ulcers, such as primary herpes, ocular cicatrical pemphigoid, and erythema multiforme
B. Lower alimentary tract
(1) Ulcerative colitis as in Sjögren disease
(2) Mild colitis, such as that due to an adenovirus
C. Stomach
(1) Indigestion as in Sjögren syndrome and acne rosacea

2. Articular diseases
A. Psoriasis arthropathica
B. Reiter disease (polyarthritis enterica)
C. Riley-Day syndrome (congenital familial dysautonomia)
D. Sjögren syndrome (secretoinhibitor syndrome)

3. Conjunctival discharge
A. Mucoid
*(1) Ocular cicatricial pemphigoid
(2) Ulcers, such as primary herpes, ocular cicatrical pemphigoid, and erythema multiforme
B. Mucopurulent (see p. 186)
(1) Angular blepharoconjunctivitis
(2) Ulcers, such as primary herpes, ocular cicatrical pemphigoid, and erythema multiforme
(3) Gonococcal
(4) Inclusion conjunctivitis (acute stage)
(5) Meningococcal
(6) Reiter disease (polyarthritis enterica)
(7) Trachoma
(8) Vernal conjunctivitis
C. Serous
(1) Adenovirus
(2) Herpes simplex
(3) Herpes zoster
(4) Inclusion conjunctivitis (later)
(5) Molluscum contagiosum
(6) Trachoma (later)
(7) Warts

4. Conjunctival inflammation
A. Cicatizing (see p. 194)
*(1) Ocular cicatricial pemphigoid
(2) Ulcers, such as primary herpes, ocular cicatrical pemphigoid, and erythema multiforme
(3) Erythema multiforme (Stevens-Johnson syndrome)
(4) Diphtheria
(5) Fuchs-Salzmann-Terrien syndrome (allergic reactions from drugs)
(6) Radiation bums
*(7) Sjögren keratoconjunctivitis sicca
(8) Thermal bums
(9) Trachoma
B. Diffuse catarrhal
   (1) Adenovirus
   (2) Bacterial conjunctivitis
   (3) Erythema multiforme (Stevens-Johnson syndrome)
   (4) Onchocerciasis syndrome (river blindness)
   (5) Reiter disease (polyarthitis enterica)
   (6) Superior limbic keratoconjunctivitis
   (7) Vaccinia

C. Follicular (see p. 192, 193)
   (1) Adenovirus
   (2) Herpes simplex
   (3) Herpes zoster
   (4) Inclusion conjunctivitis
   (5) Molluscum contagiosum
   (6) Trachoma

D. Giant papillary, such as in vernal and atopic conjunctivitis, and related to contact users, prosthesis, and exposed sutures

E. Papillary
   (1) Sjögren syndrome (secretoinhibitor syndrome)
   (2) Trachoma

5. Corneal conditions
A. Deep keratitis, disciform or irregular
   *(1) Herpes simplex
   (2) Herpes zoster and other viral diseases
   (3) Corneal dystrophy (e.g., lattice)
   (4) Harlequin syndrome

B. Thinned facets because of previous ulcerative or other lesions
   (1) Acne rosacea (ocular rosacea)
   (2) Erythema multiforme (Stevens-Johnson syndrome)
   (3) Gorlin-Chaudhry-Moss syndrome
   (4) Herpes simplex
   (5) Sjögren keratoconjunctivitis sicca

C. Vascularization
   (1) Acne rosacea (ocular rosacea)
   (2) Molluscum contagiosum
   (3) Ocular cicatricial pemphigoid
   (4) Phlyctenular disease (see p. 214)
   (5) Sjögren keratoconjunctivitis sicca
   (6) Trachoma
   (7) Vaccinia

D. Trauma
   (1) Chemical injury
   (2) Contact lens related
   (3) Foreign body under upper eyelid
   (4) Mild, such as eye rubbing
   (5) Ultraviolet photokeratopathy
E. Thygeson superficial punctate keratitis (SPK)

6. Diseases of the lids
   A. Dermatitis
      (1) Psoriasis
      (2) Seborrheic blepharitis
   B. Ectropion (see p. 78-79)
      (1) Exposure keratopathy
      (2) Neuroparalytic keratopathy
   C. Folliculitis (see p. 94)
      (1) Blepharitis due to *Demodex folliculorum*
      (2) Seborrheic blepharitis
      (3) Staphylococcal blepharitis
   D. Lid retraction (see p. 62-64)
      *(1) Endocrine exophthalmos
      *(2) Exposure keratopathy
   E. Madarosis, such as that associated with leprosy (stiff immobile lids)
   F. Nodules
      (1) Acne rosacea (ocular rosacea)
      (2) Molluscum contagiosum
      (3) Papilloma
      (4) Warts
   G. Trichiasis or entropion with traumatic keratitis
   H. Vesicles or ulcers
      (1) Herpes simplex
      (2) Herpes zoster
      (3) Ocular cicatricial pemphigoid
      (4) Vaccinia
   I. Floppy eyelid syndrome

7. Diseases of the skin associated with punctate keratitis
   A. Acne rosacea (ocular rosacea)
   B. CRST (calcinosis, cutis, Raynaud phenomenon, sclerodactyly, and
telangiectasia syndrome (calcinosis)
   C. Erythema multiforme (Stevens-Johnson syndrome)
   D. Follicular hyperkeratosis of the palms and soles
   E. Hypertrichosis
   F. Ichthyosis
   G. Incontinentia pigmenti
   H. Leprosy (Hansen disease)
   I. Melkersson-Rosenthal syndrome (Melkersson idiopathic fibroedema)
   J. Ocular cicatricial pemphigoid
   K. Psoriasis

8. Genitourinary diseases associated with punctate keratitis
   A. Erythema multiforme (Stevens-Johnson syndrome)
   B. Inclusion blennorrhea
   C. Ocular cicatricial pemphigoid
   D. Reiter disease (polyarthrosis enterica)
9. Keratitis associated with use of drugs, including the following:

- acebutolol
- acetophenazine
- acetyldigitoxin
- acyclovir
- adenine arabinoside
- adrenal cortex injection
- alcohol
- aldosterone
- allopurinol
- amantadine
- amphotericin B
- antazoline
- antipyrine
- aspirin
- atenolol
- auranofin
- aurothioglucose
- aurothioglycanide
- bacitracin
- benoxinate
- benzalkonium
- betamethasone
- betaxolol
- botulinum A toxin
- bromonidine tartrate
- brinzolamide
- brompheniramine
- bupivacaine
- butacaine
- butaperazine
- capcetabine
- carbimazole
- carphenazine
- chlorambucil
- chloramphenicol
- chlorhexidine
- chloroform
- chloroprocaine
- chlorpheniramine
- chlorpromazine
- chlorprothixene
- chlortetracycline
- chrysarobin
- ciprofloxacin
- cocaine
- colchicine
- cortisone
- cycloheximide
- cytarabine
- deslanoside
- dexamethasone
- dextran
- dibucaine
- diclofenac
- diethazine
- diethylcarbamazine
- digitoxin
- dimethindene
- dipivalyl epinephrine (DPE)
- dipivefrin
- dorzolamide
- dyclonine
- emedastine difumarate
- emetine
- epinephrine
- ether
- ethopropazine
- etidocaine
- etretinate
- F3T
- firaxetine hydrochloride
- fluoxuridine
- flumethasone
- fluorometholone
- fluorouracil
- fluphenazine
- fluprednisolone
- flurbiprofen
- fluvoxamine maleate
- framycetin
- gentamicin
- gitalin
- gold Au 198
- gold sodium thiomalate
- gold sodium thiosulfate
- guanethidine
- hexachlorophene
- hydrocortisone
- idoxuridine
- IDU
- indomethacin
- iodide and iodine solutions and compounds
- isoniazid
- isotretinoin
- labetalol
- lanatoside C
- levobunolol
- lidocaine
- medryson
- mepivacaine
- mesoridazine
- methdilazine
- methimazole
- methotrexate
- methotrimethazine
- methoxasen
- methylprednisolone
- methylthiouracil
- metipranol
- metoprolol
- minoxidil
- nadolol
- naphazoline
- neomycin
- niacin
- ofloxacin
- oral contraceptives
- ouabain
- oxprenolol
- oxyphenbutazone
- paramethasone
- penicillamine
- perazine
- pericyazine
- perphenazaine
- phenacaine
- pheniramine
- phenylbutazone
- phenylephrine
pilocarpine  sulfacetamide  thiethylperazine
pindolol  sulfachlorpyridazine  thimerosal
piperacetazine  sulfacytine  thiopropazate
piperocaine  sulfadiazine  thioproperazine
polymyxin B  sulfadimethoxine  thioridazine
prednisonolone  sulfamerazine  thiotepa
prilocaine  sulfameter  thiothixene
procaine  sulfamethazine  timolol
prochlorperazine  sulfamethizole  tobramycin
promazine  sulfamethoxazole  trichloroethylene
promethazine  sulfamethoxypyridazine  trifluoperazine
proparacaine  sulfanilamide  triflupromazine
propiomazine  sulfaphenazole  trifluorothymidine
propofol  sulfapyridine  trimedazine
propoxycaine  sulfasalazine  trimethylprazin
propylthiouracil  sulfathiazole  triflupromazine
quinacrine  sulfisoxazole  trioxsalen
radioactive iodides  sulindac  tripelennamine
rubella virus vaccine  tetracaine  vidarabine
(live)  tetracycline  vinblastine
smallpox vaccine  tetracycline  vinblastine
sodium salicylate  tetrahydrozoline

10. Limbal conditions associated with punctate keratitis
   A. Focal necrotic lesions
      (1) Herpes simplex
      (2) Phlyctenular disease
      (3) Vaccinia
   B. Follicles
      (1) Acne rosacea (ocular rosacea)
      (2) Herpes simplex
      (3) Inclusion conjunctivitis
      (4) Molluscum contagiosum
      (5) Trachoma
      (6) Other viral infections
   C. Nodules and plaques
      (1) Avitaminosis A (Bitot spots)
      (2) Bowen disease (dyskeratosis)
      (3) Intraepithelial melanoma
      (4) Limbal vernal conjunctivitis

11. Punctate keratitis preceded by lymphadenopathy
   A. Adenovirus
   B. Herpes simplex
      C. Herpes zoster
      D. Inclusion conjunctivitis
      E. Trachoma
F. Vaccinia
12. Respiratory diseases
   A. Adenovirus infections
   B. Myxovirus infections (influenza, Newcastle disease, mumps)
   C. Recurrent herpes complicating any fever


**MORPHOLOGIC CLASSIFICATION OF PUNCTATE CORNEAL LESIONS**

(CLASSIFICATION BY ANATOMIC LOCATION)

1. Punctate epithelial erosions-fine, very slightly depressed spots scarcely visible without staining with fluorescein
   A. Warts
   B. Artificial-silk keratitis
   C. Staphylococcal blepharoconjunctivitis (lower cornea)
   *D. Keratoconjunctivitis sicca (interpalpebral area)
   *E. Exposure keratitis (interpalpebral area)
   F. Neuroparalytic keratitis (see p. 246)
   G. Ocular medications (especially those with preservatives)
   H. Trichiasis
   I. Trauma, mild (e.g., eye rubbing)

2. Punctate epithelial keratitis-very small, whitish flecks on the surface of the epithelium
   A. Fine
      (1) Scattered-staphylococcal blepharitis; viral keratitis, especially trachoma and molluscum contagiosum, sometimes inclusion conjunctivitis, and not infrequently herpetic keratitis and rubella and rubella
      (2) Confluent-keratitis sicca, exposure keratitis, vernal conjunctivitis, topical steroid-induced and early viral keratitis
B. Coarse

*(1) Thygeson superficial punctate keratitis (characteristic)
(2) Herpes zoster
(3) Adenovirus infections
(4) Early herpes simplex
(5) Acne rosacea (lower cornea)
(6) Encephalitozoon hellem

C. Areolar-spots have enlarged to occupy a large area

(1) Herpes simplex
(2) Thygeson superficial punctate keratitis
(3) Herpes zoster
(4) Vaccinia

3. Filamentary keratitis or keratopathy-formation of fine epithelial filaments that are attached at one end

*A. Keratoconjunctivitis sicca (frequent)
B. Infections, such as that due to adenovirus, herpes, vaccinia, acne rosacea, molluscum contagiosum, rubella, rubeola, and staphylococcus
C. Trauma, such as wounds, abrasions, exposure to shortwave diathermy, and prolonged eye patching
D. Edema of cornea, such as that due to recurrent erosions or wearing of contact lens
E. Sarcoid with infiltration of conjunctiva and lacrimal gland
F. Heerfordt syndrome and Mikulicz syndrome
G. After irradiation-of the lacrimal gland
H. Keratoconus (see p. 288)
I. Neuropathic keratopathy (anesthesia of cornea, p. 246)
J. Conjunctival cicatrizatization, such as that associated with ocular cicatricial pemphigoid, erythema multiforme, ocular psoriasis, and advanced trachoma
K. Degenerative condition of corneal epithelium, such as in advanced glaucoma
L. Superior limbic keratoconjunctivitis
M. Hereditary hemorrhagic telangiectasis (Rendu-Osler-Weber disease)
N. Aerosol keratitis
O. Diabetes mellitus
P. Ectodermal dysplasia
Q. Following cataract or corneal transplant surgery
R. Following patching
S. Use of diphenhydramine hydrochloride (Benadryl)
T. Idiopathic

4. Punctate subepithelial keratitis-punctate epithelial keratitis may progress to combine epithelial and subepithelial lesions followed by healing of the epithelial component, leaving a punctate subepithelial keratitis typical of viral punctate keratitis

A. Areolar or stellate lesions-grayish white

(1) Herpes simplex (usually)
(2) Herpes zoster
(3) Vaccinia
(4) Infectious mononucleosis
(5) Epstein-Barr virus infection
(6) Dimmer keratitis
(7) Brucellosis
(8) Onchocerciasis

B. Fine or medium-sized lesions, typically
   (1) Adenovirus, especially types 3 and 7-grayish white

C. Yellowish tinge-typical of trachoma, inclusion conjunctivitis, acne rosacea, and marginal “catarrhal infiltrates” associated with staphylococcal blepharitis, *Neisseria* conjunctivitis, and Reiter disease

5. Punctate opacifications of Bowman membrane-gray, homogeneous, thickened spots, often with irregular edges
   A. Salzmann degeneration
   B. Punctate lesion of trachoma, measles, or phlyctenular disease


**SICCA KERATITIS (DRY EYE WITH SECONDARY CORNEAL CHANGES)**

1. Boeck sarcoid (Schaumann syndrome)
2. Dermatitis herpetiformis
3. Diabetes mellitus (Willis disease)
4. Herpes simplex
5. Lye burns
   *6. Ocular cicatricial pemphigoid
7. Polychondritis
   *8. Sjögren syndrome (secretinhibitor syndrome)
9. Trachoma
10. Vitamin A deficiency (xerosis)
11. Stevens-Johnson syndrome


**WHITE RINGS OF THE CORNEA (COATS DISEASE)**

These rings are made up of a series of tiny white dots that may coalesce at the level of Bowman membrane or just below it.
1. Congenital
2. Trauma
   * A. Foreign body, usually metal
   B. Occupational—in working with limestone, there may be deposition of some of
      the substance's components, especially calcium oxide, in the cornea
3. Intraocular disease
4. Iron deposition


DRY SPOTS OF THE CORNEA (PRE CORNEAL TEAR FILM DRYING IN SPOT-WISE FASHION)

The precorneal tear film is best examined by using fluorescein and cobalt-blue
filtered light. Patients may have difficulty wearing contact lenses or may have corneal
pain. Normal tear-film breakup time is greater than seconds and averages 25 to 30
seconds.

1. Associated with corneal dellen (see p. 280)
2. Chemical burns
3. Chronic bacterial or viral conjunctivitis
4. Congenital alacrima
5. Instillation of topical anesthetic
   *6. Keratitis sicca
   *7. Ocular cicatricial pemphigoid
8. Ocular pemphigus (chronic cicatricial conjunctivitis)
9. Sleep apnea syndrome
10. Sometimes in elderly persons without obvious pathology
11. Stevens-Johnson syndrome (erythema multiforme)
12. Vitamin A deficiency

Dohlman CH. The function of the corneal epithelium in health and disease. *Invest

Lemp MA, Hamill JR. Factors affecting tear film break-up in normal eyes. *Arch


ANTERIOR EMBRYOTOXON (ARCUS)

In this condition, white or gray substance is deposited at level of the Descemet
membrane and Bowman membrane initially and then in the stroma with a clear limbal
interval.
1. Age—may be present normally in a white patient older than 40 years of age or in a black patient older than 35 years of age
2. Alagille syndrome
3. Alport syndrome (hereditary nephritis)
4. Associated with corneal disease, such as interstitial keratitis
5. Contralateral carotid occlusive disease—when unilateral
6. Familial hypercholesterolemia (type II, familial beta-lipoproteins and type III, familial hyper-beta and pre-beta lipoproteins [carbohydrate-induced hyperlipemia])
7. Hereditary-autosomal dominant or autosomal recessive inheritance
8. Isolated phenomenon
9. Long exposure to irritating dust or chemicals
10. Ocular anomaly association, such as blue sclera (see p. 231), megalocornea (see p. 255), or aniridia (see p. 364-365)
11. Secondary to ocular disease, such as large corneal scars, sclerokeratitis, limbal dermoid, nevus, or epithelial cyst
12. Schnyder crystalline dystrophy


**BOWMAN MEMBRANE FOLDS**

1. Bullous keratopathy
2. Idiopathic
3. Inflammation
4. Lowering of intraocular pressure, such as occurs in association with phthisis bulbi


**Extracted Table Anterior corneal abnormalities**

**DELAYED CORNEAL WOUND HEALING**

Delayed corneal wound healing because of drugs, including the following:

- adenine
- adrenal cortex injection
- aldosterone
- alpha chymotrypsin(?)
- amphotericin B
- arabinoside
- azathioprine
- bacitracin
- beclomethasone
- benoxinate
- benzalkonium
- betamethasone
- butacaine
- chymotrypsin (?)
- cocaine
- colchicine
- cortisone
- cytarabine
- desoxycorticosterone
- dexamethasone
- dibucaine
- dyclonine
- F3T
- floxuridine
Fundocortisone  medrysone  proparacaine
Fluorometholone  meprednisone  sulfacetamide
Fluorouracil  methylprednisolone  sulfamethizole
Fluprednisolone  paramethasone  sulfisoxazole
Flurbiprofen  penicillamine(?)  tetracaine
ganciclovir  phenacaine  thiopeta
gentamicin  phenylephrine(?)  triamcinolone
Hydrocortisone  piperocaine  trifluorothymidine
Idoxuridine  prednisolone  trifluridine
Iodine solution  prednisone  vidarabine


ANTERIOR CORNEAL MOSAIC
A pattern of fluorescein pooling in corneal epithelial grooves can be induced in any normal eye by pressure on the cornea.

1. Exophthalmos as in dysthyroid eye disease with corneal compression against the eyelids
2. Exposure to a high-pressure fire extinguisher jet
3. Pressure on the cornea, either directly on the cornea or indirectly through the lids


LINEAR OPACITY IN SUPERFICIAL CORNEAL STROMA
1. Arc-like at superior limbus
   A. Poorly fit contact lens
   B. Well-fitting soft contact lens with tight eyelids
2. Central
   A. Amiodarone
   B. Chloroquine and hydroxychloroquine
   C. Microcystic epithelial dystrophy
   D. Phenothiazine


SUPERFICIAL VERTICAL CORNEAL STRIATIONS - EPITHELIAL WRINKLES CAN BE ACCENTUATED WITH FLUORESCEIN

1. Corneal surgery with corneal indentation or low intraocular pressure
2. Graves disease
3. Scarred lids
4. Soft contact lens


DENDRITIC CORNEAL LESIONS (AREA OF STAINING OF CORNEA IN A BRANCHING PATTERN)

* 1. Corneal erosions, in which the epithelium may become loose
* 2. Herpes simplex
* 3. Herpes zoster
4. Use of soft contact lenses
* 5. Acanthamoeba keratitis
6. Latanoprost


BULLOUS KERATOPATHY (TERMINAL STAGES OF SEVERE OR PROLONGED EPITHELIAL EDEMA SECONDARY TO ENDOTHELIAL DAMAGE)

1. Anterior-posterior corneal incisions for myopia
2. Anterior synechiae
3. Associated with progressive facial hemiatrophy (Parry-Romberg syndrome)
4. Birth trauma (forceps injury)
5. Chronic uveitis, especially herpes simplex or herpes zoster
6. Congenital corneal dystrophy
7. Congenital glaucoma
8. Congenital hereditary endothelial dystrophy
9. Corneal hydrops (acute keratoconus)
10. Epithelial downgrowth
* 11. Following cataract surgery with or without intraocular implantation
12. Following perforating wounds, especially when the lens capsule or vitreous is adherent to the cornea
*13. Fuchs epithelial-endothelial dystrophy
14. Immunologic reaction after keratoplasty or endothelial decompensation
15. Iridocorneal endothelial syndrome
16. Long-standing glaucoma
17. Posterior polymorphous dystrophy
18. Prolonged inflammation of corneal stroma, such as in disciform or interstitial keratitis (rare)
19. Silicone oil in anterior chamber


**NUMMULAR KERATITIS (COIN-SHAPED LESIONS OF CORNEA)**

1. Brucellosis
2. Dimmers nummular keratitis
3. Epidemic keratoconjunctivitis
4. Herpes zoster
5. Infectious mononucleosis-Epstein-Barr virus
6. Onchocerciasis (River blindness)
7. Varicella
8. Herpes simplex


**DEEP KERATITIS**

1. Behçet disease (dermatostomatoophthalmic syndrome)
2. Deep pustular keratitis
3. Disciform keratitis
4. Herpes zoster
5. Keratitis profunda
6. Stromal herpes
7. Vaccinia
8. Varicella

INTERSTITIAL KERATITIS (CORNEAL STROMAL INFLAMMATION, NOT PRIMARILY ON ANTERIOR OR POSTERIOR SURFACES OF STROMA)

1. After burns
   A. Acid
   B. Alkali
2. Deep punctate
   A. Influenza
   B. Local trauma
   C. Mumps
   *D. Ophthalmic zoster

*3. Luetic (syphilis)
4. Nonluetic
   A. Acanthamoeba
   B. rosacea (ocular rosacea)
   C. Brucellosis (Bang disease)
   D. Cogan I syndrome (nonsyphilitic interstitial keratitis)
   E. Epstein-Barr
   F. Filariasis
   G. Herpes simplex
   H. Hodgkin disease (lymph node disease)
   I. Leishmania species
   J. Measles
   K. Microsporida
   L. Mycosis fungoides
   M. Mumps
   N. Onchocerciasis
   O. Recurrent fever
   P. Roberts pseudothalidomide syndrome
   Q. Sarcoidosis (Schaumann syndrome)
   R. Sleeping sickness (von Economo syndrome)
   S. Steroid therapy
   T. Topical anesthetic abuse
   U. Trypanosomiasis
   V. Tuberculosis (scrofulous keratitis)
   W. Viral as metaherpetic keratitis
   X. Corneal opacification after forceps delivery
   Y. Human T-lymphotropic virus

5. Sclerosing keratitis
   A. Scleritis
       (1) Foci or some local process
       (2) Hennebert syndrome (luetic otitic nystagmus syndrome)
       (3) Sarcoidosis syndrome (Schaumann syndrome)
       (4) Syphilis (acquired lues)
       (5) Tuberculosis
   B. Sclerocornea
6. With chemical poisons
   A. Arsenic
   B. Gold

7. With corneal ring abscess
   A. Anterior segment necrosis
      (1) After circular diathermy
      (2) After a "string" encircling procedure for retinal detachment
      (3) After multiple extraocular muscle surgery
   B. Bacillus subtilis
   C. Bacterium pyocyaneum
   D. Pneumococci
   E. Proteus species

8. With skin disease
   A. Herpes zoster
   B. Incontinentia pigmenti (Bloch-Sulzberger syndrome)
   C. Lichen planus
   D. Molluscum contagiosum
   E. Palmoplantar keratosis
   F. Pityriasis rubra pilaris
   G. Psoriasis


PANNUS (SUPERFICIAL VASCULAR INVASION CONFINED TO A SEGMENT OF THE CORNEA OR EXTENDING AROUND THE ENTIRE LIMBUS)
* 1. Acne rosacea
*2. Allergic marginal infiltration
*3. Anoxic contact lens overwear syndrome
4. Ariboflavinosis keratopathy
*5. Contact lens usage
6. Deerfly fever (tularemia)
7. Degenerative-blind degenerative eyes; often associated with bullous keratopathy
8. Dermatitis herpetiformis (Duhring-Brocq disease)
9. Drugs including the following:
   benoxinate        butacaine        cocaine
   benzalkonium      chlorhexidine    dibucaine
dyclonine  oxyphenbutazone  tetracaine
F3T  phenacaine  thimerosal
ibuprofen  phenylbutazone  trifluridine
idoxuridine  piperocaine  urokinase(?)
IDU  proparacaine  vidarabine
iodine solution  silicone

10. Fuchs corneal dystrophy (degenerative pannus)
11. Glaucoma (degenerative pannus)
12. *Haemophilus influenzae*
13. Histiocytosis X (Hand-Schüller-Christian syndrome)
14. Hypoparathyroidism
15. Inclusion conjunctivitis in infants and adults (micropannus) (chlamydia)
16. Keratoconjunctivitis sicca
17. Leishmaniasis
18. Leprosy (Hansen disease)
19. Linear nevus sebaceous of Jadassohn
20. Lyell disease (toxic epidermal necrolysis or scalded skin syndrome)
21. Lymphopathia venereum
22. Molluscum contagiosum
23. Ocular cicatricial pemphigoid
24. Onchocerciasis (river blindness)
25. Papilloma (wart)
26. Pellagra (avitaminosis B12)
27. Pemphigus foliaceus (Cazenave disease)
28. Phlyctenular keratoconjunctivitis (see p. 280)
29. Siemens disease (keratosis follicularis spinulosa decalvans)
*30. Staphylococcal keratoconjunctivitis (micropannus)
31. Stevens-Johnson syndrome (mucocutaneous ocular syndrome)
*32. Superior limbic keratoconjunctivitis (micropannus)
33. Terrien disease (senile marginal atrophy)
34. Trachoma
35. Tuberculosis
36. Vaccinia
37. Vernal conjunctivitis (micropannus)
38. Vitamin B12 deficiency (Addison pernicious anemia syndrome)


<table>
<thead>
<tr>
<th>CORNEAL OPAcity-DIFFUSE</th>
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<tbody>
<tr>
<td>1. Acromesomelic dysplasia</td>
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<td>*2. Birth trauma</td>
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<td>3. Cockayne syndrome</td>
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<td>*4. Congenital hereditary endothelial dystrophy</td>
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<td>5. Congenital hereditary stromal dystrophy</td>
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<td>6. Cystinosis</td>
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<td>7. Fabry syndrome</td>
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<td>8. Fetal rubella effects</td>
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<td>9. GM gangliosidosis type 1</td>
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<td>10. Hurler syndrome</td>
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<td>11. Infection</td>
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<td>12. Maroteaux-Lamy syndrome</td>
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<td>13. Morquio syndrome</td>
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<td>14. Mucolipidosis III</td>
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<td>15. Mucolipidosis IV</td>
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<td>16. Multiple sulfatase deficiency</td>
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<td>17. MPS VII</td>
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<td>18. Pachyonychia congenita syndrome</td>
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<td>19. Pena-Shokeir type II syndrome [cerebrooculofacial-skeletal (COFS) syndrome]</td>
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<td>20. Rutherford syndrome</td>
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<td>21. Scheie syndrome</td>
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<td>*22. Sclerocornea</td>
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<td>23. Seip syndrome</td>
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<td>24. Sialidosis, Goldberg type</td>
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<td>25. Trisomy syndrome</td>
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<td>26. 18q syndrome</td>
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<thead>
<tr>
<th>CORNEAL OPAcity-LOCALIZED, CONGENITAL</th>
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<tbody>
<tr>
<td>1. Acromegaloid changes, cutis verticis gyrata, and corneal leukoma</td>
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<tr>
<td>2. Aniridia</td>
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<td>3. Autosomal dominant colomba</td>
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<td>4. Cataract microcornea syndrome</td>
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<td>*5. Dermoid limbal, central, and ring</td>
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<td>6. Fetal alcohol syndrome</td>
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7. Fetal rubella effects
8. Fetal transfusion syndrome
9. Fucosidosis
10. Group 13-trisomy phenotype
11. Keratoconus posticus circumspectus
12. Meesman syndrome
13. Peters anomaly and short stature
14. Pilay syndrome (ophthalmomandibulomelic dysplasia)
15. Radial aplasia, anterior chamber cleavage syndrome
16. Richner-Hanhart syndrome
17. Rieger syndrome
18. Trisomy syndrome
19. Waardenburg syndrome
20. Wedge-shaped stromal opacity
21. 4p syndrome
22. 11q syndrome
23. 18q syndrome


**DEEP CORNEAL STROMAL DEPOSITS**

*1. Cornea farinata
2. Deep filiform dystrophy
3. Deep punctiform dystrophy associated with ichthyosis
4. Fleck corneal dystrophy
5. Gold (chrysiasis)
6. Lattice corneal dystrophy
7. Macular corneal dystrophy
8. Polymorphic amyloid degeneration


**INTRACORNEAL HEMORRHAGE**

1. Associated with intraocular surgery
2. Diseases of cornea, such as corneal ulcers and chemical burns
3. Microbial keratitis
4. Migration from subconjunctival hemorrhage
5. Mooren ulceration
6. Ocular trauma
7. Spontaneous in contact lens wearers


**CENTRAL POSTERIOR STROMAL CORNEAL DEPOSITS**

1. Bence Jones proteinuria
2. Dysproteinemia
3. Filiform corneal dystrophy
4. Immunoglobulin deposition
   A. Abnormal gamma globulin
   B. Benign monoclonal gammopathy
5. Multiple myeloma


**DELEN**

Dellen is characterized by shallow corneal excavation near the limbus, usually on the temporal side; the base of the lesion is hazy and dry.

1. Following the wearing of contact lens
2. In elderly persons-limbal vasosclerosis
3. Lagophthalmos
4. Lengthy administration of cocaine
5. Postcataract section
6. Swelling of perlimbal tissues
   A. Allergic conjunctival edema
   B. Episcleritis
   *C. Filtering bleb
   D. Limbal tumor
   *E. Postoperative advancement of rectus muscle
   F. Postoperative retinal detachment
   G. Pinguecula
   H. Subconjunctival effusion or injection
7. With hemeralopia
PHLYCTENULAR KERATOCONJUNCTIVITIS

This condition is characterized by a localized conjunctival, limbal, or corneal nodule about 1 to 3 mm in size.

1. Delayed hypersensitivity to bacterial protein, particularly tuberculoprotein and staphylococci; lymphopathia venereum and coccidioidomycosis may also be allergens.
2. Malnutrition.
3. Secondary infection of the conjunctiva, especially from *Staphylococcus aureus*, pneumococcus, Koch-Weeks bacillus, chlamydia, coccidioidomycosis, and gonorrhea.
4. Systemic infection:
   A. Bang disease (Brucellosis)
   B. Candidiasis
   C. Neurodermatitis
   D. Mikulicz-Radecki syndrome (dacryosialoadenopathy)
   E. Trachoma
   F. Sjögren syndrome (secretinhibitor syndrome)


CORNEAL RING LESION

1. Acanthamoebic keratitis
2. Associated with rheumatoid arthritis-inferior
3. Associated with Sjögren syndrome (secretinhibitor syndrome)
4. *Capnocytophaga ochracea*
5. Double-ring formation-allergic keratitis
6. Marginal dystrophy-degenerative chronic corneal lesion with stromal thinning and intact epithelium
7. Marginal ulceration-secondary to massive granuloma of sclera or necrotizing nodular scleritis (see p. 237)
8. Mooreau ulcer-deeply undermined central edges and chronic course with inflammation, painful
9. Ring abscess-rapidly destructive purulent lesion in the deepest parts of the cornea
10. Ring ulcer-see marginal corneal ulcers (p. 283-286)
11. Steroid use in furrow dystrophy
12. Terrien marginal degeneration-usually begins superiorly
13. Wegener granulomatosis (Wegener syndrome)


**CORNEOSCLERAL KERATITIS**

1. Boeck sarcoid (Schaumann syndrome)
2. Gout (hyperuricemia)
3. Leprosy (Hansen disease)
4. Infections (e.g., pseudomonas)
5. Malformations, such as in sclerocornea (see p. 301)
6. Sarcoma
7. Syphilis (acquired lues)
8. Trisomy 13 (trisomy D)
9. Tuberculosis
10. Wegener granulomatosis


**CENTRAL CORNEAL ULCER**

1. Bacterial origin
   *A. Diplococcus pneumoniae* (pneumococcus)-infiltrated gray-white or yellow disc-shaped central ulcer typically associated with diffuse keratitis, severe iridocyclitis, and hypopyon; follows corneal abrasion; occurs especially in the presence of chronic dacryocystitis; enhanced by general debility
   *B. Beta-hemolytic streptococcus and other streptococcus species
   *C. Pseudomonas aeruginosa* but may also have *Pseudomonas acidovorans, Pseudomonas stutzeri, Pseudomonas mallei*, and *Pseudomonas pseudomallei*-primary corneal involvement, rapid spread often to panophthalmitis, large hypopyon, thick, greenish pus; may be contaminant of eserine and fluorescein often is associated with contact use.
   D. *Escherichia coli*
   E. *Moraxella liquefaciens* (diplococcus of Petit)-morphologically resembles diplobacillus of Morax-Axenfeld, which is never seen in central corneal ulcers
   F. *Klebsiella pneumoniae*
   G. *Proteus vulgaris*
   H. Actinomyces
   I. Tuberculous-secondary to conjunctival or uveal infections
   J. *Serratia marcescens*-gram-negative coccobacillus
   *K. Staphylococcus aureus, S. epidermidis*, and other *Staphylococcus* species
   L. Mima polymorpha
   M. Dysgonic fermenter-2
   N. Others

2. Viral origin
3. Mycotic origin—follows corneal trauma, such as foreign bodies in the cornea or corneal abrasions caused by vegetable matter, or diseases, such as radiation keratitis, exposure keratitis, herpes zoster, and ocular pemphigus; chronic course; shallow crater; absent corneal vascularization; may follow treatment with antibiotics or, more likely, treatment with steroid-antibiotic combinations
   A. *Absidia corymbifera*
   B. Aspergillus species
   C. *Blastomyces dermatitidis*
   D. Candida albicans
   E. Cephalosporium species
   F. Fusarium solani
   G. Nocardia species
   H. Others

4. Acquired immune deficiency syndrome (AIDS) related

5. Atopic

6. Basement membrane abnormalities as microcysts, evidence of map, dot fingerprints, or anterior stromal dystrophies, trauma history, other dystrophy

7. Brittle cornea syndrome

6. Chemical-latex keratitis, alkali/acid burn

8. Dry eyes, including Sjögren syndrome

9. Exposure as lagophthalmos, lid abnormalities, inadequate blink, facial palsy, proptosis, thyroid disease

10. Extrusion of anterior chamber intraocular lens

11. Factitious

12. Hypogammaglobulinemia

13. Medicamentosus as drops

14. Neurotrophic

15. Thermal/radiation burns

16. Sjögren syndrome

17. Soluble tyrosine aminotransferase (STAT) deficiency


**MARGINAL CORNEAL ULCERS**

1. Ring ulcers—often bilateral, circumcorneal injection, and continuous ring or confluent multiple lesions
   - A. Acute leukemia
   - B. Bacillary dysentery
   - C. Brucellosis (Bang disease)
   - *D. Coalescence of several marginal ulcers*
   - E. Dengue fever
   - F. Gold poisoning
   - G. Gonococcal arthritis
   - H. Following penetrating keratoplasty
   - I. Hookworm infestation
   - J. Influenza
   - K. Last stages of trachoma, secondary to small circumferential pannus
   - *L. Mooren ulcer*
   - M. Polyarteritis nodosa (Kussmaul disease)
   - N. Porphyria
   - *O. Rheumatoid arthritis-Sjögren syndrome (secretion inhibitor syndrome)*
   - P. Scleroderma (progressive systemic sclerosis)
   - Q. *Systemic lupus erythematosus (Kaposi-Libman-Sacks syndrome)*
   - R. Tuberculosis
   - S. Wegener granulomatosis (Wegener syndrome)

2. Simple marginal ulcers—superficial crescentic gray-colored ulcer
   - A. Infection—due to *Staphylococcus* organisms, Koch-Weeks bacillus, pseudomonas, diplobacillus of Morax-Axenfeld; usually chronic
   - B. Toxic or allergic including antiinflammatory drugs
   - C. Systemic disturbances, such as
     1. Acute upper respiratory infection
     2. Bacillary dysentery
     3. Barre-Lieou syndrome (posterior cervical sympathetic syndrome)
     4. Brucellosis (Bang disease)
     5. Crohn disease (granulomatous ileocolitis)
     6. Gout (hyperuricemia)
     7. Influenza
     8. Lupus erythematosus (Kaposi-Libman-Sacks syndrome)
     9. Polyarteritis nodosa (Kussmaul disease)
Postvaccinial ocular syndrome *(11) Rheumatoid arthritis-inferior cornea


**Extracted Table Marginal corneal ulcers**

**DESCemet Membrane Folds (Usually Following Hypotony)**

*1. Trauma, such as that due to cataract or corneal surgery
2. Mechanical cause, such as firm, prolonged ocular bandaging or phthisis bulbi
3. Inflammatory condition, such as that following interstitial or herpes simplex keratitis
4. Diabetes (8%-33%)
5. Ochronosis
6. Toxic
   A. Quinone and hydroquinone-vertical folds
   B. Formaldehyde 26%
   C. Experimental cold injury to cornea
   D. Digitoxin
7. Idiopathic


**DESCemet Membrane Tears (Haab Striae)**

*1. Acute hydrops of the cornea, such as that due to keratoconus
*2. Buphthalmos (e.g., from congenital glaucoma)
3. Conical cornea
4. Myopia with marked anteroposterior diameter
*5. Trauma, such as birth injury or contusion


**DESCEMET MEMBRANE THICKENING**

*1. Central cornea guttata*

A. Primary
B. Secondary cornea guttata
   *(1) Congenital luetic interstitial keratitis
   (2) Endothelial cell insult
      a. Breaks in Descemet membrane, including scrolls of Descemet membrane in healed syphilitic interstitial keratitis
      b. Chandler syndrome (iridocorneal endothelial syndrome)
      c. Cogan-Reese syndrome (iris-nevus syndrome)
   *d. Corneal dystrophy, including Fuchs syndrome
   e. Posterior keratoconus syndrome
C. Transient cornea guttata associated with short-term episodes of iritis and corneal inflammation

*2. Peripheral Hassall Henle warts*


**RETROCORNEAL PIGMENTATION**

1. Endothelial phagocytosis of free melanin pigment, such as Krukenberg spindle
2. Iris melanocytes, iris pigment epithelial cells, or pigment containing macrophages in the posterior corneal surface; can follow operative or accidental ocular trauma
3. Status post hyphema


**LOW ENDOTHELIAL CELL COUNT (DIMINISHED NUMBER OF CORNEAL ENDOTHELIAL CELLS)**

1. Acute and chronic uveitis
*2. Corneal endothelial dystrophy
3. Following cataract or other intraocular surgery
4. Cornea guttata, endothelial dystrophy, and Fuch dystrophy


**SNAIL TRACKS OF CORNEA**

This condition involves irregular, discontinuous grayish white streaks or patches, usually orientated horizontally and obliquely on the corneal endothelium.

1. Corneal buttons preserved in corneal storage medium
2. Following ocular surgery
3. Ocular trauma


**KERATOCONUS (CONICAL CORNEA)**

Keratoconus is characterized by noninflammatory ectasia of the cornea in its axial part, with considerable visual impairment because of development of a high degree of irregular myopic astigmatism. Keratoconus may be associated with

1. Acute hydrops of the cornea
2. Alagille syndrome
3. Anetoderma and bilateral subcapsular cataracts
4. Angelman syndrome
5. Aniridia
6. Apert syndrome (acrodysplasia)
7. Asthma, hay fever
*8. Atopic dermatitis, keratosis plantaris, and palmaris
9. Autographism
10. Avellino dystrophy
11. Blue sclerotics, including van der Hoeve syndrome (osteogenesis imperfecta) (see blue sclera, p. 231)
12. Chandler syndrome (iridocorneal endothelial syndrome)
13. Congenital hip dysplasia
14. Wearing of contact lens
15. Crouzon syndrome
16. Deep filiform corneal dystrophy
17. Ehlers-Danlos syndrome (fibrodysplasia elastica generalisata, cutis hyperelastica)
18. Essential iris atrophy
19. Facial hemiatrophy
*20. Familial
21. False chordae tendineae of left ventricle
22. Fleck corneal dystrophy
23. Focal dermal hypoplasia (Goltz syndrome)
24. Fuchs corneal endothelial dystrophy
25. Grönblad-Strandberg syndrome (pseudoxanthoma elasticum)
26. Hereditary history
27. Hyperextensible joints and mitral valve prolapse
28. Hyperornithemia
29. Infantile tapetoretinal degeneration of Leber
30. Iridocorneal dysgenesis
31. Iridoschisis
32. Joint hypermobility
33. Kurz syndrome
34. Laurence-Moon-Biedl syndrome (retinitis-polydactyly-adiposogenital syndrome)
35. Little syndrome (nail-patella syndrome)
36. Lymphogranuloma venereum
37. Marfan syndrome (arachnodactyly dystrophia mesodermalis congenita)
38. Measles retinopathy
39. Microcornea
40. Mongolism (Down syndrome)
41. Mulvihil-Smith syndrome
42. Neurocutaneous angiomatosis
43. Neurodermatitis
44. Neurofibromatosis (von Recklinghausen syndrome)
45. Noonan syndrome (male Turner syndrome)
46. Ocular hypertension
47. Pellucid marginal corneal degeneration
48. Posterior ectasia following laser in situ keratomileusis (generally if stromal bed less than 250 μ)
49. Posterior lenticus
50. Posterior polymorphous dystrophy
51. Retinal disinsertion syndrome
52. Retinitis pigmentosa
53. Retinopathy of prematurity
54. Rieger syndrome
55. Tourette disease
56. Thalassemia syndrome
*57. Trauma, such as rubbing of eyes, birth injury, or contusion
58. Vernal catarrh
59. Vernal conjunctivitis
60. 18q syndrome


**CORNEA PLAN A (DECREASED CORNEAL CURVATURE)**

1. Isolated
2. Marfan syndrome
*3. Sclerocornea


**STAPHYLOMA OF CORNEA (CORNEAL STRETCHING WITH INCARCERATION OF UVEAL TISSUE)**

1. Advanced keratoconus (see p. 288)
2. Avitaminosis A with keratomalacia
3. Congenital
*4. Following corneal ulcer (see p. 281, 283), neuroparalytic keratitis, corneal leprosy, and severe corneal injury
5. Mucoviscidosis (cystic fibrosis of the pancreas)


**WHORL-LIKE CORNEAL LESIONS**

*1. Amiodarone toxicity
2. Amodiaquine hydrochloride administration
3. Atabrine administration
*4. Chloroquine and hydroxychloroquine toxicity
5. Chlorpromazine administration
*6. Fabry disease (diffuse angiokeratosis)
7. Incontinentia pigmenti
8. Indomethacin administration
9. Meperidine hydrochloride
10. Quinacrine administration
11. Urethana dministration


**CORNEAL DERMIOIDS**

These congenital corneal limbal lesions grow slowly. Tumors are yellowish, elevated, and variable in size; they consist of fibrofatty tissue covered by epidermal rather than by conjunctival epithelium and may contain ectodermal derivatives such as hair follicles, sebaceous glands, and sweat glands. Trauma, irritation, and puberty hasten their growth.

1. Bloch-Sulzberger syndrome (incontinentia pigmeneti)
2. Cri-du-chat syndrome (cat-cry syndrome)
3. Duane retraction syndrome
4. Multiple dermoids of the cornea associated with miliary aneurysms of the retina
5. Neurocutaneous syndrome (ectomesodermal dysgenesis)
6. Nevus sebaceous of Jadassohn (linear nevus sebaceous of Jadassohn)
*7. Oculoauriculovertebral dysplasia (Goldenhar syndrome)*
8. Organoid nevus syndrome
9. Ring dermoid syndrome-autosomal dominant
10. Sporadic
11. Thalidomide teratogenicities


**Extracted Table Corneal Dermoids**

**CORNEAL PROBLEMS ASSOCIATED WITH KERATOTIC SKIN LESIONS**

1. Ectodermal dysplasia (anhidrotic)
*2. Ichthyosis*
3. Keratosis follicularis
4. Keratosis follicularis spinulosa decalvans
5. Keratosis plantaris and palmaris
6. Pityriasis rubra pilaris
CORNEAL PROBLEMS ASSOCIATED WITH LID EXCRESCENCES
1. Keratosis folliculosis
2. Lipid proteinosis
*3. Molluscum contagiosum
4. Verruca vulgaris

CORNEAL DISEASE ASSOCIATED WITH LENTICULAR PROBLEMS
1. Aberfeld syndrome (ocular and facial abnormalities syndrome) - cataracts, microcornea
2. Acrodermatitis chronica atrophicans-keratomalacia, corneal opacification, cataracts
3. Addison syndrome (idiopathic hypoparathyroidism) - keratoconjunctivitis, corneal ulcers, keratitic moniliasis, cataracts
4. Amiodarone-corneal deposits, anterior subcapsular cataracts
5. Amyloidosis-amyloid deposits of cornea, corneal dystrophy, pseudopodia lenti
6. Anderson-Warburg syndrome (oligophrenia-microphthalmos syndrome)-corneal opacification and lenticular destruction with a mass visible behind the lens
7. Andosky syndrome (atopic cataract syndrome)-atopic keratoconjunctivitis, keratoconus, uveitis, subcapsular cataract
8. Aniridia-microcornea and subluxated lenses
9. Anterior chamber cleavage syndrome (Reese-Ellsworth syndrome) - corneal opacities, anterior pole cataract
10. Anterior segment ischemia syndrome-corneal edema, corneal ulceration, cataract
11. Apert syndrome (absent digits cranial defects syndrome) - exposure keratitis, cataracts, ectopia lenti
12. Arteriovenous fistula (arteriovenous aneurysm) - bullous keratopathy, cataract
13. Aspergillosis-corneal ulcer, keratitis, cataract
*14. Atopic disease (atopic eczema, Besnier prurigo)-keratoconjunctivitis, keratoconus, cataract
15. Autosomal dominant - cataracts and microcornea
16. Avitaminosis C (scurvy)-keratitis, corneal ulcer, cataract
17. Chickenpox (varicella)-corneal ulcer, corneal opacity, keratitis, cataract
18. Chlorpromazine-corneal and lens opacities
19. Cholera-keratomalacia, cataract
20. Chromosome partial deletion (short-arm) syndrome-cataracts, corneal opacities
21. Cockayne syndrome (Mickey Mouse syndrome)-cataracts, band keratopathy, corneal dystrophy
22. Congenital spherocytic anemia (congenital hemolytic jaundice) - congenital cataract, ring-shaped pigmented corneal deposits
23. Crouzon syndrome (Parrot-head syndrome) - exposure keratitis, cataract, corneal dystrophy
24. Cryptophthalmia syndrome (cryptophthalmos-syndactyly syndrome) - cornea differentiated from sclera, lens absence to hypoplasia, dislocation, and calcification
25. Cytomegalic inclusion disease (cytomegalovirus)-cataract, corneal opacities
26. Darier-White syndrome (keratosis follicularis)-keratosis, corneal subepithelial infiltrations, corneal ulceration, cataract
27. Dermatitis herpetiformis (Duhring - Broca disease) - corneal vascularization, cataract
28. Diphtheria-keratitis, corneal ulcer, cataract
29. Ehlers-Danlos syndrome (fibrodysplasia elastica generalisata)-microcornea, keratoconus, lens subluxation
30. Electrical injury-corneal perforation, necrosis of cornea, anterior or posterior subcapsular cataracts
31. Exfoliation syndrome (capsular exfoliation syndrome)-cataract, dislocated lens, corneal dystrophy, lens capsule exfoliation
32. Folling syndrome (phenylketonuria)-corneal opacities, cataracts
33. Fuchs syndrome (I) (heterochromic cyclitis syndrome)-secondary cataract, edematous corneal epithelium
34. Goldscheider syndrome (epidermolysis bullosa)-bullous keratitis, corneal subepithelial blisters to corneal perforation, cataract
35. Gorlin-Goltz syndrome (multiple basal cell nevi syndrome)-cataract, corneal leukoma
36. Grönblad-Strandberg syndrome (elastorrhexis)-keratoconus, cataract, subluxation of lens
37. Hallermann-Streiff syndrome (oculomandibulofacial dyscephaly)-cataracts, microcornea
38. Hanhart syndrome (recessive keratosis palmoplantaris)-dendritic corneal lesions, keratitis, corneal haze, corneal neovascularization, cataract
39. Heerfordt syndrome (uveoparotid fever)-band keratopathy, keratoconjunctivitis, cataract
40. Hereditary ectodermal dysplasia syndrome (Siemens syndrome)-keratosis, corneal erosions, corneal dystrophy, cataract, lens luxation
*41. Herpes simplex-keratitis, corneal ulcer, cataract
*42. Herpes zoster-keratitis, corneal ulcer, cataract
43. Histiocytosis X (Hand-Schüller-Christian syndrome)-pannus, bullous keratopathy, corneal ulcer, cataract
44. Hodgkin disease-keratitis, cataract
45. Homocystinuria syndrome-dislocated lens, cataract, keratitis
46. Hutchinson-Gilford syndrome (progeria)-cataract, microcornea
47. Hydatid cyst (echinococcosis)-keratitis, corneal abscess, cataract
48. Hypervitaminosis D-band keratopathy, cataract
49. Hypoparathyroidism-keratitis, cataract
50. Hypophosphatasia (phosphoethanolaminuria)-cataract, corneal subepithelial calcifications
51. Influenza-keratitis, cataract
52. Jadassohn-type anetodermal-keratoconus, cataract
53. Jadassohn-Lewandowsky syndrome (pachyonychia congenita)-corneal dyskeratosis, cataract
54. Juvenile rheumatoid arthritis (Still disease)-band keratopathy, cataract
55. Kussmaul disease (periarteritis nodosa)-corneal ulcer, cataract
56. Kyrie disease (hyperkeratosis penetrans)-subcapsular cataracts, subepithelial corneal opacities
57. Leri syndrome (carpal tunnel syndrome)-corneal clouding, cataract
58. Listerellosis (listeriosis)-keratitis, corneal abscess and ulcer, cataract
59. Little syndrome (nail-patella syndrome)-microcornea, keratoconus, cataract
60. Lowe syndrome (oculocerebrorenal syndrome) -cloudy cornea, cataracts, megalocornea, corneal dystrophy
61. Miliaria-keratitis, cataract
62. Marchesani syndrome (brachymorphia with spherophakia) -lenticulinar myopia, ectopia lentis, megalocornea, corneal opacity
63. Marfan syndrome (arachnodactyly-dystrophia-mesodermalis congenita)-lens dislocation, cataract, megalocornea, lenticular myopia
64. Matsoukas syndrome (oculocerebroarticuloskeletal syndromes)-cataract, corneal sclerosis
65. Meckel syndrome (dysencephalia splanchnocystica syndrome) -sclerocornea, microcornea, cataract
66. Morbilli (rubeola, measles)-corneal ulcer, cataract
67. Mucolipidosis IV (ML IV)-corneal clouding, cataract
68. Nematode ophthalmia syndrome (toxocariasis)-cataract, larvae present in the cornea
69. Neurodermatitis (lichen simplex chronicus)-keratoconjunctivitis, atopic cataracts, keratoconus
70. Ocular toxoplasmosis (toxoplasmosis)-keratitis, cataract
71. Oculodental syndrome (Peter syndrome)-macrocornea, opacities of the corneal margin, ectopic lentis, corneoscleral staphyloma
72. O'Donnell-Pappas syndrome-presenile cataract, peripheral corneal pannus
73. Paget syndrome (osteitis deformans)-corneal ring opacities, cataract
74. Passow syndrome (status dysraphicus syndrome)-neuroparalytic keratitis, zonular cataract
75. Pemphigus foliaceus (Cazenave disease)-pannus, corneal infiltration, cataract
76. Pigmentary ocular dispersion syndrome (pigmentary glaucoma)-Krukenberg spindle, equatorial pigmentation of lens capsule
77. Pseudohyoparathyroidism (Seabright-Bantan syndrome)-punctate cataracts, keratitis
78. Radiation-corneal ulcer, punctate keratitis, cataracts, exfoliation of lens capsule
79. Refsum syndrome (phytanic acid oxidase deficiency)-corneal opacities, cataracts
80. Relapsing polychondritis-corneal ulcer, cataracts, keratoconjunctivitis sicca
81. Retinal disinsertion syndrome-lens subluxation, keratoconus
82. Retinopathy of prematurity- cataracts, corneal opacification
83. Rieger syndrome (dysgenesis mesodermalis corneae et irides)-microcornea, corneal opacities in Descemet membrane, dislocated lens
84. Romberg syndrome (facial hemiatrophy)-neuroparalytic keratitis, cataracts
85. Rubella syndrome (German measles)-corneal haziness, cataracts, microcornea
86. Sabin-Feldman syndrome-posterior lenticonus, microcornea
87. Sanfilippo-Good syndrome (MPS III)-deposits in cornea and lens
88. Schafer syndrome (keratosis palmoplantaris syndrome)-lesions in the lower portion of the cornea, cataract
89. Schaumann syndrome (sarcoidosis syndrome)-keratitis sicca, band-shaped keratitis, complicated cataract
90. Scheie syndrome (MPS IS)-diffuse haze to marked corneal clouding, cataracts
91. Stannus cerebellar syndrome-corneal vascularization, corneal opacities, cataracts
92. Stevens-Johnson syndrome (erythema multiforme exudativum)-keratitis, corneal ulcers, cataracts, pannus
93. Stickler syndrome (hereditary progressive arthroophthalmopathy)-keratopathy, cataracts
94. Thioridazine-corneal and lens opacities
95. Toxic lens syndrome-pigment precipitation on the surface of an intraocular lens, chronic uveitis
96. Trisomy syndrome-corneal and lens opacities
97. Turner syndrome (gonadal dysgenesis )-corneal nebulae, cataracts
98. Ultraviolet radiation-band keratopathy, keratitis, discoloring of lens
99. van Bogaert-Scherer-Epstein syndrome (familial hypercholesterolemia syndrome)-lipid keratopathy, cataract, juvenile corneal arcus
100. von Recklinghausen syndrome (neurofibromatosis)-nodular swelling of corneal nerves, cataracts
101. Waardenburg syndrome (intercilioiridodermatoauditive dysplasia)-microcornea, cornea plana, lenticous
102. Wagner syndrome (hereditary hyaloideoretinal degeneration and palatoschisis)-corneal degeneration, including band-shaped keratopathy, cataracts
103. Ward syndrome (nevus jaw cyst syndrome)-congenital cataracts, congenital corneal opacities
104. Wegener syndrome (Wegener granulomatosis)-corneal ulcer, corneal abscess, cataract
105. Weil disease (leptospirosis)-keratitis, cataract
106. Werner syndrome (progeria of adults)-juvenile cataracts, bullous keratitis, trophic corneal defects
107. Yersiniosis-corneal ulcer, cataract
108. Zellweger syndrome (cerebrohepatorenal syndrome of Zellweger)-corneal opacities, cataract


CORNEAL DISEASE ASSOCIATED WITH RETINAL PROBLEMS

1. Abdominal typhus (enteric fever)-corneal ulcer, retinal detachment, central retinal artery emboli
2. *Acanthamoeba-keratitis*, pannus, corneal ring abscess, retinal perivasculitis
3. African eyeworm disease-keratitis, central retinal artery occlusion, macular hemorrhages
4. Amyloidosis-amyloid corneal deposits, corneal dystrophy, retinal hemorrhages
5. Anderson- Warburg syndrome (oligophrenia-microphthalmos syndrome) -corneal opacification, malformed retina with retina pseudotumors
6. Angioedema (hives)-central serous retinopathy, corneal edema
7. Anterior segment ischemia syndrome-corneal edema midperiphery retinal hemorrhages
8. Apert syndrome (acrodysplasia)-exposure keratitis, retinal detachment
9. Arteriovenous fistula-bullous keratopathy, retinal hemorrhages
10. Aspergillosis-corneal ulcer, keratitis, retinal hemorrhages, retinal detachment
11. Atopic dermatitis-keratoconus and retinal detachment
12. Avitaminosis C-retinal hemorrhages, keratitis, corneal ulcer
13. *Bacillus cereus-ring* abscess of cornea, necrosis of retina
14. Bang disease (brucellosis)-keratitis, chorioretinitis, macular edema
15. Behçet syndrome (dermatostomata-ophthalmic syndrome)-keratitis, posterior corneal abscess, retinal vascular changes
16. Bietti disease (Bietti marginal crystalline dystrophy)-marginal corneal dystrophy, retinitis punctate albcens
17. Candidiasis-keratitis, corneal ulcer, retinal atrophy, retinal detachment
18. Carotid artery syndrome-corneal ulcer, loss of corneal sensation, retinal edema, engorgement of retinal veins
19. Chickenpox (varicella)-corneal ulcer, corneal opacity, retinitis, hemorrhagic retinopathy
20. Chloroquine-corneal epithelial pigmentation, macular lesions
21. Chronic granulomatous disease of childhood-keratitis, destructive chorioretinal lesions
22. Cockayne syndrome (dwarfism with retinal atrophy and deafness)-pigmentary degeneration, band keratopathy, corneal dystrophy
23. Crohn disease (granulomatous ileocolitis)-marginal corneal ulcers, keratitis, macular edema, macular hemorrhages
24. Cryoglobulinemia-deep corneal opacities, venous stasis
25. Cystinosis (aminoaciduria)-crystals in cornea and pigment in retina
26. Dengue fever-keratitis, corneal ulcer, retinal hemorrhages
27. Diffuse keratoses syndrome-corneal nodular thickening in the stroma worse in fall, retinal phlebitis
28. Diphtheria-keratitis, corneal ulcer, central artery occlusion
29. Disseminated lupus erythematosus (Kaposi-Libman-Sacks syndrome)-keratitis, keratoconjunctivitis sicca, corneal ulcer, central retinal vein occlusion, retinal detachment
30. Ehlers-Danlos syndrome (cutis hyperelastica)-keratoconus and retinitis pigmentosa
31. Electrical injury-corneal perforation, retinal edema, retinal hemorrhages, pigmentary degeneration, retinal holes, dilatation of retinal veins
32. Fabry disease (diffuse angiookeratosis)-whorl-like changes in cornea, central retinal artery occlusion, tortuosity of retinal vessels
33. Goldscheider syndrome (epidermolysis bullosa)-bullous keratitis with opacities, retinal detachment
34. Gronblad-Strandberg syndrome (systemic elastodystrophy)-angioid streaks of the retina, macular hemorrhages, retinal detachment, keratoconus
35. Hamman-Rich syndrome (alveolar capillary block syndrome)-keratomalacia ischemic retinopathy, cystic macular changes
36. Heerfordt syndrome (uveoparotid fever)-band keratopathy, retinal vasculitis
37. Hennebert syndrome (luetic otitic nystagmus syndrome)-interstitial keratitis, disseminated syphilitic chorioretinitis
38. Histiocytosis X (Hand-Schüller-Christian syndrome)-retinal hemorrhage, retinal detachment, bullous keratopathy, corneal ulcer, pannus
39. Hodgkin disease-keratitis, retinal hemorrhages
40. Hollenhorst syndrome (chorioretinal infarction syndrome)-hazy cornea, serous retinal detachment, pigmentary retinopathy
41. Hunter syndrome (MPS II)-splitting or absence of peripheral Bowman membrane, stromal haze, pigmentary retinal degeneration, narrowed retinal vessels
42. Hurler-Scheie syndrome (MPS IH-S)-corneal clouding, pigmentary retinopathy
43. Hurler syndrome (gargoylism)-diffuse corneal haziness, retinal pigmentary changes, megalocornea, retinal detachment
44. Hydatid cyst (echinococcosis)-keratitis, abscess of cornea, retinal detachment, retinal hemorrhages
45. Hyperlipoproteinemia-arcus juvenilis, lipemia retinalis, xanthomata of retina
46. Hyperparathyroidism-band keratopathy, vascular engorgement of retina
47. Hypovitaminosis A-keratomalacia with perforation, corneal opacity, retinal degeneration
48. Idiopathic hypercalcemia (blue-diaper syndrome)-band keratopathy, optic atrophy, papilledema
49. Indomethacin-corneal deposits, reduced retinal sensitivity
50. Influenza-keratitis, retinal hemorrhage
51. Japanese River fever (typhus)-keratitis, retinal hemorrhages
52. Juvenile rheumatoid arthritis (Still disease)-band keratopathy, macular edema
53. Kahler disease (multiple myeloma)-crystalline deposits of cornea, central retinal artery occlusion, retinal microaneurysms
54. Kussmaul disease (periarteritis nodosa)-retinal detachment, pseudoretinitis pigmentosa, corneal ulcer
55. Leber tapetoretinal dystrophy syndrome (retinal aplasia)-keratoconus, salt-and-pepper or “bone corpuscle” pigmentation, yellowish-brown or gray macular lesions
56. Lubarsch-Pick syndrome (primary amyloidosis)-amyloid corneal deposits, retinal hemorrhages
57. Lymphogranuloma venereum disease (Nicolas-Favre disease)-keratitis, pannus, corneal ulcer, keratoconus, tortuosity of retinal vessels, retinal hemorrhages
58. Marfan syndrome (arachnodactyly dystrophy mesodermalis congenita)-keratoconus, retinitis pigmentosa
59. Meckel syndrome (dysencephalia splanchnocystica syndrome) -sclerocornea, microcornea, retinal dysplasia
60. Meningococcemia-keratitis, retinal endophlebitis
61. Mikulicz-Radeski syndrome (dacryosialoadenopathy)-keratoconjunctivitis, retinal candlewax spots
62. ML IV (mucolipidosis IV)-corneal clouding, corneal opacities, retinal atrophy
63. Morbilli (measles-rubeola)-keratitis, corneal ulcer, pigmentary retinopathy, central retinal artery occlusion
64. Mucormycosis (phycomycosis)-corneal ulcer, striate keratopathy, retinitis, central retinal artery thrombosis
65. Mycosis fungoides syndrome (malignant cutaneous reticulosis syndrome)-keratoconjunctivitis, retinal edema, retinal hemorrhage
66. Myotonic dystrophy syndrome - corneal epithelial dystrophy, loss of corneal sensitivity, tapetoretinal degeneration, macular red spot, macular degeneration, chorioretinitis
67. Neurofibromatosis (von Recklinghausen syndrome)-nodular swelling nerves, hamartoma of retina
68. Norrie disease (atrophia oculi congenita)-malformation of sensory cells of retina, corneal nebulae
69. Oculodental syndrome (Peter syndrome)-corneoscleral staphyloma, megalocornea, corneal marginal opacities, macular pigmentation
70. Onchocerciasis syndrome-punctate keratitis, sclerosing keratitis, chorioretinitis, retinal degeneration
71. Paget syndrome (osteitis deformans)-corneal ring opacities, retinal hemorrhages, pigmentary retinopathy, macular changes resembling Kuhnt-Junius degeneration
72. Phenothiazine-epithelial and endothelial pigment, retinal pigmentation
73. Pierre Robin syndrome (micrognathia-glossoptosis)-retinal disinsertion, megalocornea
74. Plasma lecithin (cholesterol acyltransferase deficiency)-corneal stromal opacities, retinal hemorrhages
75. Porphyria cutanea tarda-keratitis, retinal hemorrhages, cotton-wool spots, macular edema
76. Postvaccinial ocular syndrome-corneal vesicles, and marginal ulcers, chorioretinitis, central serous retinopathy, central retinal vein thrombosis
77. Progressive systemic sclerosis-marginal corneal ulcers with cicatization, cotton-wool spots, retinal hemorrhages
78. Radiation-corneal ulcer, punctate keratitis, keratoconjunctivitis sicca, retinal hemorrhage, macular degeneration, macular holes with vascularization
79. Refsum syndrome (phytanic acid oxidase deficiency)-band keratopathy, retinitis pigmentosa
80. Relapsing fever-interstitial keratitis, retinal hemorrhage
81. Relapsing polychondritis-corneal ulcer, retinal detachment, retinal artery thrombosis, keratoconjunctivitis sicca
82. Renal failure-cotton-wool spots, band keratopathy
83. Rendu-Osler syndrome (hereditary hemorrhagic telangiectasis)-intermittent filamentary keratitis, small retinal angiomata, retinal hemorrhages
84. Retinal disinsertion syndrome-bilateral keratoconus, retinal detachment
85. Retinoblastoma-corneal tumor, retinal neovascularization
86. Rothmund syndrome (telangiectasia-pigmentation cataract syndrome)-corneal lesions, retinal hyperpigmentation
87. Rubella syndrome (Gregg syndrome)-microcornea, pigmentary retinal changes
88. Sabin-Feldman syndrome-microcornea, chorioretinitis or atrophic degenerative chorioretinal changes
89. Sanfilippo-Good syndrome (MPS III)-slight narrowing of retinal vessels, acid mucopolysaccharide deposits in cornea
90. Schaumann syndrome (sarcoidosis syndrome)-mutton fat keratic precipitates, keratitis sicca, band-shaped keratitis, inflammatory retinal exudates
91. Scheie syndrome (MPS IS)-diffuse to marked corneal clouding, tapetoretinal degeneration
92. Schwartz syndrome (glaucoma associated with retinal detachment)-retinal detachment, microcornea
93. Shy-Gonatas syndrome (orthostatic hypotension syndrome)-keratopathy, corneal ulcer, lattice-like white opacities in the area of Bowman membrane, retinal pigmentary degeneration
94. Smallpox-keratitis, congenital corneal clouding, chorioretinitis
95. Stannus cerebellar syndrome (riboflavin deficiency)-corneal vascularization, superficial diffuse keratitis, corneal opacities, brownish retinal patches
96. Stickler syndrome (hereditary progressive arthroophthalmopathy)-keratopathy, chorioretinal degeneration, total retinal detachment
97. Sturge-Weber syndrome (neurooculocutaneous angiomatosis)-retinal detachment, increased corneal diameter with cloudiness
98. Syphilis (acquired lues)-keratitis, retinal hemorrhages, retinal proliferation
99. Temporal arteritis syndrome (Hutchinson-Horton-Magath-Brown syndrome)-retinal detachments, narrowing of retinal vessels, central retinal artery occlusion, corneal hyposthesia
100. Trisomy 13 (Patau syndrome)-malformed cornea, retinal dysplasia
101. Tuberculosis keratitis, pannus, corneal ulcer, retinitis
102. Ullrich syndrome (dyscraniopygophalangy)-cloudy cornea, corneal ulcers, chorioretinal coloboma
103. Ultraviolet radiation-photokeratitis, band keratopathy, herpes simplex keratitis, recurrent corneal erosions, retinal degeneration
104. Vaccinia keratitis, pannus, corneal perforation, central serous retinopathy, pseudoretinitis pigmentosa
105. van Bogaert-Scherer-Epstein (primary hyperlipidemia)-arcus juvenilis of the cornea, lipid keratopathy, retinopathy with yellowish deposits
106. Vitreous tug syndrome-vitreous strands attached to corneal wound or scar, circumscribed retinal edema, posterior retinal detachment
107. von Gierke disease (glycogen storage disease type I)-corneal clouding, discrete non-elevated, yellow flecks in macula
108. Waardenburg syndrome (embryonic fixation syndrome)-microcornea, cornea plana, hypopigmentation and hypoplasia of retina
109. Wagner syndrome (hyaloideoretinal degeneration)-corneal degeneration, band-shaped keratopathy, hyaloideoretinal degeneration, narrowing of retinal vessels, retinal detachment, avascular preretinal membranes
110. Waldenström syndrome (macroglobulinemia syndrome)-crystalline corneal deposits, keratoconjunctivitis sicca, retinal venous thrombosis, retinal microaneurysms, cotton-wool spots
111. Weil disease (leptospirosis)-keratitis, retinitis
112. Werner syndrome (progeria of adults)-bullous keratitis, paramacular retinal degeneration
113. Wiskott-Aldrich syndrome (sex-linked draining ears, eczematoid, dermatitis, bloody diarrhea)-corneal ulcers, retinal hemorrhages
114. Yersiniosis corneal ulcer, retinal hemorrhages
115. Zellweger syndrome (cerebrohepatorenal syndrome)-corneal opacities, narrowing of retinal vessels, retinal holes without detachment, tapetoretinal degeneration
116. Zieve syndrome (hyperlipemia hemolytic anemia-icterus syndrome)-cloudy cornea, corneal ulcers, retinal lipemia


**CORNEAL DISEASES ASSOCIATED WITH DEAFNESS**
1. Atopic dermatitis-limbal keratitis, conjunctivitis
*2. Cogan syndrome (nonsyphilitic interstitial keratitis)-interstitial keratitis
3. Meniere disease-iritis, glaucoma
*4. Polyarteritis nodosa (Kussmaul disease)-paralimbal keratitis, corneoscleral ulceration
5. Sarcoidosis syndrome (Schaumann syndrome)-primary stromal keratitis, keratoconjunctivitis sicca
*6. Syphilis (lues)-interstitial keratitis
7. 3-methyl-pentyn-3-yl acid phthalate (Whipcide, trichuricidal agent)-keratitis, uveitis, stromal keratitis
8. Tuberculosis-interstitial keratitis
10. Wegener granulomatosis-necrotizing sclerokeratitis


**TRIGGER MECHANISMS FOR RECURRENT HERPES SIMPLEX KERATITIS**
*1. Corticosteroids (topical)
*2. Emotional disturbances
*3. Exposure to sunlight (ultraviolet)
*4. Fever (most common)
5. Gastrointestinal upsets
6. Ingestion of food to which patient is allergic
7. Mechanical trauma
8. Menses


**PREDISPOSING FACTORS IN KERATOMYCOSIS**

1. Antibiotics
2. Steroids
3. Trauma


**SCLEROCORNEA**

This type of developmental corneal abnormality has ill-defined limbus due to extension of opaque scleral tissue into the cornea. Vision varies with involvement. Somatic abnormalities include craniofacial, digital, skin, and testis abnormalities; deafness and mental retardation are seen.

1. Associated ocular abnormalities including:
   A. Abnormalities of Descemet membrane, endothelium, and corneal stroma
   B. Aniridia
   C. Cataract
   D. Coloboma
   E. Cornea plana-occurrence
   F. Dysgenesis of angle and iris
   G. Esotropia
   H. Glaucoma
   I. Iridocorneal synechiae
   J. Microphthalmia
   K. Nystagmus
   L. Persistent pupillary membrane
   M. Posterior embryotoxon

2. Associated syndromes, including the following:
   A. Axenfeld syndrome
   B. Cross syndrome
   C. Dandy-Walker syndrome
   D. Hallermann-Streiff syndrome
   E. Hurler syndrome
   F. Hypomelanosis of Ito
   G. Lobstein syndrome
POSTOPERATIVE CORNEAL MELT

This condition is characterized by central or peripheral thinning of corneal stroma after pterygium, refractive surgery, keratoplasty, glaucoma surgery, strabismus surgery, cataract surgery, or retina surgery.

1. AIDS
2. Collagen disorders
   a. Polyanarteritis nodosa
   b. Rheumatoid arthritis
   c. Scleroderma
   d. Systemic lupus erythematosus
3. Delayed corneal wound healing due to drugs, including adenine arabinoside, adrenal cortex injections, aldosterones, alpha-chymotrypsin, azathioprine, betarnethasone, butacaine, cocaine, cortisone, dexamethasone, fluorometholone, fluprednisolone, hydrocortisone, prednisolone, sulfacetamide, sulfamethizole, tetracaine, triamcinolone
4. Drug use
   a. Steroid therapy
   b. Stevens-Johnson syndrome
   c. Topical anesthetics abuse
   d. Topical nonsteroidal antiinflammatory agents
5. Marginal ring ulcer
   a. Acute leukemia
   b. Bacillary dysentery
   c. Brucellosis
   d. Dengue fever
   e. Gonococcal arthritis
   f. Hookworm infestation
   g. Influenza
   h. Possibly porphyria trigger mechanisms
6. Ocular predisposing factors
   a. Phlyctenular keratoconjunctivitis
   b. Superior limbic keratoconjunctivitis
c. Trachoma

d. Terrien disease

e. Vernal conjunctivitis

7. Preexisting corneal dellen

8. Skin conditions

  a. Acne rosacea
  b. Benign mucosa; pemphigoid
  c. Dermatitis herpetiformis
  d. Ectodermal dysplasia
  e. Herpes zoster
  f. Ichthyosis
  g. Incontinentia pigment
  h. Leprosy
  i. Lichen planus
  j. Lyell disease
  k. Lymphopathia venereum
  l. Molluscum contagiosum
  m. Palmoplantar keratosis
  n. Pemphigus foliaceus
  o. Pityriasis rubra pilaris
  p. Psoriasis

9. Systemic diseases

  a. Brucellosis
  b. Deerfly fever
  c. Filariasis
  d. Gout
  e. H. influenzae
  f. Hodgkin disease
  g. Hypoparathyroidism
  h. Leprosy
  i. Lymphopathia venereum
  j. Mumps
  k. Sarcoidosis
  l. Trypanosomiasis
  m. Tuberculosis
  n. Upper respiratory infection

10. Vitamin deficiency

  a. Ariboflavinosis keratopathy
  b. Pellagra
  c. Vitamin A deficiency
  d. Vitamin B<sub>2</sub> deficiency

CORNEAL MUCOUS PLAQUES

These plaques are abnormal collections of a mixture of mucous, epithelial cells, and proteinaceous and lipoidal material that adhere firmly to the corneal surface.

1. Local radiation exposure
2. Herpes zoster
3. Keratoconjunctivitis sicca
4. Other forms of keratitis
5. Vernal keratoconjunctivitis


Intraocular Pressure

CONTENTS
Glaucoma suspect, infant 305
Conditions simulating congenital glaucoma 305
Syndromes and diseases associated with glaucoma 307
Glaucoma suspect, adult 312
Elevated intraocular pressure measurement with normal-appearing optic disc 313
Secondary open-angle glaucoma 314
Unilateral glaucoma 315
Glaucoma associated with displaced lens 317
Glaucoma and elevated episcleral venous pressure 318
Glaucoma associated with shallow anterior chamber 318
Glaucoma in aphakia or pseudophakia 319
Medications and chemicals that may cause elevated intraocular pressure 320
Primary low-tension glaucoma 323
Neovascular glaucoma 324
Hypotony 325
Glaucoma associated with uveitis 326
Glaucoma associated with intraocular tumors 327

GLAUCOMA SUSPECT, INFANT

1. Amblyopia ex anopsia
2. Corneal edema (see p. 250)
3. Corneal enlargement
4. Cupping and atrophy of optic disc
5. Deep anterior chamber
6. Epiphora, photophobia, and blepharospasm (see p. 65-66)
7. Iridodonesis and subluxation of lens (see p. 374)
8. Iris processes
9. Tears in Descemet membrane (see p. 287)


CONDITIONS SIMULATING CONGENITAL GLAUCOMA

1. Blue sclera
   A. Albright hereditary osteodystrophy (pseudohypoparathyroidism)
   B. Craniofacial dysostosis (Crouzon disease)
C. de Lange syndrome
D. Ehlers-Danlos syndrome
E. Hilling syndrome
F. Hallermann-Streiff syndrome (oculomandibulodysecephaly)
G. Incontinentia pigmenti (Bloch-Sulzberger syndrome)
H. Juvenile Paget disease (hyperphosphatasia, hereditary)
I. Lowe (oculocerebrorenal) syndrome
J. Marfan syndrome
K. Turner (XO, gonadal dysgenesis) syndrome
L. van der Hoeve syndrome (osteogenesis imperfecta)
M. Werner syndrome

2. Corneal opacity
   A. Congenital malformations
      (1) Anterior corneal staphyloma
      (2) Cornea plana
      (3) Incontinentia pigmenti (Bloch-Sulzberger syndrome)
      (4) Norrie disease
      *(5) Peters anomaly
      (6) Riley-Day syndrome (congenital familial dysautonomia)
      *(7) Sclerocornea
      (8) Trisomy 13-15 syndrome (Patau syndrome)
   B. Edema
      *(1) Birth injury, such as breaks in Descemet membrane
      (2) Congenital hereditary corneal edema
      (3) Fetal uveitis
      *(4) Infectious keratitis (congenital syphilis, interstitial keratitis, rubella,
          variola, varicella, gonorrhea, mumps, and others)
      (5) Keratitis (chemical)
      *(6) Keratoconus
   C. Metabolic disorders/dystrophies
      (1) Congenital hereditary stromal dystrophy
      (2) Corneal amyloidosis (Lubarsch-Pick syndrome)
      (3) Corneal lipidosis
      (4) Cystinosis (Lignac-Fanconi syndrome)
      (5) Fabry disease
      (6) Hyperlipidemia
      (7) Mucopolysaccharidoses (MPS)
          a. Hunter syndrome (MPS IIB)
          b. Hurler syndrome (MPS IH)
          c. Maroteaux-Lamy syndrome (MPS VI)
          d. Morquio syndrome (MPS IV)
          e. Sanfilippo syndrome (MPS IIC)
          f. Scheie syndrome (MPS IS)
      (8) Porphyria (congenital)
      (9) von Gierke glycogen storage disease

3. Epiphora (excessive tearing)
SYNDROMES AND DISEASES ASSOCIATED WITH GLAUCOMA

1. Ocular disease
   A. Corneal endothelial disorders
      (1) Fuchs endothelial dystrophy
      (2) Iridocorneal endothelial (ICE) syndrome
         a. Chandler syndrome
         b. Cogan-Reese (iris-nevus) syndrome
         c. Progressive iris atrophy
      (3) Posterior polymorphous dystrophy
   B. Developmental glaucoma with associated ocular anomalies
      (1) Aniridia
      (2) Axenfeld-Rieger syndrome
      (3) Congenital ectropion uveae
      (4) Congenital iris hypoplasia
      (5) Megalocornea
      (6) Microcornea
      (7) Peters anomaly
      (8) Sclerocornea
   C. Elevated episcleral venous pressure (see p. 235)
   D. Iris disorders
      (1) Iridocyclitis
      * (2) Pigmentary glaucoma
   E. Lens disorders
      (1) Cataract
a. Lens-particle glaucoma  
b. Phacoanaphylaxis  
c. Phacolytic (lens protein) glaucoma  
d. Phacomorphic (intumescent lens) glaucoma  
(2) Dislocation of the lens  
*(3) Exfoliation syndrome  
F. Medications or chemicals  
(1) Corticosteroids  
(2) Others  
G. Myopia  
H. Ocular hemorrhage  
(1) Degenerated ocular blood  
   a. Ghost cell glaucoma  
   b. Hemolytic glaucoma  
   c. Hemosideric glaucoma  
*(2) Hyphema  
   a. Blunt trauma  
   b. Intraocular surgery  
1. Intraoperative  
2. Postoperative  
   c. Penetrating trauma  
   d. Spontaneous  
1. Anterior segment neovascularization  
2. Intraocular tumor  
3. Pupillary vascular tufts  
   (3) Orbital hemorrhage (massive)  
   (4) Vitreous hemorrhage (massive)  
I. Ocular inflammation  
(1) Choroiditis and retinitis  
   a. Cytomegalic inclusion retinitis  
   b. Sympathetic ophthalmia  
   c. Toxocariasis  
   d. Vogt-Koyanagi-Harada syndrome  
(2) Episcleritis  
*(3) Iridocyclitis  
   a. Acute anterior iridocyclitis  
   b. Ankylosing spondylitis  
   c. Behçet disease  
   d. Fuchs heterochromic cyclitis  
   e. Glaucomatocyclitic crisis (Posner-Schlossman syndrome)  
   f. Infectious diseases  
1. Acquired immunodeficiency syndrome (AIDS)  
2. Congenital rubella  
3. Disseminated meningococcemia  
4. Hansen disease (leprosy)  
5. Hemorrhagic fever with renal syndrome
6. Onchocerciasis (also keratitis)
7. Syphilis
   g. Juvenile rheumatoid arthritis (JRA)
   h. Pars planitis
   i. Precipitates on the trabecular meshwork (Grant syndrome)
   j. Reiter syndrome
   k. Sarcoid
   l. Trauma
(4) Keratitis
   a. Adenovirus type 10
   b. Herpes simplex
   c. Herpes zoster
   d. Interstitial keratitis
(5) Scleritis
*J. Ocular surgery
   (1) Aphakia or pseudophakia (see p. 405)
   (2) Corticosteroid induced
   (3) Cyclodialysis cleft (sudden closure)
   (4) Epithelial downgrowth
   (5) Malignant (ciliary block) glaucoma
   (6) Penetrating keratoplasty
      a. Distortion of angle structures
      b. Graft rejection
(7) Vitreoretinal procedures
   a. Intravitreal gas
   b. Pars plana vitrectomy
   c. Retinal photocoagulation
   d. Scleral buckling surgery
   e. Silicone oil
K. Ocular trauma
   (1) Chemical burns (acid, alkali, and others)
   (2) Contusion injuries
      a. Angle recession
      b. Hyphema
      c. Iritis
      d. Lens damage or dislocation
      e. Trabecular damage
   (3) Penetrating injuries
      a. Epithelial downgrowth
      b. Hyphema
      c. Lens damage or dislocation
      d. Peripheral anterior synechiae
   (4) Radiation damage
   (5) Retained intraocular foreign body (iron, copper)
   (6) Retrobulbar hemorrhage (massive)
L. Ocular tumors
(1) Benign tumors of the anterior uvea
   a. Adenomas
   b. Cysts (primary versus secondary)
   c. Iris nevi
   d. Leiomyomas
   e. Melanocytomas
   f. Melanoses
(2) Histiocytosis X
(3) Leukemias
(4) Lymphomas
(5) Metastatic tumors
   a. Carcinomas (most commonly, breast carcinoma in females and lung carcinoma in males)
   b. Melanomas
(6) Multiple myeloma
(7) Ocular tumors of childhood
   a. Juvenile xanthogranuloma
   b. Medulloepithelioma (diktyoma)
   c. Retinoblastoma
(8) Orbital tumors
(9) Primary uveal melanomas
(10) Retrobulbar tumors

M. Retinal, vitreous, and choroidal disorders
(1) Angle closure
   a. Acute choroidal hemorrhage
   b. Central retinal vein occlusion (CRVO)
   c. Ciliochoroidal effusion

1. AIDS
2. Arteriovenous malformations
3. Inflammatory conditions
4. Nanophthalmos
5. Surgery
6. Trauma
7. Tumors
8. Uveal effusion syndrome
   d. Hemorrhagic retinal and choroidal detachment
   e. Iris retraction syndrome with retinal detachment (Campbell)
   f. Persistent hyperplastic primary vitreous (PHPV)
   g. Postoperative panretinal photocoagulation
   h. Postoperative scleral buckle
   i. Retinal dysplasia
   j. Retinopathy of prematurity (retrolental fibroplasia)
*(2) Neovascular glaucoma (see p. 324)
(3) Retinitis pigmentosa
(4) Rhegmatogenous retinal detachment (Schwartz syndrome)

2. Systemic disorders
A. AIDS
B. Angioneurotic edema (giant urticaria)
C. Ankylosing spondylitis (Marie-Strümpell disease)
D. Aortic arch syndrome
E. Behçet disease
F. Carotid artery occlusive disease
G. Carotid-cavernous fistula
H. Cavernous sinus thrombosis
I. Crouzon disease (craniofacial dysostosis)
J. Cushing disease
K. Developmental glaucoma as part of a syndrome
   (1) Bing-Neel [macroglobulinemia and central nervous system (CNS)] syndrome
   (2) Chondrodystrophy, joint dislocation, glaucoma, and mental retardation
   (3) Chromosomal abnormalities
      a. Chromosome partial deletion (long-arm) syndrome
      b. Pericentric inversion of chromosome II
      c. Trisomy 21 (Down syndrome)
      d. Trisomy 16-18 (Edward syndrome)
      e. Trisomy F (17-18)
      f. Trisomy 13-15 (Patau syndrome)
      g. Turner syndrome (XO, gonadal dysgenesis)
      h. 9p syndrome
   (4) Cockayne syndrome
   (5) Congenital rubella syndrome
   (6) Cretinism (juvenile hypothyroidism)
   (7) Cystinosis
   (8) Dental-ocular-cutaneous syndrome
   (9) Diamond-Blackfan syndrome
   (10) Ehlers-Danlos syndrome
   (11) Familial histiocytic dermatarthritis syndrome
   (12) Fetal alcohol syndrome
   (13) Gorlin-Goltz (multiple basal cell nevi) syndrome
   (14) Hallermann-Streiff syndrome (oculomandibulofacial dyscephaly)
   (15) Homocystinuria
   (16) Kartagener syndrome (sinusitis-bronchiectasis-situs inversus)
   (17) Kimmelstiel- Wilson syndrome
   (18) Klinefelter syndrome
   (19) Klippel-Trenaunay-Weber syndrome
   (20) Krabbe syndrome
   (21) Krause syndrome (congenital encephalophthalmic dysplasia)
   (22) Lowe (oculocerebrorenal) syndrome
   (23) Marfan syndrome (arachnodactyly dystrophia mesodermalis congenita)
   (24) Meyer-Schwickerath-Weyers syndrome (oculodentodigital dysplasia)
   (25) Miller (Wilms aniridia) syndrome
(26) MPS
   a. Hurler syndrome (MPS IH)
   b. Maroteaux-Lamy syndrome (MPS VI)
   c. Morquio syndrome (MPS IV)
(27) Nieden (telangiectasia-cataract) syndrome
(28) Pierre Robin syndrome (micrognathia-glossoptosis) syndrome
(29) Prader-Willi syndrome (hypotonia, hypogonadism, obesity, and mental retardation)
(30) Rubella syndrome
(31) Rubinstein-Taybi (broad thumb) syndrome
(32) Silverman (battered-child) syndrome
(33) Stickler syndrome (hereditary progressive arthroophthalmopathy)
(34) Treacher-Collins syndrome
(35) Ullrich syndrome (dyscraniopygophalangy)
(36) Waardenburg syndrome
(37) Wagner syndrome
(38) Weber-Christian disease
(39) Weil-Marchesani syndrome
(40) X-linked mental retardation (XLMR) syndrome
(41) Zellweger (cerebrohepatorenal) syndrome

*L. Diabetes
M. Epidemic dropsy (argemone oil poisoning)
N. Giant cell arteritis
*O. Graves disease
P. Hemorrhagic fever with renal syndrome (nephropathia epidemicca)
*Q. Herpes simplex
*R. Herpes zoster
S. Histiocytosis X
T. JRA
U. Juvenile xanthogranuloma
V. Leukemia
W. Lymphoma
X. Medications or chemicals
Y. Metastatic carcinoma
Z. Metastatic melanoma
AA. Multiple myeloma
BB. Phakomatoses
   (1) Nevus of Ota (oculodermal melanocytosis)
   (2) Sturge-Weber syndrome (encephalotrigeminal angiomatosis)
   (3) von Hippel-Lindau disease
   (4) von Recklinghausen neurofibromatosis
CC. Reiter syndrome
DD. Retinoblastoma
EE. Retrobulbar tumors
FE Sarcoidosis
GG. Sickling disorders
HH. Superior vena cava (superior mediastinal) syndrome
II. Syphilis
JJ. Systemic corticosteroids
KK. Vogt-Koyanagi-Harada syndrome


**GLAUCOMA SUSPECT, ADULT**

1. Advanced age
   *2. Applanation reading 21 mm Hg or greater
3. Asymmetric intraocular pressures (IOPs)
4. Black race
5. Contusion-angle deformity glaucoma in the fellow eye
   *6. Diabetes mellitus
7. Diurnal fluctuation in IOP of 10 mm Hg or greater
8. Endothelial dystrophy of the cornea
   *9. Exfoliative syndrome (see p. 400)
10. Family history of glaucoma
   *11. Hemorrhage at optic disc margin
12. High myopia
13. IOP elevation following use of corticosteroids
14. Krukenberg spindle or dense trabecular pigment band
   *15. Prominent cupping of optic disc
      A. Asymmetry of cup-to-disc ratio
      B. Cup-to-disc ratio 0.4
      C. Cupping-to-disc margin
      D. Vertical elongation of cup
16. Retinal detachment (see p. 487)
   *17. Retinal vein occlusion (see p. 468-472)
18. Schiötz scale reading 4.0/5.5 or 6.25/7.5 or less
19. Thyrotropic exophthalmos
   *20. Visual field changes suggestive of glaucoma

ELEVATED INTRAOCULAR PRESSURE MEASUREMENT WITH NORMALAPPEARING OPTIC DISC

1. Acromegaly
2. Anesthesia
   A. Ketamine
   B. Nitrous oxide with intravitreal gas
   C. Succinylcholine
*3. Blepharospasm
4. Caffeine intake
5. Cardiopulmonary bypass surgery
*6. Dysthyroid ophthalmopathy
7. Elevation in hemoglobin concentration
8. Excessive water intake
9. High scleral rigidity and indentation (e.g., Schiötz) tonometry
10. Horizontal gaze position
11. Hyperthermia
12. Hyperthyroid
13. Marked emotional stress
14. Mechanical factors in checking IOP (e.g., by patient's hair interfering with applanation tonometer arm)
15. Medications or chemicals
   *A. Corticosteroids
   B. Cycloplegics
   C. Others
*16. Normal variation (ocular hypertension)
*17. Preglaucoma (IOP sufficiently elevated to cause damage to the optic nerve, but damage is not yet visible ophthalmoscopically)
18. Reduced gravity
19. Tight collar, short neck, obesity
20. Tobacco smoking
*21. Tonometer in need of calibration
22. Valsalva maneuver
23. Voluntary widening of palpebral fissure


SECONDARY OPEN-ANGLE GLAUCOMA
1. Corneal endothelial disorders
   A. Fuchs endothelial dystrophy
   B. Posterior polymorphous dystrophy

2. Elevated episcleral venous pressure (see p. 235)

3. Iris disorders
   A. Iridoschisis
   *B. Pigmentary glaucoma

4. Lens disorders
   A. Cataract
      (1) Lens particle glaucoma
      (2) Phacoanaphylaxis
      (3) Phacolytic (lens protein) glaucoma
   B. Displaced lens (see p. 317-318)
   *C. Exfoliation syndrome

5. Medications or chemicals
   *A. Corticosteroids
   B. Cycloplegic effect
   C. Others

6. Ocular hemorrhage

7. Ocular inflammation

8. Ocular surgery
   A. Alpha-chymotrypsin (enzyme glaucoma)
   *B. Corticosteroid induced
   C. Distortion of anterior chamber angle from limbal or keratoplasty sutures
   D. Early postoperative elevation of IOP following cataract surgery (especially in eyes with preexisting glaucoma)
   E. Hemorrhage
      (1) Degenerated ocular blood
         a. Ghost cell glaucoma
         b. Hemolytic glaucoma
         c. Hemosideric glaucoma
      *(2) Hyphema
      (3) Internal wound neovascularization (late postoperative hyphema)
      (4) Pseudophakia
         a. Anterior chamber intraocular lens (IOL), including uveitis, glaucoma, hyphema (UGH) syndrome
         b. Iris-fixated IOL
         c. Posterior chamber IOL (usually sulcus fixation)
      (5) Retrobulbar hemorrhage (massive)
   F. Inflammation
   G. Intravitreal gas
   H. Neodymium: yttrium-aluminum-aluminum-garnet (Nd:YAG) laser capsulotomy
   I. Pseudophakic pigmentary dispersion (e.g., with posterior chamber implant)
   J. Retained lens cortex
   *K. Retained viscoelastic
L. Silicone oil
M. Sudden closure of cyclodialysis cleft
N. Vitreous filling anterior chamber

9. Ocular trauma
   A. Chemical burns (acid, alkali, other)
   B. Contusion
      (1) Angle recession
      * (2) Hyphema
      (3) Iritis
      (4) Trabecular damage
   C. Radiation damage
   D. Retained intraocular foreign body (iron, copper)
   E. Retrobulbar hemorrhage (massive)

10. Ocular tumors
11. Retinal, vitreous, and choroidal disorders
    * A. Neovascular glaucoma-open-angle stage (see p. 324)
    B. Retinitis pigmentosa
    C. Rhegmatogenous retinal detachment (Schwartz syndrome)

12. Systemic, such as ocular amyloidosis


UNILATERAL GLAUCOMA

1. Corneal endothelial disorders
   A. Fuchs endothelial dystrophy with angle closure due to thickened peripheral cornea
   B. Iridocorneal endothelial (ICE) syndrome
      (1) Chandler syndrome
      (2) Iris-nevus (Cogan-Reese) syndrome
      (3) Progressive iris atrophy

2. Elevated episcleral venous pressure (see p. 235)

3. Lens disorders
   A. Cataract
      (1) Lens-particle glaucoma
      (2) Phacoanaphylaxis
      (3) Phacolytic (lens protein) glaucoma
      (4) Phacomorphic (intumescent lens) glaucoma
B. Displacement of the lens
   (1) Buphthalmos
   (2) Cataract (mature or hypennature)
   (3) Exfoliation syndrome
   (4) Intraocular tumor
   (5) PHPV
   (6) Sturge-Weber syndrome (encephalotrigeminal angiomatosis)
   (7) Trauma
   (8) Uveitis
C. Exfoliation syndrome (see p. 40)

4. Medications or chemicals
   A. Alpha-chymotrypsin (enzyme glaucoma)
   B. Chemical burns
   *C. Corticosteroids (topical or periocular)
   D. Cycloplegics (angle closure or open angle)
   E. Nitrous oxide inhalation with intraocular gas
   F. Urokinase (intraocular)
   G. Others

5. Ocular hemorrhage
6. Ocular inflammation
   *7. Ocular surgery
8. Ocular trauma
9. Ocular tumors
10. Retinal, vitreous, and choroidal disorders
    A. Angle closure
       (1) Acute choroidal hemorrhage
       *(2) CRVO
       (3) Ciliochoroidal effusion
          a. Arteriovenous malformations
          b. Inflammatory conditions
          c. Nanophthalmos
          d. Surgery
          e. Trauma
          f. Tumors
          g. Uveal effusion syndrome
       (4) Hemorrhagic retinal and choroidal detachment
       (5) PHPV
       *(6) Postoperative panretinal photocoagulation
       (7) Postoperative scleral buckle
       (8) Retinal dysplasia
       (9) Retinopathy of prematurity (retrolental fibroplasia)
   *B. Neovascular glaucoma (see p. 324)
      *(1) Diabetic retinopathy
      (2) Extraocular vascular disorders
         a. Carotid-cavernous fistula
         b. Carotid occlusive disease
c. Giant cell arteritis

(3) Ocular disorders-miscellaneous
   *a. Chronic glaucoma
   b. Endophthalmitis
   c. Intraocular malignancy
   d. Iris melanoma
   e. PHPV
   f. Photoradiation or helium ion irradiation for uveal melanoma
   g. Pseudophakia
   h. Sympathetic ophthalmia
   *i. Uveitis (chronic)

(4) Retinal disorders-miscellaneous
   a. Coats disease
   b. Eales disease
   c. Optic-nerve glioma with venous stasis
   d. Retinal detachment (usually chronic)
   e. Retinal vascular occlusive disorders

1. Retinal artery occlusion central or branch
   *2. Retinal vein occlusion central or branch
      f. Retinoblastoma
      g. Retinopathy of prematurity (retrolental fibroplasia)
      h. Retinoschisis
      i. Sickle cell retinopathy
   C. Open-angle glaucoma associated with rhegmatogenous retinal detachment
      (Schwartz syndrome)


GLAUCOMA ASSOCIATED WITH DISPLACED LENS

1. Alport syndrome
2. Aniridia
3. Axenfeld-Rieger syndrome
4. Buphthalmos
   *5. Cataract (mature or hypermature)
6. Cornea plana
7. Crouzon disease (craniofacial dysostosis)
8. Ectopia lentis et pupillae
9. Ehlers-Danlos syndrome
   *10. Exfoliation syndrome (see p. 400)
11. High myopia
GLAUCOMA AND ELEVATED EPISCLERAL VENOUS PRESSURE

1. Arteriovenous fistulas
   A. Carotid-cavernous sinus fistulas
      (1) Spontaneous
      (2) Traumatic
   B. Orbital-meningeal shunts
   C. Orbital varices
      *D. Sturge-Weber syndrome (encephalotrigeminal angiomatosis)

2. Idiopathic elevation of episcleral venous pressure
   *A. Familial
   B. Sporadic

3. Venous obstruction
   A. Cavernous sinus thrombosis
      *B. Congestive heart failure
   C. Episcleral
      (1) Chemical burns (acid, alkali, and others)
      (2) Radiation
D. Jugular venous obstruction
E. Ocular amyloidosis
F. Orbital
  *(1) Dysthyroid
  (2) Orbital vein thrombosis
  (3) Phlebitis
  (4) Pseudotumor
  (5) Retrobulbar tumor
G. Pulmonary venous obstruction
H. Superior vena cava (superior mediastinal) syndrome


**GLAUCOMA ASSOCIATED WITH SHALLOW ANTERIOR CHAMBER**

*1. Primary-angle closure*
   A. Plateau iris syndrome
   *B. Relative pupillary block (most common)*

*2. Secondary-angle closure*
   *A. CRVO (central retinal vein occlusion)*
   B. Choroidal hemorrhage (acute)
   C. Ciliochoroidal effusion
     (1) AIDS
     (2) Arteriovenous malformations
     *(3) Inflammation*
     (4) Nanophthalmos
     (5) Trauma
     (6) Tumor
     (7) Uveal effusion syndrome

D. Cystinosis
E. Drug-induced acute transitory myopia (diuretics, sulfonamides, and others)
F. Elevated episcleral venous pressure associated with arteriovenous fistula
G. Fuchs endothelial dystrophy-with peripheral corneal thickening
H. Hemorrhagic retinal and choroidal detachment
I. Hyperglycemia (acute)
J. Inflammation
   (1) Episcleritis
   (2) Iridocyclitis with posterior synechiae and iris bombe
   (3) Posterior scleritis
K. Intraocular tumor (posterior segment melanoma, metastatic carcinoma, retinoblastoma, medulloepithelioma, and others)
L. Lens dislocation (see p. 401-404)
M. Luetic interstitial keratitis
N. Malignant (ciliary block) glaucoma
O. Maroteaux-Lamy syndrome (MPS VI)
P. Multiple cysts of the iris and ciliary body
Q. Nanophthalmos
R. Pars plana vitrectomy
S. PHPV
*T. Phakic or aphakic pupillary block
U. Phakomorphic (intumescent lens) glaucoma
V. Postoperative panretinal photocoagulation
W. Postoperative scleral buckle
X. Pupil dilatation, including topical dilatation and systemic decongestants, bronchodilators, gastroenterologic and genitourinary disorders.
Y. Retinal dysplasia
Z. Retinopathy of prematurity (retrolental fibroplasias)


**GLAUCOMA IN APHAKIA OR PSEUDOPHAKIA**

1. Alpha-chymotrypsin (enzyme glaucoma)
2. Ciliary-block (malignant) glaucoma
*3. Corticosteroid induced
4. Degenerated intraocular blood
   A. Ghost cell glaucoma
   B. Hemolytic glaucoma
   C. Hemosideric glaucoma
5. Distortion of the anterior chamber angle by limbal sutures
*6. Early postoperative pressure elevation (especially in eyes with preexisting glaucoma)
7. Epithelial downgrowth
8. Fibrous proliferation
*9. Following Nd:YAG capsulotomy
10. Hyphema
   A. Internal wound neovascularization (late postoperative hyphema)
   B. Pseudophakia
      (1) Anterior chamber IOL (including the UGH syndrome)
(2) Iris-fixated IOL
(3) Posterior-chamber IOL (usually sulcus fixation)

11. Inflammation
12. Peripheral anterior synechiae
13. Primary open-angle glaucoma
14. Pseudophakic pigmentary dispersion
*15. Pupillary block
*16. Retained lens cortex
*17. Retained viscoelastic
18. Vitreous filling the anterior chamber
19. Vitreous hemorrhage (massive)


**MEDICATIONS AND CHEMICALS THAT MAY CAUSE ELEVATED INTRAOCULAR PRESSURE**

1. Anesthetic agents
   A. Ketamine
   B. Nitrous oxide (inhalation, especially in eyes with retinovitreal surgery and intraocular gas)
2. Anticholinergics/parasympatholytics
   A. Antidepressants
      (1) Amitriptyline (Elavil)
      (2) Imipramine (Tofranil)
      (3) Nortriptyline (Pamelor)
      (4) Protriptyline (Vivactil)
      (5) Trimipramine (Surmontil)
   B. Antihistamines
      (1) Anazolene (Vasocon-A)
      (2) Brompheniramine (Dimetane)
      (3) Cyclizine (Marezine)
      (4) Cyproheptadine (Periactin)
      (5) Diphenhydramine (Benadryl)
      (6) Orphenadrine (Norgesic)
      (7) Tripelennamine (Pyribenzamine)
   C. Antiparkinson medications
      (1) Biperiden (Akineton)
      (2) Cycrimine (Pagitane)
      (3) Trihexyphenidyl hydrochloride (Artane)
   D. Antispasmodic agents
(1) Dicyclomine (Bentyl)
(2) Diphenamid methylsulfate (Prantal)
(3) Hexocyclium methylsulfate (Tral)
(4) Hyoscyamine (Donnatal, Donnagel)
(5) Mepenzolate (Cantil)
(6) Methscopolamine bromide (Pamine)
(7) Oxyphenonium bromide (Antrenyl)
(8) Propantheline bromide (Pro-Banthine)
(9) Tridihexethyl chloride (Pathilon)

*E. Cycloplegics
    (1) Atropine
    (2) Cyclopentolate (Cyclogyl)
    (3) Homatropine
    (4) Tropicamide (Mydriacyl)
    (5) Scopolamine (Hyoscine)

F. Miscellaneous
    (1) Atropine (systemic)
    (2) Glycopyrrolate (Robinul)

G. Phenothiazine
    (1) Doxepin (Sinequan)
    (2) Haloperidol (Haldol)
    (3) Prochlorperazine (Compazine)
    (4) Promethazine (Phenergan)
    (5) Triflupromazine (Vesprin)

H. Poisoning
    (1) Belladonna
    (2) Jimson weed

3. Argemone oil (epidemic dropsy)
4. Caffeine
5. Carbon dioxide inhalation
6. Carmustine injection
7. Chemical burns
   A. Acid
      (1) Chromic acid
      (2) Hydrochloric (muriatic) acid
      (3) Sulfuric (battery) acid
   B. Alkali
      (1) Ammonium hydroxide (ammonia)
      (2) Calcium hydroxide (lime)
      (3) Sodium hydroxide (lye)
   C. Dibent[b.f][1,4] oxazepine (CR tear gas)
   D. Formaldehyde gas (in aqueous solution formalin)

8. CNS stimulants/anorexics
   A. Dextroamphetamine
   B. Methamphetamine
   C. Phenmetrazine (Preludin)
D. Phentermine (Ionamin)

*9. Corticosteroids

* A. Ocular (topical)
(1) Dexamethasone (Decadron, Maxidex)
(2) Fluorometholone (FML, Flarex)
(3) Prednisolone acetate (Pred Forte)
(4) Prednisolone sodium phosphate (Inflamase)

*B. Subconjunctival depot injection
(1) Methylprednisolone acetate
(2) Triamcinolone

C. Systemic
(1) Betamethasone (Celestone)
(2) Cortisone acetate
(3) Dexamethasone (Decadron)
(4) Hydrocortisone (Cortef, Solu-Cortef)
(5) Methylprednisolone (Medrol)
(6) Prednathasone (Haldron)
(7) Prednisolone
(8) Prednisone (Deltasone)
(9) Triamcinolone (Aristocort)

10. Idiopathic lens swelling
A. Acetylsalicylic acid (aspirin)
B. Sulfanilamide
C. Others

11. Intraocular injection
A. Alpha-chymotrypsin (enzyme glaucoma)
B. Urokinase
*C. Viscoelastic (Healon, others)

12. Methylphenidate (Ritalin)

13. Miotics
A. Carbachol
B. Demecarium (Humorsol)
C. Echothiophate (phospholine iodide)
D. Pilocarpine

14. Succinylcholine (Anecline)

15. Sympathomimetics
A. Ephedrine
B. Mydriatics
(1) Dipivalyl epinephrine (Propine)
(2) Epinephrine (many products)
(3) Hydroxyamphetamine (paredrine)
(4) Phenylephrine (Neo-Synephrine)
C. Naphazoline (Naphcon)
D. Pheniramine maleate (Naphcon-A)
E. Phenylephrine (Neo-Synephrine)
F. Tetrahydrozoline (Visine)
16. Testosterone
17. Vasodilators
   A. Elevation of IOP following subconjunctival injection
      (1) Bamethan (Bupatol)
      (2) Isoxsuprine (Vasodilan)
      (3) Tolazoline (Priscoline)
      (4) Triaziquone (Trenimon)
   B. Amyl nitrite (Vaporole)
18. Water (excessive intake)
    May be potentiated by monoamine oxidase inhibitors such as phenelzine, pargyline, or tranylcypromine.


**PRIMARY LOW-TENSION GLAUCOMA**

1. Nonglaucomatous optic nerve disorders resembling glaucomatous damage
   A. Developmental abnormalities
      (1) Colobomas of the optic-nerve head, including optic pits
      *(2) Large physiologic cups
      (3) Tilted discs
   B. Nonglaucomatous causes of acquired cupping
      (1) Compressive lesions
         a. Aneurysm
         b. Chiasmic arachnoiditis
         c. Cyst
         d. Tumor
      *(2) Ischemic optic neuropathy (especially arteritic)
   *C. Nonglaucomatous causes of nerve fiber bundle defects on visual field testing
      (1) Chorioretinal lesions
         a. Chorioretinitis
         b. Retinal vascular occlusions
         c. Tumors
      (2) Optic-nerve head lesions
         a. Colobomas
         b. Drusen
         c. Other
      (3) Posterior lesion of the visual pathway
         a. Meningioma
         b. Pituitary tumor
         c. Pseudotumor
         d. Other
2. Undetected high-pressure glaucoma
   A. Corneal edema giving false low measurement of IOP with applanation (e.g., Goldman or Perkins) tonometry
   B. Intermittent elevation of IOP causing damage (IOP normal at time of examination)
      (1) Glaucomatocyclitic crisis (Posner-Schlossman syndrome)
      (2) Intermittent angle closure
      (3) Others
   C. Low scleral rigidity giving false low measurement of IOP with indentation (e.g., Schiötz) tonometry
   *D. Prior elevation in pressure resulting in optic nerve damage
      (1) Burned-out open-angle glaucoma
      (2) Corticosteroids
      (3) Pigmentary glaucoma
      (4) Trauma
      (5) Uveitis
   E. Wide diurnal variation (multiple measurements at different times of day required to rule-out high-pressure glaucoma)


**NEOVASCULAR GLAUCOMA**

*1. Diabetic retinopathy
2. Extraocular vascular disorders
   A. Aortic arch syndrome
   *B. Carotid artery occlusive disease
   C. Carotid-cavernous fistula
   D. Giant cell arteritis
3. Ocular disorders-miscellaneous
   A. Chronic glaucoma
   B. Endophthalmitis
   C. Iris melanoma
   D. PHPV
   E. Pseudophakia
   F. Sympathetic ophthalmia
   *G. Uveitis
4. Retinal disorders-miscellaneous
   A. Choroidal melanoma
   B. Coats exudative retinopathy
   C. Eales disease
D. Metastatic carcinoma
E. Norrie disease
F. Optic nerve glioma with subsequent venous stasis retinopathy
G. Photoradiation or helium ion irradiation for uveal melanoma
*H. Retinal detachment (usually chronic)
I. Retinal vascular occlusive disorders
   (1) Branch retinal artery occlusion.
   *(2) Branch retinal vein occlusion
   *(3) Central retinal artery occlusion
   *(4) Central retinal vein occlusion
J. Retinoblastoma
K. Retinopathy of prematurity (retrolental fibroplasia)
L. Retinoschisis
M. Sickle cell retinopathy
N. Stickler syndrome (inherited vitreoretinal degeneration)
O. Syphilitic retinal vasculitis


**HYPOTONY**

*1. Essential hypotension
2. Secondary hypotony
   A. Cartilaginous-arthritic-ophthalmic deafness
   B. Ciliochoroidal detachment
      (1) Chorioretinal inflammation
      (2) Ocular neoplasm
      *(3) Trauma, including ocular surgery
   *C. Cyclitis
   *D. Cyclodialysis
E. Decreased IOP from medications and chemicals
   (1) Alcuronium
   (2) Aminophylline (intravenous)
      *(3) Carbonic anhydrase inhibitors (e.g., acetazolamide, methazolamide, ethoxzolamide)
   (4) Cardiac glycosides (digitoxin, digoxin, lanatoside-C, ouabain)
   (5) Dibenamine
   (6) Dihydroergotoxine (Hydergine)
   (7) HPMPC (cidofovir)
   (8) Hyperosmotics (urea, glycerin, mannitol, ascorbic acid, glycerol, ethanol, trometamol)
   (9) Isosorbide
   (10) Pargyline (Eutonyl)
   (11) Phentolamine (Regitine)
(12) Propranolol (Inderal)
(13) Thiopental (Pentothal)

*F. Deep anesthesia
G. Deep coma and severe cerebral disease
*H. Dehydration-severe (e.g., cholera, dysentery, diabetic coma)
*I. Diabetic coma
*J. Glaucoma medications (beta-blockers, sympathomimetics, miotics, carbonic anhydrase inhibitors)
K. Hilding syndrome
L. Homer syndrome
M. Hyperosmolarity
N. Intestinal perforation or obstruction
O. Intraocular lens mal position
P. Irradiation
Q. Morquio-Brailsford syndrome (MPS IV)
R. Myotonic dystrophy
S. Ocular ischemia
*T. Ocular trauma with or without visible ciliary body injury
*U. Perforating ocular trauma
*V. Phthisis
W. Postencephalitic syndrome
*X. Postoperative surgical procedures especially for glaucoma
Y. Raeder syndrome
*Z. Retinal detachment
*AA. Systemic hypotension-severe (circulatory collapse, medications)
BB. Uremic coma
*CC. Wound leak


GLAUCOMA ASSOCIATED WITH UVEITIS

1. Arthritis
2. Juvenile rheumatoid arthritis
3. Ankylosing spondylitis, Reiter syndrome, psoriatic arthritis
6. Fuchs heterochromic uveitis
7. Posner-Schlossman syndrome (glaucomatocyclitic crisis)
8. Herpes simplex
9. Herpes zoster
10. Hansen disease
11. Rubella
12. Mumps and other viral infections
13. Gnathostomiasis
14. Traumatic uveitis
15. Phacoanaphylactic glaucoma
16. Phacolytic glaucoma
17. Pseudophakic-inflamatory glaucoma
18. Intermediate uveitis (pars planitis)
19. Posterior uveitis
   a. Ocular toxoplasmosis
   b. Acute retinal necrosis
   c. AIDS
   d. Other posterior uveitides
20. Panuveitis
   a. Sarcoidosis
   b. Behçet syndrome
   c. Sympathetic ophthalmia
   d. Vogt-Koyanagi-Harada syndrome
   e. Congenital syphilis
   f. Acquired syphilis
   g. Tuberculosis
   h. Onchocerciasis
21. Masquerade syndromes
   a. Intraocular neoplasia (uveal malignant melanoma, intraocular lymphoma, and others)
   b. Retinal detachment
22. Open-angle glaucoma
23. Angle-closure glaucoma
24. Combined-mechanism glaucoma


GLAUCOMA ASSOCIATED WITH INTRAOCULAR TUMORS

1. Iris
   A. Nevus
   B. Melanocytoma
   C. Iris pigment epithelium adenoma
   D. Malignant melanoma
   E. Metastatic
2. Ciliary body
   A. Medulloepithelioma
   B. Melanocytoma
C. Malignant melanoma
D. Metastatic

3. Choroid
   A. Malignant melanoma
   B. Metastatic

4. Optic nerve
   A. Melanocytoma
   B. Metastatic

5. Retina-retinoblastoma

6. Metastatic
   A. Carcinoma
   B. Cutaneous melanoma
   C. Breast
   D. Lung
   E. Kidneys
   F. Testicles
   G. Prostate
   H. Pancreas
   I. Colon
   J. Gastrointestinal

7. Others
   A. Leukemia
   B. Lymphoma
   C. Phakomatoses as Sturge-Weber or neurofibromatosis
   D. Multiple myelomas
   E. Juvenile xanthogranuloma


Hypopyon 329
Hyphema 331
Spontaneous hyphema 332
    Diagnostic table 334
Spontaneous hyphema in infants 333
    Diagnostic table 336
Plasmoid aqueous 338
Cholesterolosis of the anterior chamber 338
Gas bubbles in the anterior chamber 338
Pigmentation of trabecular meshwork 338
Pigment liberation into the anterior chamber with dilatation of pupil 339
Grading of anterior chamber angle width 339
Blood in Schlemm canal 339
Deep anterior chamber angle 340
Narrow anterior chamber angle 340
Irregular depth of the anterior chamber 341
Peripheral anterior synechiae 341
Neovascularization of anterior chamber angle 342
Iris processes 343
White mass in anterior chamber 343

HYPOPYON (PUS IN ANTERIOR CHAMBER)

1. Hypopyon ulcer-corneal ulcer with pus in the anterior chamber
   A. Acanthamoeba
   B. Acquired immunodeficiency syndrome (AIDS)
   C. Aspergillus species
   D. Candida albicans
   E. Chemical injury
   F. Diplococcus pneumoniae
   G. Escherichia coli
   H. Fusarium species
   I. Herpes simplex
   J. Herpes zoster
   K. Measles
   L. Moraxella
   M. Neisseria gonorrhoeae
   N. Proteus vulgaris
   *O. Pseudomonas aeruginosa
   P. Serratia species
   Q. Smallpox
   R. Spitting-cobra venom
S. Staphylococcus
*T. Streptococcus

2. Severe acute iridocyclitis
3. Repeated corneal transplantation of human amniotic membrane
4. Necrosis of intraocular tumors or metastasis
5. Retained intraocular foreign bodies, including toxic lens syndrome
6. Endophthalmitis—at time of surgical treatment, accidental trauma, in drug users, or spontaneous occurrence (see p. 223)
   A. Acanthamoebae
   B. Actinomycosis
   C. Amebiasis
   D. Aspergillosis species
   E. Bacterial including bacillus cereus
   F. Behçet syndrome
   G. C. albicans
   H. Coccidioidomycosis
   I. Coenurosis
   J. Cysticercosis
   K. Fusarium species
   L. Hydatid cyst
   M. Influenza
   N. Listeria monocytogenes
   O. Lockjaw (Clostridium tetani)
   P. Metastatic bacterial endophthalmitis
   Q. Moraxella species
   R. Mucor species
   S. Mycobacterium avium
   *T. Pseudomonas species
   U. Relapsing fever
   V. Serratia marcescens
   W. Saprophytic fungi
   *X. Staphylococcus
   *Y. Streptococcus
   Z. Sterile hypopyon

   (1) Behçet syndrome (oculobuccogenital syndrome)
   (2) Endotoxin contamination of ultrasonic bath
   (3) Following cyanoacrylate sealing of a corneal perforation
   (4) Following refractive surgery
   (5) Histiocytosis X (Hand-Schüller-Christian syndrome)
   (6) Intraocular lens or instrument polishing compounds or sterilization techniques
   *(7) Juvenile rheumatoid arthritis
   (8) Laser iridotomy
   (9) Leukemia
   *(10) Reaction to lens protein
   (11) Rough intraocular lens edges
(12) von Bechterev-Strumpel syndrome (rheumatoid spondylitis)
AA. Stevens-Johnson syndrome (dermatostomatitis)
BB. Tight contact lens or contact lens overwear syndrome
CC. Tuberculosis
DD. Weil disease (leptospirosis)
EE. Yersiniosis

6. Drugs, including the following:
   - benoxinate
   - butacaine
   - cocaine
   - colchicine (?)
   - dibucaine
   - dyclonine
   - ferrocholinate
   - ferrous fumarate
   - ferrous gluconate
   - ferrous succinate
   - ferrous sulfate
   - iodide and iodine solutions and compounds
   - iron dextran
   - iron sorbitex
   - phenacaine
   - piperocaine
   - polysaccharide-iron complex
   - radioactive iodides
   - rifabutin
   - tetracaine urokinase
   - urokinase

7. Vitreous "fluff-ball"
8. Following refractive surgery
9. Pseudohypopyon
   A. Ghost cell glaucoma with khaki-colored cells
   B. Accidental intraocular steroid injection
10. Acute angle-closure glaucoma
11. Non-Hodgkin lymphoma
12. Pars plana vitrectomy and silicone oil injection


**HYPHEMA (BLEEDING INTO THE ANTERIOR CHAMBER)**
1. Trauma
   A. Following laser iridectomy or strabismus surgery in aphakia
   B. Honan balloon use in Fuchs heterochromic iridocyclitis
   *C. Tear of ciliary body-post contusion deformity of anterior chamber
   *D. To ciliary body, such as cyclodialysis
   E. To iris, such as in iridodialysis or intraocular lens irritation
   F. After airbag inflation
   G. Metallic intraocular foreign body during magnetic resonance imaging
2. Overdistention of vessels
   A. Obstruction of central retinal vein
   B. Sudden lowering of high intraocular pressure
3. Fragility of vessel walls
   A. Acute gonorrheal iridocyclitis
   B. Acute herpes iridocyclitis
   C. Acute rheumatoid iridocyclitis
   D. Ankylosing spondylitis
4. Blood abnormality
   A. Anemias
   B. Association with use of aspirin
   C. Hemophilia
   D. Leukemia
   E. Purpura
   F. Sickle cell disease
5. Metabolic disease
   A. Diabetes mellitus (Willis disease)
   B. Scurvy (avitaminosis C)
6. Neovascularization of iris (see rubeosis iridis, p. 366-367)
7. Vascularized tumors of iris (see pigmented and nonpigmented iris lesions, p. 374-375)
   A. Angioma
   B. Iris vascular tufts
   *C. Juvenile xanthogranuloma (JXG)
   D. Lymphosarcoma
   E. Retinoblastoma
8. Wound vascularization following cataract extraction
9. Persistent pupillary membrane hemorrhage


SPONTANEOUS HVPEMA

1. Delayed following glaucoma surgery
2. Diseases of blood or blood vessels
   A. Hemophilia
   B. Leukemia
   C. Malignant lymphoma
   D. Purpura
   E. Scurvy
3. Fibrovascular membranes in retrolenticular or zonular area
   A. Persistent primary vitreous
   B. Retinoschisis
   C. Retinopathy of prematurity
4. Systemic hypertension
5. Hydrophthalmos
6. Iatrogenic
7. Intraocular neoplasms
   *8. JXG-yellow nodules of skin and iris
9. Malignant exophthalmos
10. Microbial keratitis, especially Moraxella
11. Occult trauma or trauma with late effect
   *12. Rubeosis iridis
13. Severe iritis with or without
   A. Behçet disease (dermatostomatoophthalmic syndrome)
   B. Diabetes mellitus (Willis disease)
   C. Gonococcal infection
   D. Herpes zoster or herpes simplex
14. Use of warfarin, heparin, aspirin, or alcohol
15. Vascular anomalies of iris
16. Wound vascularization following cataract extraction


SPONTANEOUS HYPHEMA IN INFANTS

1. Acute rheumatoid iridocyclitis
2. Blood dyscrasias, such as anemia, leukemia, and disseminated intravascular coagulation
3. Iritis
*4. JXG
5. Perinatal asphyxia
6. Persistent hyperplastic primary vitreous
7. Retinoblastoma
8. Retinoschisis
9. Retinopathy of prematurity
*10. Trauma without history (consider child abuse)


**Extracted Table Spontaneous Hyphema**

**Extracted Table Spontaneous Hyphema in Infants**

PLASMOID AQUEOUS (AQUEOUS WITH A HIGH PROTEIN CONTENT)

1. Rheumatoid arthritis
2. Serum sickness
3. Infection with gonococcus
4. Following paracentesis or intraocular operation, such as cataract extraction
5. Severe corneal ulceration
6. Trauma


**CHOLESTEROLOSIS OF THE ANTERIOR CHAMBER**

In this condition, cholesterol crystals develop in the anterior chamber; usually in a blind eye following trauma, but can be associated with hyphema or secondary glaucoma. It is also associated with the following:

*1. Chronic uveitis*
2. Eales disease (periphlebitis)
3. Lens subluxation
4. Mature or hypermature cataract
5. Microphthalmia
*6. Phthisis bulbi
7. Retinal detachment
8. Traumatic cataract
9. Vascular disorders
10. Vitreous hemorrhage


**GAS BUBBLES IN THE ANTERIOR CHAMBER**

1. *Clostridium perfringens*
2. *E. coli*
3. Yttrium-aluminum-garnet (YAG) laser treatment to the anterior segment
4. Postoperative intraocular surgery


**PIGMENTATION OF TRABECULAR MESHWORK**

1. In elderly individuals-inferior nasal or faint band circumferential
*2. Pseudoexfoliation of lens with or without glaucoma-unilateral or bilateral
*3. Pigmentary glaucoma
*4. Krukenberg spindle without glaucoma
*5. Malignant melanoma-one eye
6. Cyst of pigment layer of iris-unilateral
7. Previous intraocular operation, inflammation, or hyphema-scattered, mostly in lower angle
8. Nevus-dense, isolated patch
9. Open-angle glaucoma-patchy band, whole circumference
10. Following gamma irradiation for malignancy of nasal sinus


**PIGMENT LIBERATION INTO THE ANTERIOR CHAMBER WITH DILATATION OF PUPIL**

1. Diabetes mellitus (Willis disease)
2. Exercise
3. Hurler disease (mucopolysaccharidoses IH)
4. Low-tension glaucoma with pigment dispersion


**GRADING OF ANTERIOR CHAMBER ANGLE WIDTH (USUALLY DETERMINED BY GONIOSCOPY)**

1. Grade 0: No angle structures visible-narrow angle, complete or partial closure (angle closure)
2. Grade 1: Unable to see posterior one half of trabecular meshwork-extremely narrow angle (probably capable of angle closure)
3. Grade 2: Part of Schlemm canal is visible-moderately narrow angle (may be capable of angle closure)
4. Grade 3: Posterior portion of Schlemm canal is visible-moderately open angle (incapable of angle closure)
5. Grade 4: Ciliary body is visible-open angle (incapable of angle closure)


**BLOOD IN SCHLEMM CANAL (REVERSAL OF NORMAL PRESSURE GRADIENT)**

*1. Artifact of goniolens flange occluding the episcleral veins in one or more quadrants
2. High episcleral venous pressure
   *A. Carotid-cavernous sinus fistula (Red-eyed shunt syndrome)
   *B. Dural-cavernous fistula
   C. Mediastinal tumors
   D. Orbital arteriovenous fistula
   E. Sturge-Weber syndrome (meningocutaneous syndrome)
   F. Superior vena cava obstruction (superior vena cava syndrome)
   G. Tetralogy of Fallot
3. Low intraocular pressure
   A. Following trabeculectomy
B. Hypotony (see p. 325)
C. Intraocular inflammation

4. Normal eye


DEEP ANTERIOR CHAMBER ANGLE

1. Normal variation
2. Aphakia
3. Myopia
4. Megalocornea or conical cornea including keratoconus (see p. 288)
5. Congenital glaucoma
6. Posterior dislocation of the lens (see p. 401-404)
7. Recession of anterior chamber angle


NARROW ANTERIOR CHAMBER ANGLE (MAY BE CAPABLE OF ANGLE CLOSURE GLAUCOMA)

1. Normal variation
*2. Predisposition to angle closure
3. Anterior dislocation of the lens
4. Hyperopia
5. Spherophakia and microcornea
6. Postoperative intraocular operation with leaking wound (see hypotony, p. 325)
7. Choroidal detachment (see p. 532-535)
*8. Pupillary block
9. Loss of aqueous from perforating ulcer, corneal wound, or staphyloma (see hypotony, p.325)
10. Intumescent senile cataract
11. Traumatic cataract that fluffs up
12. Primary hyperplastic primary vitreous (PHPV)
13. Peripheral anterior synechiae (see p. 341)
*14. Posterior entrapment of aqueous humor (malignant glaucoma or ciliary-block glaucoma)
15. Drugs, including the following:
- acetazolamide
- acetylcholine
- alpha-chymotrypsin
- demecarium
- dichlorphenamide
- echothiophate
- edrophonium
- ethoxzolamide
- isoflurophate
- methazolamide

- neostigmine
- physostigmine
- pilocarpine
- sulfacetamide
- sulfachlorpyridazine
- sulfadiazine
- sulfadimethoxine
- sulfamerazine
- sulfameter
- sulfamethazine

- sulfamethizole
- sulfamethoxazole
- sulfamethoxypyridazine
- sulfanilamide
- sulfaphenazole
- sulfapyridine
- sulfasalazine
- sulfathiazole
- sulfisoxazole

*16. Plateau iris

17. Diffuse ciliary body or iris tumor


**IRREGULAR DEPTH OF THE ANTERIOR CHAMBER**

1. Partial dislocation of lens
2. Tumor of iris or ciliary body
3. Peripheral anterior synechiae on one side of the chamber (see p. 341)
4. Iris bombe or pupillary block
5. Ruptured lens capsule with swelling on one side
6. Anatomic narrowing superiorly
7. Subacute angle-closure glaucoma
8. Cyclodialysis and traumatic recession of chamber angle


**PERIPHERAL ANTERIOR SYNECHIAE (ADHESION OF IRIS TISSUE ACROSS ANTERIOR CHAMBER STRUCTURES IN VARIABLE AMOUNTS NOTED WITH GONIOSCOPY)**

1. Bridge corneoscleral trabecular meshwork to Schwalbe line or anterior to Schwalbe line (uncommon)
A. Anterior chamber cleavage syndrome
   (1) Axenfeld syndrome (posterior embryotoxon)
   (2) Congenital central anterior synechiae
   (3) Following intraocular lens implantation
   (4) Reiger syndrome (dysgenesis mesostromatolysis)

B. Essential iris atrophy (see p. 373)
C. Iris bombe from occlusion of pupil
D. Iris or ciliary body tumor pushing iris into contact with cornea
E. Local adhesion with of epithelium or fibrous ingrowth
F. Penetrating injury of the cornea
G. Postoperative flat anterior chamber

2. Synechiae of iris limited to ciliary band, scleral spur, and trabecular meshwork (common)
   *A. Following cataract surgery, intraocular implantation, refractive surgery, or laser trabeculoplasty
   *B. Intraocular inflammation
   *C. Neovascular glaucoma from fibrovascular membrane (see p. 324)
   *D. Sequelae to angle-closure glaucoma


**NEOVASCULARIZATION OF ANTERIOR CHAMBER ANGLE (NEWLY FORMED VESSELS EXTEND INTO THE TRABECULAR MESHWORK)**

1. Anterior chamber angle
   A. Congenital pupillary iris lens membrane with goniodysgenesis
   B. Traumatic chamber angle

2. Iris tumors
   A. Hemangioma
   B. Melanoma
   C. Metastatic carcinoma

3. Ocular vascular disease
   *A. Central retinal artery thrombosis
   *B. Central retinal vein thrombosis (see p. 468)
   C. Hemiretinal branch vein occlusion (HBVO)

4. Postinflammatory
   A. Anterior chamber implants
   B. Fungal endophthalmitis
C. Radiation  
D. Retinal detachment operation  
E. Uveitis, chronic  

5. Proximal vascular disease  
A. Aortic arch syndrome (Takayasu syndrome)  
B. Carotid cavernous fistula  
C. Carotid ligation  
D. Carotid occlusive disease  
E. Cranial arteritis (temporal arteritis syndrome)  

6. Retinal disease  
A. Coats disease (Leber miliary aneurysms)  
*B. Diabetic retinopathy  
C. Eales disease (periphlebitis)  
D. Glaucoma, chronic  
E. Melanoma of choroid  
F. Norrie disease (fetal iritis syndrome)  
G. Persistent hyperplastic primary vitreous  
H. Retinal detachment  
I. Retinal hemangioma  
J. Retinal vessel occlusion  
K. Retinoblastoma  
L. Retrolental fibroplasia  
M. Sickle cell retinopathy (Herrick syndrome)  


**IRIS PROCESSES (PECTINATE LIGAMENTS IN ANTERIOR CHAMBER ANGLE)** 

1. Achondroplasia, diastrophic dwarfism, cartilage-hair hypoplasia, and spondyloepiphyseal dysplasia, anterior chamber cleavage syndrome, Axenfeld syndrome, Reiger syndrome, Peter anomaly.  
2. Congenital glaucoma-may be associated with congenital microcoria and goniodysgenesis  
3. Congenital scoliosis  
4. Legg-Perthes disease (coxa plana)  
5. Marfan syndrome (hypoplastic form of dystrophia mesodermalis congenita)
6. Mucopolysaccharidoses (including Hunter syndrome, Hurler syndrome, Scheie syndrome, and Sanfilippo-Good syndrome)
7. Myopic patients
8. Normal, especially in brown-eyed persons
9. Pigmentary ocular dispersion syndrome


**WIDTE MASS IN ANTERIOR CHAMBER**

* 1. Endophthalmitis
2. Ocular aspergillosis
3. Sterile inflammation following surgery or trauma
* 4. Tumor

11
Pupil

MYDRIASIS (DILATED PUPIL, USUALLY > 5 MM)

1. Physiologic
   A. Larger pupils in women than in men
   B. Larger pupils in myopes than in hypermetropes
   C. Larger pupils in blue irides than in brown irides
   D. Larger pupils in adolescents and middle-aged persons than in very young or old persons
   E. Surprise, fear, pain, strong emotion, or vestibular stimulation
   F. General anesthesia of stages I, II, and IV
   G. Autosensory pupillary reflex-stimulation of middle ear
   H. Auditory pupillary reflex-tuning fork adjacent to ear
   I. Vestibular pupillary reflex-stimulation of labyrinth by heat, cold, or rotation
   J. Vagotonic pupillary reflex-stimulation on deep inspiration

2. Drugs, including the following:
   acetaminophen       allobarbital       anisotropine
   acetenilid          alprazolam         antazoline
   acetophenazine      alseroxylon        antimony lithium
   acetylcholine       amantadine         thiomalate
   adipheneine         ambutonium          antimony potassium
   adrenal cortex injection amitriptyline   tartrate
   albuterol           amobarbital        antimony sodium
   alcohol             amoxapine          antimony sodium
   aldosterone         amphetamine       tartrate
   alkavervir          amyl nitrite       aprobarbital
stibophen  thiopropazate  trihexyphenidyl
syrosingopine  thioproperazine  trimeprazine
talbutal  thioridazine  trimethaphan
temazepam  thiothixene  trimethidinium
tetraethylammonium  tolazoline  trimethaphan
tetrahydrocannabinol  tranylcypromine  tripelennamine
tetrahydrozoline  trazodone  tropicamide
thiamylal  triamcinolone  urethan
thiethylperazine  triazolam  veratrum viride
thiocarbanidin  tridihexethyl  alkaloids
   (THC) (?)  trifluoperazine  vinbarbital
thioglycollate  trifluperidol
thiopental  triflupromazine

3. Toxins, including after-shave lotion, arsenic, *Clostridium botulinum* (gas gangrene), tetanus (lockjaw), cannabis, adrenergic agents (such as nasal sprays or asthma therapy in newborns), paraaminosalicylic acid, lead, carbon monoxide, organic phosphorus, bovine milk protein in infants with allergic malabsorption, *Datura stramonium* (Jimson weed), *Datura wrightii* (moonflower), and *Solanaceae* (nightshade), nitrocompounds and aminocompounds of benzene, carbon disulfide, and papaverine.

4. Ocular causes (mydriasis) (see fixed pupil section p. 348)
   A. Glaucoma, usually acute
   B. Glaucomocylitic crisis (Posner-Schlossman syndrome)
   C. Hollenhorst syndrome (chorioretinal infarction syndrome)
   *D. Iritis; uveitis
   E. Intraocular foreign body (iron mydriasis)
   F. Iris atrophy
   G. Iris sphincter rupture
   *H. Paralytic mydriasis following trauma
   I. Photocoagulation complications
   J. Retinoblastoma

5. Lesions of ciliary ganglion causing internal ophthalmoplegia (e.g., dilated pupil and absent accommodation)
   A. Adie tonic pupil
   B. Congenital lesion
   C. Herpes zoster
   D. Orbital floor fracture repair
   E. Systemic lupus erythematosus (disseminated lupus erythematosus)
   F. Varicella (chickenpox)
   G. Yellow fever

6. Acute or chronic ophthalmoplegias (see p. 168-171)
7. Third-nerve lesion-also ptosis and ophthalmoplegia on affected side (see p. 168-171)
8. Coma because of alcohol ingestion, eclampsia, diabetes, uremia, epilepsy, apoplexy, or meningitis-the pupils are equally dilated and do not constrict with stimulation
9. Midbrain tumors, in which dilated pupils, paralysis of vertical gaze (especially upward gaze), and retraction nystagmus are manifested
A. Craniopharyngioma
B. Parinaud syndrome (paralysis of upgaze movements)

10. Epidural or subdural hematoma
11. Paralytic parasympathetic lesions
12. Irritative sympathetic lesion-pupillary dilatation widening of palpebral aperture and slight exophthalmos
   A. Irritative lesion, such as tumor, encephalitis, or syringomyelia of the hypothalamus, midbrain, medulla, or cervical cord
   B. Thoracic lesions, such as cervical rib, aneurysms of the thoracic vessels, mediastinal tumors, or tubercular pleurisy
   *C. Cervical lesions, including nasopharyngeal tumors, thyroid swelling, or cervical nodes
   D. Rabies (hydrophobia)
   *E. Trauma
   F. Visceral disease
   G. Aortic dilatation or exudative endocarditis (Roque sign)
   H. Acute abdominal conditions, such as appendicitis, cholecystitis, or colitis (Moschowitz sign)
   I. Psychiatric patients with pressure over McBurney point (Meyer phenomenon)

13. Tumors, injury, or hemorrhage of frontoparietal, parietal, temporal, or temporooccipital area-contralateral mydriasis and ipsilateral defect in the visual field
14. Fractured skull
15. Acute autonomic neuropathy
16. Acute pandysautonomia
17. Avitaminosis B\textsubscript{2} (pellagra)
18. Chorea
19. Clivus edge syndrome
20. Cranio cervical syndrome (whiplash injury)
21. Foramen lacerum syndrome (aneurysm of internal carotid artery syndrome)
22. Hemiacrososmia syndrome (hemifacial or unilateral hypertrophy)
23. Iron deficiency anemia
24. Lockjaw (tetanus)
25. Mycosis fungoides syndrome (Sézary syndrome)
26. Optic canal syndrome
27. Parkinson syndrome (shaking palsy)
28. Prematurity
29. Pulseless disease
30. Reye syndrome (acute encephalopathy syndrome)
31. Rollet syndrome (orbital apex-sphenoidal syndrome)
32. Suprarenal-sympathetic syndrome (adrenal medulla tumor syndrome)
33. Temporal arteritis
34. Weber syndrome (cerebellar peduncle syndrome)
35. Wernicke syndrome (I) (avitaminosis B\textsubscript{1} thiamine deficiency)
36. Zellweger syndrome (cerebrohepatorenal syndrome)


**RELATIVE FIXED, DILATED PUPIL**

1. Midbrain damage-vascular accidents, tumors, degenerative and infectious diseases
   A. Dorsal (Edinger-Westphal nucleus and its connections)-rare, involves both pupils, pupillary near reaction often retained, and often associated with supranuclear vertical gaze palsy (upgaze)
   *B. Ventral (fascicular part of third nerve)-associated with other neurologic deficits, such as Nothnagel syndrome, Benedikt syndrome, Weber syndrome, and involves other extraocular components of the third nerve

*2. Damage to the third nerve (from interpeduncular fossa to ciliary ganglion)
   A. Basal aneurysms
   B. Supratentorial space-occupying masses, causing displacement of the brainstem or transtentorial herniation of the uncus; patient is stuporous or comatose
   C. Basal meningitis-often bilateral internal ophthalmoplegia
   *D. Ischemic oculomotor palsy
   E. Parasellar tumor (e.g., pituitary adenoma, meningioma, craniopharyngioma, nasopharyngeal carcinoma, or distant metastases)
   F. Parasellar inflammation (e.g., Tolosa-Hunt syndrome, temporal arteritis, herpes zoster)

*3. Damage to the ciliary ganglion
   A. Viral ciliary ganglionitis or involvement of the ciliary nerves, such as from herpes zoster
   B. Orbital trauma or tumor
C. Trauma from inferior oblique surgery
D. Trauma from retrobulbar injections
4. Damage to short ciliary nerves
   A. Blunt trauma to the globe may injure the ciliary plexus at the iris root (traumatic iridoplegia)
   B. Choroidal trauma or tumor
*5. Damage to the iris
   A. Degenerative or inflammatory diseases of the iris
   B. Posterior synechiae
   C. Acute rise of intraocular pressure (hypoxia or sphincter damage)
   D. Blunt injury to the globe with sphincter damage (traumatic iridoplegia)
   E. Pharmacologic blockade by atropinic substances
   F. Following cataract surgery
6. Total blindness, including cortical blindness (see p. 632-636)
   A. Bilateral optic nerve
      (1) Anterior ischemic optic neuropathy
      (2) Avulsion (traumatic)
      (3) Optic neuritis
   B. Bilateral retina
      (1) Acute retinal necrosis
      (2) Central retinal artery occlusion
      (3) Central retinal vein occlusion
      (4) Retina detachment


**Miosis (Small Pupil) (Usually <2 MM)**

1. Physiologic
   A. Smaller pupil in men than in women
   B. Smaller pupil in hypermetropes than in myopes
   C. Smaller pupil in brown irides than in blue irides
   D. Smaller pupil in very young or old than in adolescents and middle-aged persons
   E. Sleep, fatigue, coma
   F. Stage III anesthesia
   G. Near vision (synkinesis with convergence and accommodation)
   H. Vestibular stimulation
2. Drugs, including the following:

aceclidine  
adropinopon (DFP)  
dronabinol  
dropieridol  
edotocinophate

acetophenazine  
ephedrine (?)  
edetoarginine  
epitocaine

acetylcholine  
ergot  
ephorbidine  
ergocaine

alcohol  
ergotamine  
ether  
ergometrine

allobarbital  
alseroxylon  
ambenonium  
amobarbital

aprobarbital  
ahalperidol  
barbital  
heptabarital

bacoil  
halothane  
bromide  
hexaethalon

bromisovalum  
hexamethylenamine  
bupivacaine  
hexamethylenamine

butabarbital  
hexamethylenamine  
butilbital  
hexamethylenamine

butallylonal  
hexamethylenamine  
butaperazine  
hexamethylenamine

buthanechol  
hexamethylenamine  
bathanechol  
hexamethylenamine

carbromal  
procarbazine  
caricoprolol  
procarbazine

carphenezine  
procarbazine  
chloral hydrate  
procarbazine

chloroform  
procarbazine  
chlorproprazine  
procarbazine

chlorothiazene  
procarbazine  
clonidine  
procarbazine

codeine  
procarbazine  
cyclobarbital  
procarbazine

cycloprobarbital  
procarbazine  
demecarium  
procarbazine

deserpidine  
procarbazine  
diacetilmorphine  
procarbazine

dibucaine  
procarbazine  
diethazine  
procarbazine

digalactides (?)  
procarbazine  
diisopropyl  
procarbazine

fluorophosphate  
procarbazine  
disopropyl  
procarbazine

(DFP)  
procain  
dronabinol  
procain

droperidol  
procain  
edothiophate  
procain

neostigmine  
nioalamide  
nitrous oxide  
opium

oxprenolol  
oxymorphone  
paraldehyde  
perazocine

pentobarbital  
perazine  
pericyazine  
phenazepine

phenacylprazine  
phenol  
phenelzine  
phenobarbital

phenoxybenzamine  
phenylephrine  
physostigmine  
pilocarpine

piperacetafine  
piperazine  
piperacetafine  
pilocarbin

pipericaine  
pilocarbin  
primidone  
probarbital

procaine  
procachrome  
prochlorperazine  
procholine

promazine  
procholine  
promethazine  
propiomazine

propoxycaine  
propranolol  
pyridostigmine  
radioactive iodies (??)

rauwolfa serpentina  
rescinnamine reserpine  
secobarbital  
sulindac

syrosoingopine  
talbutal  
tetracaine  
tetrahydrocannabinol

thiamylal  
thiamylal  
thesethylperazine
thiopental  thiopropazate  thioproperazine  thioridazine  thiothixene
tolazoline  tranzylicpromine  trifluoperazine  trifluperidol  triflupromazine
trimeprazine  vinbarbital  vitamin A

3. Ocular causes
   *A. Accommodative spasm (hysteria)
   B. Corneal irritation, such as keratitis or corneal injury
   C. Conjunctival irritation
   D. Congenital miosis (absent dilator muscle)
   E. Dislocated lenses
   F. Iritis
   *G. Posterior iris synechiae, usually irregular
   H. Retinitis pigmentosa

4. Central nervous system defects
   A. Acute pontine angle lesion, such as hemorrhage or tumor associated with disturbed conjugate gaze
   B. Arteriosclerotic and degenerative disease of the cerebrum
   C. Encephalitis
   D. Facial tetanus
   E. Giant cell (temporal arteritis)
   F. Infections or tumors of the cavernous sinus or superior orbital fissure
   G. Purulent meningitis
   H. Severe hypoxia

5. "Cluster headache" or histamine cephalgia-ptosis; miosis; red, watering eye on side of headache
6. Raeder paratrigeminal syndrome-ipsilateral miosis and pain-may be associated with third-nerve paralysis or corneal anesthesia
   A. Extracranial aneurysm of internal carotid
   B. Idiopathic
   C. Meningioma
   D. Migraine
   E. Posttrauma

7. Argyll Robertson pupil-small and irregular; reacts better to accommodation than to light
   A. Aberrant regeneration of the third nerve
   B. Carbon disulfide poisoning
   C. Cerebral aneurysm
   D. Chronic alcoholism
   *E. Diabetes mellitus (Willis disease)
   F. Encephalitis
   G. Friedreich ataxia
   H. Malaria
   I. Midbrain tumors, such as pinealomas and craniopharyngioma
   J. Multiple sclerosis (disseminated sclerosis)
K. Senile and degenerative diseases of the central nervous system
L. Syphilis (acquired lues)
M. Syringomyelia
N. Trauma to skull or orbit
8. Ataxia, spastic with congenital miosis-dominant
9. Babinski-Nageotte syndrome (medulla tegmental paralysis)
10. Coenurosis
11. Craniocervical syndrome (whiplash injury)
12. Dejerine-Klumpke syndrome (lower radicular syndrome)
13. Devic syndrome (neuromyelitis optica)
*14. Diabetes mellitus
15. Eaton-Lambert syndrome (myasthenic syndrome)
16. Elevated intracranial pressure
*17. Homer syndrome (cervical sympathetic paralysis syndrome)
18. Jugular foramen syndrome (Vernet syndrome)
19. Lowe syndrome (oculocerebrorenal syndrome)
20. Marfan syndrome (arachnodactyly dystrophia mesodermalis congenita)
21. Morquio syndrome (mucopolysaccharidosis IV)
22. Myotonic dystrophy (Curschmann-Stewart syndrome)
23. Naffziger syndrome (scalenum anticus syndrome)
24. Pancoast syndrome (superior pulmonary sulcus syndrome)
25. Parkinsonism (shaking palsy)
26. Psychogenic diseases, such as schizophrenia, dementia precox, or hysteria
27. Refsum syndrome (phytanic acid storage disease)
28. Retroparotid space syndrome (Villaret syndrome)
29. Romberg syndrome (facial hemiatrophy)
30. Spider bites
31. Stormorken syndrome (thrombocytopenia bleeding tendency)
32. Tetanus (lockjaw)
33. von Herrenschwand syndrome (sympathetic heterochromia)
34. Wallenberg syndrome (dorsolateral medullary syndrome)
35. Wernicke syndrome (avitaminosis B1, thiamine deficiency)


PARADOXICAL PUPILLARY REACTION (CONSTRICTS WHEN LIGHT IS WITHDRAWN)

1. Best disease
2. Congenital achromatopsia
3. Congenital stationary night blindness
4. Leber congenital amaurosis
5. Optic nerve hypoplasia
6. Retinitis pigmentosa


**ABSENCE OR DECREASE OF PUPILLARY REACTION TO LIGHT**

This type of absence or decreased pupillary reaction to light is caused by drugs, including the following:

- acetaminophen
- acetanilid
- acetophenazine
- alcohol
- allobarbital
- alprazolam
- amitriptyline
- amobarbital
- amoxapine
- amoxicillin
- amphetamine
- ampicillin
- antazoline
- antimony lithium thiomalate
- antimony potassium tartrate
- antimony sodium tartrate
- antimony sodium thioglucollate
- aprobarbital
- aspirin
- atropine
- baclofen
- barbital
- belladonna
- benztropine
- biperiden
- bromide
- bromisovalum
- brompheniramine
- butabarbital
- butalbital
- butallylional
- butaperazine
- butethal
- calcifediol
- calcitriol
- carbamicillin (?)
- carbinoxamine
- carbon dioxide
- carbromal
- carisoprodol
- carmustine
- carphenezine
- chloramphenicol
- chlorcyclizine
- chlordiazepoxide
- chlorprocaine
- chlorpheniramine
- chlorphenoxamine
- chlorpromazine
- chlorprothixene
- cholecalciferol
- cimetidine
- clemastine
- clomipramine
- clonazepam
- clonidine
- clorazepate
- cloxacillin (?)
- cocaine
- cyclazine
- cyclobarbital
- cyclopentobarbital
- cymicine
- desipramine
- dextroamphetamine
- dextroamphetamine
- diacetylmorphine
- diazepam
- dicloxacillin (?)
- diethazine
- dimethindene
- diphenhydramine
- diphenylpyraline
- diphtheria toxoid, adsorbed
- doxepin
- doxylamine
- emetine
- ergocalciferol
- ergot
- ethopropazine
- etidocaine
- fenfluramine
- fluphenazine
- flurazepam
- flurazepam
ANISOCORIA (INEQUALITY OF PUPILS OF ≥1 MM)

1. Central nervous system
   *A. Adie (tonic) pupil
B. Aneurysm of the aorta or carotid artery  
C. Cerebrovascular accidents  
D. Cervical rib (ipsilateral constricted pupil)  
E. Encephalitis (mild cases)  
*F. Homer syndrome (cervical sympathetic paralysis syndrome)  
G. Pontine lesions  
H. Tabes dorsalis  
I. Third-nerve paresis  
J. Trigeminal neuralgia (tic douloureux)  
K. Wernicke hemianopic pupil

2. Drugs, including the following:

<table>
<thead>
<tr>
<th>Drug</th>
<th>Drug</th>
<th>Drug</th>
</tr>
</thead>
<tbody>
<tr>
<td>alcohol</td>
<td>diphenylpyraline</td>
<td>oral contraceptives</td>
</tr>
<tr>
<td>antazoline</td>
<td>disulfiram</td>
<td>phenelzine</td>
</tr>
<tr>
<td>bromide</td>
<td>doxylamine</td>
<td>pheniramine</td>
</tr>
<tr>
<td>bromisovalum</td>
<td>dronabinol</td>
<td>phenylpropanolamine</td>
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<tr>
<td>brompheniramine</td>
<td>ethchlorvynol</td>
<td>prilocaine</td>
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<tr>
<td>bupivacaine</td>
<td>etidocaine</td>
<td>propanil</td>
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<tr>
<td>carbinoxamine</td>
<td>hashish</td>
<td>propoxycaine</td>
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<tr>
<td>carbromal</td>
<td>isocarboxazid</td>
<td>psilocybin</td>
</tr>
<tr>
<td>chloroprocaine</td>
<td>jimsonweed</td>
<td>pyrilamine</td>
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<tr>
<td>chlorpheniramine</td>
<td>lidocaine</td>
<td>scopolamine</td>
</tr>
<tr>
<td>clemastine</td>
<td>LSD</td>
<td>tetrahydrocannabinol</td>
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<tr>
<td>contraceptives</td>
<td>lysergide</td>
<td>tranylcypromine</td>
</tr>
<tr>
<td>dextrompheniramine</td>
<td>marijuana</td>
<td>trichloroethylene</td>
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<tr>
<td>dextrophiramine</td>
<td>mepivacaine</td>
<td>tripelennamine</td>
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<tr>
<td>diacetylmorphine</td>
<td>mescaline</td>
<td>triprolidine</td>
</tr>
<tr>
<td>dimethindene</td>
<td>methaqualone</td>
<td></td>
</tr>
<tr>
<td>diphenhydramine</td>
<td>nialamid</td>
<td></td>
</tr>
</tbody>
</table>

3. Ocular conditions

A. Artificial eye (pseudoanisocoria)  
B. Cornea, such as keratitis or abrasion  
C. Glaucoma, including pigmentary dispersion  
*D. Iris, such as iritis, synechiae, iris atrophy, or iris sphincter rupture  
E. Ocular trauma  
F. Spastic miosis

4. Physiologic

A. Anisometropia-larger pupil with the more myopic eye  
B. Familial  
C. Lateral illumination of one eye gives more miosis in that eye than in the other  
D. Nonfamilial-normal variation (small percentage of the population)  
E. Tournay reaction—with the eyes turned sharply to the side, dilatation of the pupil of the abducting eye and miosis of pupil of the adducting eye

5. Unilateral miosis (see p. 349)  
6. Unilateral mydriasis (see p. 349)


**IRREGULARITY OF PUPIL (INCLUDING OVAL OR PEAKED PUPIL)**

1. Adherent leukoma as one part of iris is pulled up to corneal scar, peripheral anterior synechiae, or corneal laceration with prolapse of iris
2. Alagille syndrome
3. Anterior chamber intraocular lens that is too long or erodes into uveal tissue
4. Argyll-Robertson pupil-small and irregular; reacts better to accommodation than to light; same type as seen in diabetic patients (pseudodiabetic pupil)
5. Congenital coloboma of the iris, usually below
6. Following laser iridectomy
7. Glaucoma-oval, dilated pupil
8. Injury of the iris
9. Iris tuck of anterior chamber intraocular lens
10. Iritis-usually small but pupil may be any shape with anterior or posterior synechiae
11. Long-term intraocular inflammation
12. Medication, with faster reaction of one sector of iris than of another-miosis or mydriasis
13. Operation-as sector iridectomy or peripheral iridectomy
14. Optic atrophy due to causes such as syphilis, quinine poisoning, and internal ophthalmoplegia of vascular or traumatic origin
15. Piece of anterior capsule into anterior chamber
16. Posterior chamber intraocular lens with loop of intraocular lens holding the midportion of iris peripherally
17. Posterior chamber lens with two haptics having the lens either behind the pupil with the haptics in front or having the lens anterior to the pupil with the haptics behind the iris
18. Segmental iris atrophy
19. Tumors of iris or ciliary body
20. Vitreous or zonules into corneal laceration
21. Vitreous strand from behind pupil to wound
22. Wound leak with or without prolapse of the iris

Hippus is visible, rhythmic, but irregular pupillary oscillations that are deliberate in time. It comprises 2 mm or more excursions and has no localizing significance.

1. Normal
2. Incipient cataracts
3. Central nervous system diseases, including the presence of total third cranial nerve palsy, hemiplegia, meningitis (acute), cerebral syphilis, tabes, general paralysis, myasthenia gravis, tumors of corpora quadrigemina, epileptics, Cheyne-Stokes breathing, multiple sclerosis (disseminated sclerosis), and cerebral tumors
4. Neurasthenia (nervous exhaustion, Beard disease)
5. Drugs, including the following:
   - allobarbital
   - amobarbital
   - aprobarbital
   - barbital
   - butabarbital
   - butalbital
   - butallylonal
   - butethal
   - cyclobarbital
   - cyclopentobarbital
   - heptabarbital
   - hexobarbital
   - mephobarbital
   - metharbital
   - methitural
   - methohexital
   - pentobarbital
   - pentylenetetrazol
   - phenobarbital
   - primidone
   - secobarbital
   - talbutal
   - thiamylal
   - thiopental


**TONOHAPTIC PUPIL**

Tonohaptic pupil involves a long latent period preceding both contraction to light and redilatation, followed in each instance by a short but prompt movement.

1. Catatonic state
2. Diabetes mellitus (Willis disease)
3. Diabetes insipidus
4. Dystrophia adiposogenitalis (Fröhlich syndrome) or pituitary cachexia (Simmonds disease)
5. Introverted persons of the schizophrenic group
6. Parkinsonism (shaking palsy)
7. Pigmentary retinal dystrophy
8. Postencephalitic condition
9. Schizoid state


**LEUKOKORIA (WHITE PUPIL) (SEE LESIONS CONFUSED WITH RETINOBLASTOMA, P. 502)**

1. Angiomatosis of retina (cerebelloretinal hemangioblastomatosis)
2. Astrocytoma
3. Cataract (congenital)
4. Choroidal hemangioma
5. Coats disease (retinal telangiectasia)
6. Coloboma of choroid and optic disc
7. Congenital cytomegalovirus retinitis
8. Congenital retinal detachment
9. Exudative retinitis, chorioretinitis, or both
10. Falciform fold of retina
11. Familial exudative vitreoretinopathy
12. Herpes simplex retinitis
13. High myopia with advanced chorioretinal degeneration
14. Medullation of nerve fiber layer
15. Metastatic endophthalmitis
16. Morning glory syndrome (hereditary central glial anomaly of the optic disc)
17. Nematode endophthalmitis (*Toxocara canis*)
18. Norrie disease (atrophia oculi congenita)
19. Ocular toxocariasis
20. Organized vitreous hemorrhage
21. Persistent hyperplastic primary vitreous
22. Physiologic-eye photographed at 17 to 20 degrees temporal to fixation
23. Retinal dysplasia (massive retinal fibrosis)
24. Retinoblastoma
25. Retinopathy of prematurity (ROP)
26. Retrolental membrane associated with Bloch-Sulzberger syndrome (incontinentia pigmenti)
27. Toxoplasmosis (congenital)
28. Traumatic chorioretinitis
29. Tumors other than retinoblastoma
   A. Choroidal hemangioma
   B. Combined retinal hamartoma
   C. Diktyoma
   D. Glioneuroma
   E. Leukemia
   F. Medulloepithelioma
   G. Retinal astrocytic hamartoma
H. Retinal capillary hemangioma
30. Uveitis (peripheral)
31. Vitreous organization following unsuspected penetrating wounds


Extracted Table Leukokoria (white pupil)

<table>
<thead>
<tr>
<th>LONG CILIARY PROCESSES EXTENDING INTO DILATED PUPILLARY SPACE</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Aniridia</td>
</tr>
<tr>
<td>2. Anterior rotation of ciliary processes</td>
</tr>
<tr>
<td>A. After scleral buckling operation</td>
</tr>
<tr>
<td>B. Angle closure</td>
</tr>
<tr>
<td>C. Anterior choroidal separation</td>
</tr>
<tr>
<td>D. Cyst or tumor behind iris</td>
</tr>
<tr>
<td>E. Dislocated lens</td>
</tr>
<tr>
<td>F. From adherence to limbal scar</td>
</tr>
<tr>
<td>G. Plateau iris</td>
</tr>
<tr>
<td>3. Extreme mydriasis</td>
</tr>
<tr>
<td>4. Falciform detachment of the retina</td>
</tr>
<tr>
<td>5. Incontinentia pigmenti (Bloch-Sulzberger syndrome)</td>
</tr>
<tr>
<td>6. Norrie disease (atrophia oculi congenita)</td>
</tr>
<tr>
<td>7. Persistent hyperplastic primary vitreous (PHPV)</td>
</tr>
<tr>
<td>8. Retinal dysplasia of Reese</td>
</tr>
<tr>
<td>9. Retrolental fibroplasia (RLF)</td>
</tr>
<tr>
<td>10. Surgical coloboma</td>
</tr>
<tr>
<td>11. Trisomy 13 (trisomy D)</td>
</tr>
</tbody>
</table>


**PERSISTENT PUPILLARY MEMBRANE**

1. Fetal iritis
2. Hereditary
3. Physiologic
*4. Use of oxygen therapy in nursery for premature infants


**DECENTERED PUPILLARY LIGHT REFLEX**

1. Positive angle kappa-pseudoexotropia
2. Negative angle kappa-pseudoesotropia
*3. Eccentric fixation-deep unilateral amblyopia
*4. Ectopic macula-macular displacement by retinal scarring or strands, such as retrolental fibroplasia
5. Ectopic pupil


**PUPILLARY BLOCK FOLLOWING CATARACT EXTRACTION**

1. Air pupillary block
2. Dense, impermeable anterior hyaloid membrane
3. Free vitreous block
4. Intraocular lens effectively closing off pupil and iridectomies
5. Leaky wound
6. Nonperforating iridectomy
7. Posterior vitreous detachment associated with pooling or retrovitreal aqueous
8. Postoperative iridocyclitis
9. Subchoroidal hemorrhage
10. Swollen lens material behind the iris


**AFFERENT PUPILLARY DEFECT**
The pupil of the eye has diminished vision from disease of the retina or optic nerve and will fail to react directly to light but will constrict consensually when the healthy eye is stimulated.

1. Amblyopia (rare)
2. Branch retinal artery/vein occlusion
3. Central retinal artery/vein occlusion
4. Compressive optic neuropathy
   A. Cavernous hemangiomas
   B. Cystic tumors
      (1) Cholesterol granuloma
      (2) Conjunctival orbital cysts
      (3) Dermoid cysts
      (4) Mucoceles
   C. Inflammatory and infiltrative processes
   D. Optic nerve tumors
      (1) Optic nerve gliomas
      (2) Optic nerve meningiomas
   E. Primary malignancies
   F. Sarcoidosis
   G. Solid orbital tumors
      (1) Hemangiomas
      (2) Meningiomas
      (3) Schwannoma
   H. Thyroid ophthalmopathy
   I. Trauma
5. Diabetic retinopathy (severe)
6. Hyphema
7. Macular degeneration (rarely)
8. Neovascular glaucoma
*9. Optic neuritis
*10. Optic nerve lesion
11. Radiation
12. Reticulum cell sarcoma
13. Retinal detachment
14. Toxoplasma retinochoroiditis
15. Traumatic optic neuropathy and retinopathy
16. Unilateral optic nerve hypoplasia


ANIRIDIA (ABSENCE OF IRIS, PARTIAL OR COMPLETE)

1. AGR triad - sporadic (bilateral or unilateral) aniridia, genitourinary abnormalities, and mental retardation
2. Associated ocular findings
   A. Cataracts
   B. Corneal dystrophy
   C. Ectopia lentis
   D. Glaucoma
   *E. Macular aplasia - autosomal dominant
   F. Microcornea and subluxated lenses
   G. Nystagmus
   H. Optic nerve hypoplasia
   I. Photophobia
   J. Poor foveal reflex
   *K. Strabismus
3. Associated with autosomal-recessive inheritance with fully developed macula
4. Associated with unilateral renal agenesis and psychomotor retardation
5. Beckwith-Wiedemann syndrome
6. Deletion of short arm of 11th chromosome
7. Gillespie syndrome (incomplete aniridia, cerebellar ataxia, and oligophrenia)
8. Homocystinuria syndrome
9. Marinesco-Sjögren syndrome (congenital spinocerebellar ataxia)
10. Miller syndrome (Wilms aniridia syndrome)
11. Partial trisomy 2q
12. Peters syndrome (oculodental syndrome)
13. Rieger syndrome (dysgenesis mesostromalis)
14. Ring chromosome 6
15. Scaphocephaly syndrome
16. Siemens syndrome (anhidrotic ectodermal dysplasia)
17. Traumatic
18. Ullrich syndrome (dyscraniopylophalangy)


**COLOBOMA OF IRIS**

This condition involves failure of fusions of fetal fissure in optic vesicle, usually inferior or inferonasal.

1. Acrorenooclar syndrome
2. Aicardi syndrome
3. Aniridia
4. Biemond syndrome
5. Cat eye syndrome (partial G-trisomy syndrome)
6. CHARGE association (coloboma, heart anomaly, choanal atresia, retardation, genital, and ear anomalies)
7. Chromosome partial short-arm deletion syndrome
8. Ellis-van Creveld syndrome (chondroectodermal dysplasia)
9. Epidermal nevus syndrome (ichthyosis hystrix)
10. Focal dermal hypoplasia syndrome (Goltz syndrome)
11. Hallermann-Streiff-François syndrome (dyscephalic mandibulooculofacial syndrome)
12. Hemifacial microsomia syndrome (otomandibular dysostosis)
*13. Hereditary usually dominant may be recessive
14. Hurler syndrome (mucopolysaccharidoses I)
15. Hyperchromic heterochromia
16. Jeune disease (asphyxiating thoracic dystrophy)
17. Joubert syndrome
18. Kartagener syndrome
19. Klinefelter syndrome
20. Klippel-Trenaunay-Weber syndrome (angioostehypertrophy syndrome)
1. Proximal vascular disease
   A. Aortic arch syndrome (pulseless disease; Takayasu syndrome)
B. Carotid-cavernous fistula (carotid artery syndrome)
C. Carotid ligation
D. Carotid occlusive disease
E. Cranial arteritis syndrome (giant cell arteritis)

2. Ocular vascular disease
   *A. Central retinal artery thrombosis (see p. 457)
   *B. Central retinal vein thrombosis (see p. 467)
   C. Long posterior ciliary artery occlusion
   D. Reversed flow through the ophthalmic artery

3. Retinal diseases
   A. Coats disease (retinal telangiectasia)
   *B. Diabetes mellitus
   C. Eales disease (periphlebitis)
   D. Glaucoma, chronic
   E. Melanoma of choroid
   F. Norrie disease (oligophrenia-microphthalmos syndrome)
   G. Persistent hyperplastic primary vitreous
   H. Retinal detachment
   I. Retinal hemangioma
   J. Retinoblastoma
   K. Retrolental fibroplasia
   L. Sickle cell disease (Herrick syndrome)

4. Iris tumors
   A. Hemangioma
   B. Melanoma
   C. Metastatic carcinoma

5. Postinflammatory
   A. Argon laser coreoplasty
   B. Exfoliation syndrome
   C. Fibrinoid syndrome
   D. Fungal endophthalmitis (see p. 223-225)
   E. Iris neovascularization with pseudoexfoliation
   F. Radiation
   G. Surgery for retinal detachment
   H. Uveitis, chronic

6. Vascular tufts at the pupillary margin
   A. Cataract
   B. Diabetes mellitus
   C. Myotonic dystrophy syndrome (myotonia atrophica syndrome)
   D. Ocular hypotony
   E. Respiratory failure


**HYPEREMIA OF IRIS (DILATATION OF PREEXISTING VESSELS OF THE IRIS)**

1. Corneal ulcer
2. Foreign body on the cornea
3. Injury, intraocular
   *4. Iridocyclitis
   *5. Iritis
6. Scleritis
*7. Uveitis


**HETEROCHROMIA (DIFFERENCE OF COLOR BETWEEN TWO IRIDES)**

1. Hypochromic heterochromia—abnormal eye with iris of lighter color than that of the fellow eye
   A. Anemia with unilateral iritis
   B. Chédiak-Higashi syndrome (anomalous leukocytic inclusions with constitutional stigmata)
   *C. Congenital, sporadic, or familial
   D. Conradi syndrome (epiphyseal congenital dysplasia)
   E. Fuchs syndrome (I) (heterochromic cyclitis syndrome)
   F. Gansslen syndrome (familial hemolytic icterus)
   G. Glaucomatocyclitic crisis (Posner-Schlossman syndrome)
   *H. Homer syndrome (cervical sympathetic paralysis syndrome)
   I. Hypomelanosis of Ito syndrome (incontinentia pigmenti achromiens)
   J. Infiltration of nonpigmented tumor into iris
   *K. Iris atrophy (diffuse and unilateral), including that caused by trauma, inflammation, or senility
   L. Iris coloboma
   M. Parry-Romberg syndrome (facial hemiatrophy)
   N. Status dysraphicus syndrome (Bremer syndrome)
   O. Tuberosous sclerosis hypopigmented iris spot (Bourneville syndrome)
   P. Waardenburg-Klein syndrome (embryonic fixation syndrome)
2. Hyperchromic heterochromia—abnormal eye with iris darker than that in the fellow eye
   A. Anterior or posterior chamber hemorrhage, prolonged
   B. Coloboma
   *C. Congenital, sporadic, or familial
   D. Embryonic fixation syndrome (Waardenburg-Klein syndrome)
   E. Incontinentia pigmenti (Bloch-Sulzberger syndrome)
   F. Iris abscess
   G. Iris stromal cysts
   H. Malignant melanoma of the iris or other pigmented tumors of the iris
I. Microcornea (see p. 252-253)
J. Monocular melanosis in which there are excess chromatophores in the stroma of the iris (melanosis bulbi)
*K. Neovascular, such as rubeosis iridis or hyperemia of iris, unilateral (see p. 367 or p. 366)
L. Neurofibromatosis (von Recklinghausen syndrome)
*M. Nevi of iris
N. Perforating injuries or contusion of the globe occurring before the subject is seven years of age
O. Retention of intraocular iron foreign body-siderosis
P. Severe contusion with hypertrophy of the superficial layers of the stroma of the iris
Q. Status dysraphicus (Bremer syndrome)

3. Dark central pupillary margin, pale pigment around its circumference
   A. Hereditary osteoonychodysplasia
   B. Normal iris


Extracted Table

| Hyperchromic heterochromia (abnormal eye with iris of darker color than fellow eye) |

IRIS ATROPHY

1. Anterior segment ischemia syndrome
2. Arteriovenous fistula
3. Chandler syndrome (iridocorneal endothelial syndrome)
4. Complication of light coagulation and beta radiation
5. Complication of retinal detachment operation
*6. Congenital-autosomal dominant
7. Crohn disease (granulomatous ileocolitis)
*8. Essential (progressive) atrophy
9. Glaucomatous atrophy
   A. Acute-atrophy or iridoschisis
   *B. Chronic-stromal and epithelial
10. Hallermann-Streiff-François syndrome (dyscephalic mandibulooculofacial syndrome)
11. Hilding syndrome (destructive iridocyclitis and multiple joint dislocations)
12. Homocystinuria syndrome
13. Hypothermal injury
14. Iris nevus syndrome (Cogan-Reese syndrome)
15. Ischemia
   A. Acute angle-closure glaucoma
   B. Carotid-cavernous fistula
   C. Hemoglobin sickle cell C disease
   D. Occlusive artery disease
   E. Orbital irritation
   F. Surgery angle-closure glaucoma
   *G. Trauma
16. Krause syndrome (congenital encephalo-ophthalmic dysplasia)
17. Neurogenic-tabes with stromal atrophy
18. Norrie disease (fetal iritis syndrome)
*19. Old age
20. Pierre Robin syndrome (micrognathia-glossoptosis syndrome)
21. Posterior pigment layer is swollen and degenerated
   A. Diabetes mellitus (Willis disease)
   B. Hurler syndrome (mucopolysaccharidoses I-H)
22. Postinflammatory-iritis because of diseases such as tuberculosis, syphilis (acquired lues), herpes zoster, herpes simplex, smallpox, leprosy (Hansen disease), onchocerciasis syndrome (river blindness), sporotrichosis
23. Shy-Magee-Drager syndrome (orthostatic hypotension syndrome)
24. Spontaneous progressive
   A. Congenital hypoplasia iris stroma
   B. Rieger syndrome (dysgenesis mesostromalis)
25. Takayasu syndrome (aortic arch syndrome, pulseless disease)
26. Use of quinine, chloramine, mustard gas
27. Wagner syndrome (hyaloideoretinal degeneration)
28. Xeroderma pigmentosa, including skin lesions


IRIDODONESIS (TREMULOUS IRIS)

*1. Aphakia following cataract extraction
*2. Dislocation of the lens (see p. 405)
3. Hydrophthalmos or buphthalmos (see p. 222-223)
4. Hypermature senile cataract


TUMORS ARISING FROM PIGMENT EPITHELIUM OF IRIS

1. Hyperplasia
   A. Primary (congenital)
      (1) At pupillary margin
      (2) At margins of colobomas
   B. Acquired
      (1) Region of sphincter-migrating epithelial cells appear in stroma as clump cells (equivocal origin)
      (2) Cells can reach anterior surface of iris and proliferate (velvety black in appearance)
   C. Secondary
      (1) Intraocular inflammation-pigmented cells proliferate around the papillary margin onto anterior iris surface
      (2) Long-standing glaucoma
         a. Proliferation around the pupillary margin onto the anterior iris surface
         b. Migration through stroma to anterior surface at collarette
      *(3) Trauma (including operation-proliferation of pigment epithelium on anterior surface of iris, across pupil, or on posterior surface of cornea
      *(4) Drugs, including the following: demecarium, echothiophate, edrophonium, isofluorophate, neostigmine, physostigmine, pilocarpine often associated with cystic formation

2. Neoplasia
   *A. Benign - well-differentiated epithelial cells, usually pigmented, often with pseudoacinar arrangement and cysts; may have limited locally invasive properties
   B. Malignant
      (1) Carcinoma
      (2) Local invasion, intraocular metastases
      (3) Medulloepithelioma, embryonal type (diktyoma)
Papillary cystadenoma


**PIGMENTED LESIONS OF IRIS**

1. Adenoma of iris
2. Anterior chamber intraocular lens and segmental uveal ectropion
3. Anterior staphyloma
4. Corneal or scleral perforation

*5. Cyst-congenital, spontaneous, or traumatic, including pigmentation
6. Ectopic lacrimal gland tissue

*7. Ectropion uvea
8. Epithelioma of the ciliary body
9. Exudative mass in the anterior chamber
10. Foreign body of iris, including iron with siderosis
11. Fuchs syndrome of heterochromic cyclitis with the darker normal iris considered to contain a diffuse melanoma
12. Hemangioma of the iris with pigmentation because of hemorrhage
13. Hemosiderosis because of contusions with hyphema or injuries and disease in the posterior portion of the eye with recurrent bleeding
14. Juvenile xanthogranuloma (nevoxanthoendothelioma)
15. Leiomyoma or leiomyosarcoma of the iris
16. Leukemic infiltrates and malignant lymphomas
17. Malignant melanoma of the iris
18. Metastatic carcinomas arising in the lung, breast, gastrointestinal tract, thyroid gland, prostate gland, kidney, or testicle
19. Neurofibromatosis with increased pigmentation of the iris

*20. Nevi of the iris
21. Nodular thickening and scarring of the iris
22. Pigmentary glaucoma
23. Pigment epithelial tumors of the iris
24. Segmental melanosis oculi, including congenital melanosis
25. Stromal mass in the anterior chamber
26. Uveitis, such as that because of conglomerate tuberculous lesions of the stroma or sarcoid involvement of the iris
27. Varix


**NONPIGMENTED LESIONS OF IRIS**

1. Amelanotic melanoma
2. Atypical mycobacterial panophthalmitis
3. Endothelioma
4. Exudative mass in the anterior chamber
5. Fibrosarcoma
*6. Foreign body
7. Forward extension of diktyoma
8. Hemangioma of the iris
*9. Iris cyst
10. Iris lymphoma
11. Iris nodules
   A. Ectodermal (Koeppe nodules)-pupillary margin and gray with ocular inflammation
   B. Mesodermal (Busacca nodules)-anterior surface of iris in collarette region
12. Juvenile xanthogranuloma-may be associated with diffuse infiltration of the iris
13. Lacrimal Gland Choristoma of the Iris
14. Leiomyoma or leiomyosarcoma of the iris
15. Leprosy (Hansen disease)
   A. Lepromas of the iris
   B. Leprotic pearl-minute white spots on the surface of iris
16. Metastatic carcinoma of the iris arising from the lungs, breast, gastrointestinal tract, thyroid gland, prostate gland, kidney, or testicle
17. Neurofibroma and neuroglioma
18. Sarcoid nodules-multiple, discrete, irregularly distributed over the iris
19. Seeding of tumor, such as retinoblastoma, from the posterior segment
20. Syphilis (acquired lues)
   A. Gummas-solitary, large, avascular, white lesions
   B. Papules (condylomas)-multiple, small, vascular, yellowish lesions
21. Teratoma
22. Tuberculosis
   A. Acute miliary - small grayish yellow or reddish nodules
   B. Hyalinized or fibrotic scar (Michel flecks)
   C. Tuberculoma-white-gray lesion


**CONDITIONS SIMULATING ANTERIOR UVEITIS OR IRITIS**

1. Brushfield spots
2. Fuchs syndrome (II) (Stevens-Johnson syndrome)
3. Hereditary deep dystrophy of cornea
4. Hyalinized keratitic precipitate
5. Iridoschisis-splitting of iris
6. Juvenile xanthogranuloma of the iris (nevoxanthoendothelioma)
7. Malignant lymphomas or leukemia
8. Malignant melanoma
9. Metastatic tumor arising from the lungs, breast, gastrointestinal tract, thyroid gland, prostate gland, kidney, or testicle
10. Neurofibromas of the iris
11. Pigment floaters in the anterior chamber, especially after mydriasis
12. Pseudoexfoliation of the lens capsule (glaucoma capsulare)
13. Reticulum cell sarcoma
14. Retinoblastoma
15. Scleroderma (progressive systemic sclerosis)
16. Siderosis bulbi


**SYNDROMES AND DISEASES ASSOCIATED WITH IRITIS**

1. Actinomycosis
2. Amebiasis (entamoeba histolytica)
3. Amendola syndrome (Brasilian pemphigus)
4. Anderson-Warburg syndrome (congenital progressive oculoacousticocerebral dysplasia)
5. Ankylosing spondylitis (von Beckterev-Strumpell syndrome)
6. Ascariasis
7. Aspergillosis
8. Beesting of the cornea
9. Behçet syndrome (oculobuccogenital syndrome)
10. Blastomycosis
11. Brucellosis (Bang disease)
12. Candidiasis
13. Charlin syndrome (nasociliary nerve syndrome)
14. *Chlamydia pneumoniae
15. Coccidioidomycosis
16. Cryptococcosis
17. Cysticercosis
18. Cytomegalic inclusion disease (cytomegalovirus)
19. Dengue fever
20. Endophthalmitis phacoanaphylactica
*21. Following laser iridectomy
22. Fuchs syndrome (heterochromic cyclitis syndrome)
23. Henoch-Schönlein purpura (anaphylactoid purpura)
24. Herbicide exposure-2, 4-dichlorophenoxyacetic acid
*25. Herpes simplex
*26. Herpes zoster
27. Histoplasmosis
28. Histiocytosis X (xanthomatous granuloma syndrome)
29. Hypervitaminosis D
30. Leptospirosis (Weill disease)
31. Mucormycosis (phycomycosis)
32. Mustard gas injury
33. Mycoplasma pneumoniae
34. Nocardiosis
35. Onchocerciasis syndrome (*Onchocerca volvulus)
36. Reiter syndrome (conjunctivourethrosynovial syndrome)
37. Romberg syndrome (facial hemiatrophy)
38. Rubella syndrome (Gregg syndrome)
*39. Sarcoidosis syndrome (Schaumann syndrome)
40. Sporotrichosis
*41. Still disease (juvenile rheumatoid arthritis)
42. Syphilis (acquired lues)
43. Toxoplasmosis (ocular toxoplasmosis)
44. Tuberculosis
45. Vaccinia
46. Vogt-Koyanagi-Harada syndrome (uveitis, vitiligo-alopecia-poliosis syndrome)
IRITIS (ANTERIOR UVEITIS) IN CHILDREN

1. Anterior and posterior uveitis
   A. Retinoblastoma
      *B. Sarcoidosis syndrome (Schaumann syndrome)
   C. Sympathetic ophthalmia
   D. Vogt-Koyanagi-Harada syndrome (uveitis-vitiligo-alopecia-poliosis syndrome)
*2. Chronic cyclitis (peripheral uveitis)
3. Fuchs heterochromic cyclitis
4. Iridocyclitis
   A. Acute tubulointerstitial nephritis and uveitis
   B. Ankylosing spondylitis (von Beckterev-Strumpell syndrome)
   C. Behçet syndrome (oculobuccogenital syndrome)
   D. Juvenile xanthogranuloma
   E. Kawasaki disease
   F. Leukemia
   G. Multiple sclerosis
   H. Psoriatic arthropathy
   I. Reiter syndrome
   J. Retinal capillaritis
   *K. Sarcoidosis syndrome (Schaumann syndrome)
   L. Still syndrome (juvenile rheumatoid arthritis)
   *M. Trauma
   N. Ulcerative colitis and Crohn disease
   O. Unknown
   P. Viral-associated disease
5. Keratouveitis
   *A. Herpes simplex
   *B. Herpes zoster


**Extracted Table Iritis (anterior uveitis) in children**

**NONGRANULOMATOUS UVEITIS**

1. Physical insult  
   A. Endogenous  
   B. Exogenous  

*2. Toxic insults*  
   A. Autointoxication-pteroinines, protein split products, and so forth from food poisoning  
   B. Bacterial endotoxins  
   C. Reticulum cell sarcoma of the brain  
   D. Toxins from disintegrating helminths  
   *E. Viral toxins*

3. Immediate hypersensitive reaction  
   *A. Airborne allergens*  
   B. Drugs, including the following:

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C. Foods
D. Protein antigens (anaphylaxis)

4. Delayed hypersensitive reaction
   A. Bacterial antigens
   B. Viral antigens

5. Doubtful entities-nongranulomatous uveitis
   A. Amebiasis
   B. Diabetic iritis
   C. Gouty iritis
   D. Heterochromic iridocyclitis
   *E. Sarcoidosis syndrome (Schaumann syndrome)
   F. Secondary to metabolic disease, such as biliary cirrhosis and systemic xanthomatosis
   G. Uveitis associated with collagen diseases

6. Mixed granulomatous and nongranulomatous
   A. Lens-induced uveitis
   *B. Peripheral uveitis

7. Human leukocyte antigen (HLA)-B27 associated diseases
   A. Ankylosing spondylitis
   B. Inflammatory bowel disease
   C. Psoriasis
   D. Reiter disease

8. Infections
   A. Herpes zoster/herpes simplex
   B. Lyme disease
   C. Syphilis


**GRANULOMATOUS UVEITIS**

1. Proven or probable etiology
   A. Associated with nonpyogenic systemic infections
      (1) Brucellosis (*Brucella melitensis, B. abortus, B. suis*)
      (2) Leprosy (*Mycobacterium leprae*)
      (3) Leptospirosis (*Leptospira canicola, L. icterohaemorrhagiae, L. pomona*)
      *(4) Syphilis (*Treponema pallidum*)
      *(5) Tuberculosis (*Mycobacterium tuberculosis*)
   B. Protozoan infections
      (1) Amebiasis (*Entamoeba coli, E. histolytica, Endolimax nana, Acanthamoeba hartmannella*)
      (2) Toxoplasmosis (*Toxoplasma gondii*)
      (3) Trypanosomiasis (*Trypanosoma cruzi, T. gambiense*)
   C. Fungal infections
      (1) Actinomycosis
      (2) Aspergillosis
      (3) Blastomycosis
      (4) Candidiasis (moniliasis)
      (5) Coccidioidomycosis
      (6) Cryptococcosis (*Cryptococcus neoformans* or *Torula histolytica*)
      *(7) Histoplasmosis (*Histoplasma capsulatum*)
      (8) Mycomycosis (phycomycosis)
      (9) Nocardiosis
      (10) Sporotrichosis (*Sporotrichum schenckii*)
   D. **Helminth** infestations
      (1) Ascaridiosis (*Ascaris lumbricoides*)
      (2) Cestodes
         a. Cysticercosis (*Cysticercus cellulosae*)
         b. Taeniasis (*Taenia echinococcus*)
      (3) Diptera larvae (exogenous)
      (4) Nematodes
         a. Ancylostomiasis (*Toxocara canis, Ancylostoma duodenale, Ancylostoma caninum, Necator americanus*)
         b. Onchocerciasis (*Onchocerca volvulus*)
E. Viral-herpes zoster

2. Recognized clinical and histopathologic entity, of unknown cause
   *A. Multiple sclerosis
   *B. Sarcoidosis syndrome (Schaumann syndrome)
   C. Sympathetic ophthalmia

3. Nonspecific granulomatous uveitis of unknown cause, including granulomatous ileocolitis

4. Mixed granulomatous and nongranulomatous
   A. Lens-induced uveitis
   B. Peripheral uveitis

5. Viral uveitis
   A. Proven or probable
      (1) Cytomegalic inclusion disease
      (2) Herpes simplex
      (3) Herpes zoster
      (4) Vaccinia
   B. Suspected
      (1) Behçet syndrome (oculobuccogenital syndrome)
      (2) Retinal capillaritis
      (3) Vogt-Koyanagi-Harada syndrome (uveitis-vitiligo-alopecia-poliosis syndrome)

6. Histiocytosis X (includes eosinophilic granuloma, Hand-Schüller-Christian disease and Letterer-Siwe disease)

7. Following treatment of a choroidal melanoma with proton-beam irradiation


**PIGMENTED CILIARY BODY LESIONS**

* 1. Ciliary body cyst

2. Diffuse iris melanotic lesion

*3. Drugs including the following:

<table>
<thead>
<tr>
<th>Drug</th>
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<tr>
<td>adrenal cortex injection</td>
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</tr>
<tr>
<td>echothiophate</td>
<td>methylprednisolone</td>
<td></td>
</tr>
</tbody>
</table>
4. Malignant melanoma
5. Melanocytoma of ciliary body
6. Peripheral uveal detachment
7. Posttraumatic pigmentary migration


**NEUROEPITHELIAL TUMORS OF CILIARY BODY**

1. Congenital
   A. Glioneuroma
   B. Medulloepithelioma
      *(1) Benign
      (2) Malignant
   C. Teratoid medulloepithelioma
      (1) Benign
      (2) Malignant

2. Acquired
   A. Adenocarcinoma
      *(1) Papillary
      (2) Pleomorphic
      (3) Solid
   B. Adenoma
      *(1) Papillary
      (2) Pleomorphic
      (3) Solid
   C. Mesectodermal leiomyoma
   D. Pseudoadenomatous hyperplasia


**INTERNAL OPHTHALMOPLEGIA**
Internal ophthalmoplegia is characterized by paresis of ciliary body with loss of power of accommodation and pupil dilatation because of lesions of ciliary ganglion.

1. Acute porphyria-frequently bilateral
*2. Adie syndrome (myotonic pupil)
3. Aneurysm of the posterior communicating artery at its junction with the internal carotid-unilateral
4. Congenital-rare
*5. Cycloplegic ocular medication-most common
6. During acute illness-transient
7. During blepharoplasty-transient
8. Fisher syndrome (ophthalmoplegia-ataxia-areflexia syndrome)
9. Foramen lacerum syndrome (aneurysm of internal carotid artery)
10. Histiocytosis X (Hand-Schüller-Christian syndrome)
11. Hollenhorst syndrome (chorioretinal infarction syndrome)
*12. Increased intracranial pressure
13. Infections, including chickenpox, measles, diphtheria, syphilis, scarlet fever, pertussis, smallpox, influenza, herpes zoster, botulism, sinusitis, and viral hepatitis
14. Lubarsch-Pick syndrome (amyloidosis)
15. May be early lesion of acute or chronic ophthalmoplegia
16. Metastatic tumors of choroid
17. Nasopharyngeal carcinoma-early
18. Nothnagel syndrome (ophthalmoplegia-cerebellar ataxia syndrome)
19. Partial seizures
20. Retrobulbar injections of alcohol
21. Transscleral diathermy
22. Trauma to eye or orbit


ANTERIOR SUBCAPSULAR CATARACT

1. Acrodermatitis chronica atrophicans
2. Addison syndrome (adrenal cortical insufficiency)
3. Albinism
4. Allopurinol therapy
5. Alport syndrome (hereditary nephritis)
6. Amiodarone usage
7. Andogsky syndrome (dermatogenous cataract)
8. Aniridia
9. Anterior chamber air
10. Atopic (eczema cataract)
11. Beesting of cornea
12. Cerebrohepatorenal syndrome (Smith-Lemli-Opitz syndrome)
13. Chlorpromazine therapy
14. Chromosomal 3; 18 translocation
15. Comedo cataract
16. Coughing
17. Cryotherapy
18. Electric cataract
19. Diabetes mellitus (Willis disease)
20. Facial paralysis (partial)
21. Frenkel syndrome (ocular contusion syndrome)
22. Goldscheider syndrome (epidermolysis bullosa)
23. Gyrate atrophy (ornithine ketoacid aminotransferase deficiency)
24. Head-banging (chronic)
25. Hemifacial microsomia syndrome (Francois-Haustrate syndrome)
26. Hypermature cataract with other changes
27. Hypoparathyroidism
28. Idiopathic-10% of normal population
29. Intraocular copper and iron
30. Isotretinoin
31. Jadassohn-Lewandowsky syndrome (epidermolysis bullosa)
32. Leber congenital amaurosis
33. Marinesco-Sjögren syndrome (oligophrenia syndrome)
34. Myotonic dystrophy (Curschmann-Steinert syndrome)
35. Naphthalene ingestion
37. Neurodermatitis
38. Pemphigus foliaceous (Cazenave disease)
39. Phenothiazine therapy
40. Phospholine iodide use
41. Pseudohypoparathyroidism
42. Reese-Ellsworth syndrome (anterior chamber cleavage syndrome)
43. Rothmund syndrome (telangiectasia-pigmentation-cataract syndrome)
44. Scaphocephaly
45. Schizophrenia
46. Thorazine ingestion
47. Trauma, such as contusion
48. Tyrosinosis
49. Vitrectomy for diabetic retinopathy
50. Werner syndrome (progeria of adults)
51. Wilson disease (hepatolenticular degeneration)
52. Zinc chloride (concentrated)


**Extracted Table: Anterior subcapsular cataract**

**NUCLEAR CATARACTS**

1. Alcohol
2. Arteriovenous fistula
3. Associated with photocoagulation, such as argon laser use
4. Capsular exfoliation syndrome
5. Congenital dysplasia
6. Conradi syndrome (multiple epiphyseal congenital dysplasia)
7. Coppock cataract, discoid cataract, zonular cataract - autosomal dominant
8. Hyperbaric oxygen therapy
9. Maple-syrup urine disease (branched-chain ketoaciduria)
10. Matsouka syndrome (oculocerebroarticuloskeletal syndrome)
11. Micro syndrome
12. Nuclear diffuse nonprogressive cataract - autosomal dominant, rarely recessive
*13. Nuclear sclerosis
   A. Pars plana vitrectomy for macular pucker
   B. Smoking
14. Nuclear total cataract - autosomal dominant, rarely recessive
15. Paradichlorobenzene (mothballs)
16. Perforating injuries
17. Roy syndrome II - nuclear cataract associated with smoking
*18. Rubella syndrome (German measles)
19. Siemen syndrome (hereditary ectodermal dysplasia syndrome)
20. von Gierke disease (glucose-phosphate deficiency)


**LAMELLAR (STELLATE, ZONULAR, CORTICAL, CORONARY) CATARACTS**
1. Alcohol
2. Aniridia
3. Argon laser
4. Autosomal-dominant congenital cataract
5. Congenital zonular cataract
6. Cortical cataract and congenital ichthyosis
7. Dermochonoral corneal dystrophy
8. Diabetes mellitus (Willis disease)
9. Females
10. Galactokinase deficiency (von Reuss syndrome)
11. Hagberg-Santavuori (neuronal ceroid-lipofuscinoses)
12. Hypertension
13. Hypoglycemia
14. Hypophosphatasia (phosphoethanolaminuria)
15. Iritis
16. Leiomyoma
17. Mannosidosis
18. Marfan syndrome (doliostenomelia-arachnodactyly-hyperchondroplasia-dystrophy mesodermalis congenital
19. Marshall syndrome (atypical ectodermal dysplasia)
20. Myotonic dystrophy syndrome (Curschmann-Steinert syndrome)
21. Neurofibromatosis 2 (central neurofibromatosis)
22. Nieden syndrome (telangiectasia cataract syndrome)
23. Nonwhites
24. Obesity (abdominal)
25. Organic nitrate explosives
26. Passow syndrome (status dysraphicus syndrome)
27. Riboflavin deficiency (Stannus cerebellar syndrome)
28. Roy syndrome (cataract associated with smoking)
29. Sunlight
30. Tetany cataract (hypoparathyroidism)
31. Ultraviolet-B light
32. Van Bogaert-Scherer-Epstein syndrome (primary hyperlipidemia)
33. Van der Hoeve syndrome (osteogenesis imperfecta)
34. Wagner syndrome (hyaloideoretinal degeneration)
35. Zonular cataract and nystagmus-X-linked


**PUNCTATE CATARACTS (NUMEROUS SMALL OPACITIES)**

1. Albright hereditary osteodystrophy (pseudohypoparathyroidism)
2. Argon laser
3. Autosomal dominant vitreoretinochoroidopathy (ADVIRC)
4. Cockayne syndrome (dwarfism with retinal atrophy and deafness)
5. Cretinism (hypothyroidism)
6. Galactokinase deficiency (von Reuss syndrome)
7. Hypercalcemia (adult)
8. Hyperprolactinemia
9. Incontinentia pigmenti achromians
10. Lowe syndrome (oculocerebrorenal syndrome)
11. Rothmund syndrome (telangiectasia-pigmentation-cataract syndrome)
12. Supravalvular aortic stenosis (Williams-Beuren syndrome)


**POSTERIOR SUBCAPSULAR CATARACT**

1. Complicated cataract
   A. Anterior segment involvement, such as that because of the following:
      (1) Acute and chronic corneal ulcer
      *(2) Iridocyclitis
      *(3) Chronic anterior uveitis
      (4) Acute or chronic glaucoma
   B. Posterior segment involvement such as that because of the following:
      *(1) Chronic posterior uveal inflammation
      (2) Long-standing retinal detachment
      (3) High myopia
      (4) Hereditary retinal lesions, including retinitis pigmentosa
      (5) Persistent hyperplastic primary vitreous
2. Congenital posterior polar lens changes
   A. Spurious posterior capsular cataract (Mittendorf dot)
   B. Posterior polar cataract-persistent fibrovascular sheath of lens with or without secondary cataract
   C. Posterior lenticonus
3. Aberfeld syndrome (ocular and facial abnormalities syndrome)
4. Acrodermatitis chronic a atrophicans
5. Alcoholism
6. Aniridia
7. Anterior segment ischemia syndrome
8. Aspergillosis
9. Bassen-Kornzweig syndrome (familial hypolipoproteinemia)
10. Bloch-Sulzberger syndrome (incontinentia pigmenti)
11. Buerger disease (thromboangiitis obliterans)
12. Capsular exfoliation syndrome
13. Carotid artery syndrome
14. Chromosome partial deletion (short-arm) syndrome
15. Congenital amaurosis of Leber (Leber congenital amaurosis)
16. Cushing syndrome
17. Diabetes mellitus (Willis disease)
18. Drugs, including dinitrophenol busulfan (Myleran), triparanol (MER-29), PUVA (psoralen plus ultraviolet light of A-wave length), allopurinol, indapamide, megestrol acetate, and phenothiazine usage
19. Electrical injury
20. Engelmann syndrome (diaphyseal dysplasia)
21. Fabry disease (glycosphingolipid lipidosis)
22. Familial hypogonadism syndrome
23. Frenkel syndrome (ocular contusion syndrome)
24. Fuchs syndrome
25. Glassblowers (heat) cataract
26. Gyrate atrophy (ornithine ketoacid aminotransferase deficiency)
27. Hagberg-Santavuori syndrome (neuronal ceroid-lipofuscinoses)
28. Hair dye
29. Hand-Schüller-Christian syndrome (xanthomatous granuloma syndrome)
30. Harada syndrome (uveitis-vitiligo-alopecia-poliosis syndrome)
31. Heerfordt syndrome (uveoparotitis)
32. Hemochromatosis
33. Herpes simplex
34. Hodgkin disease
35. Hypertension
36. Hypoparathyroidism
37. Ionizing radiation, such as that encountered in x-ray, radium, or neutron therapy
38. Jacobsen-Brodwall syndrome
39. Kussmaul disease (necrotizing angiitis)
40. Kyrle disease (hyperkeratosis follicularis et parafollicularis in cutem penetras)
41. Laurence-Moon-Bardet-Biedl syndrome (retinitis pigmentosa-polydactyly-adipsogenital)
42. Leprosy (Hansen disease)
43. Leri syndrome (carpal tunnel syndrome)
44. Leukemia
45. Lightning induced
46. Malaria
47. Meckel syndrome (dysencephalia splanchnocystic syndrome)
48. Myotonic dystrophy (Curschmann-Steinert syndrome)
49. Neurodermatitis (lichen simplex chronicus)
50. Neurofibromatosis 1 (von Recklinghausen syndrome)
51. Neurofibromatosis 2 (central neurofibromatosis)
52. Ocular trauma (blunt)
53. Oculootoororenoerythropoietic disease
54. O'Donnell-Pappas syndrome (foveal hypoplasia and presenile cataract-autosomal dominant)
55. Paget syndrome (osteitis deformans)
56. Pemphigus foliaceus (Cazenave disease)
57. Pernicious anemia syndrome
58. Pierre Robin syndrome (micrognathia-glossoptosis syndrome)
59. Posterior polar cataract-autosomal dominant
60. Pseudohypoparathyroidism
61. Refsum syndrome (phytanic acid storage disease)
62. Renal transplantation
63. Retinitis pigmentosa-deafness-ataxia syndrome
64. Roy syndrome I (unilateral cataract associated with smoking)
*65. Senile posterior cortical cataract
66. Sjögren syndrome (secretinhibitor syndrome)
67. Silicone oil (intraocular)
*68. Steroid usage (topical or systemic)
69. Stickler syndrome (hereditary progressive arthro-ophthalmopathy)
70. Still disease (juvenile rheumatoid arthritis)
71. Toxocariasis (Nematode ophthalmia syndrome)
72. Trisomy (Patau syndrome)
73. Tuomaala-Haapanen syndrome (similar to pseudohypoparathyroidism)
74. Turner syndrome (gonadal dysgenesis)
75. Ultraviolet-B light
76. Uric acid (increased serum levels)
77. Vitrectomy for diabetic retinopathy
78. Weil disease (leptospirosis)
79. Werner syndrome (progeria of adults)
80. Yersiniosis


Extracted Table Posterior subcapsular cataract

**IRIDESCENT CRYSTALLINE DEPOSITS IN LENS**

1. Idiopathic
2. Hypothyroid (cretinism)
3. Hypocalcemia
   A. Postoperative-removal of thyroid and accidental parathyroid removal
   B. Idiopathic hypoparathyroidism
   C. Pseudohypoparathyroidism (hypoparathyroid cretinism) or with hyperphosphatemia (Albright disease)
   D. Pseudopseudohypoparathyroidism (brachymetacarpal dwarfism)
4. Myotonic dystrophy (Curschmann-Steinert syndrome)
5. Drugs, including the following:
   acetophenazine  gold Au 198  prochlorperazine
   amiodarone  gold sodium thiomalate  promazine
   auranofin  gold sodium thiosulfate  promethazine
   aurothioglucose  mercuric oxide  propiomazine
   aurothioglycanide  mesoridazine  silver nitrate
   butaperazine  methdilazine  silver protein
   carphenazine  methotrimipazine  thiethylperazine
   chlorpromazine  mild silver protein  thiopropazate
   chlorprothixene  perazine  thioridazine
   colloidal silver  pericyazine  thiothixene
   diazepam (?)  perphenazine  trifluoperazine
   diethazine  phenylmercuric acetate  triflupromazine
   ethopropazine  phenylmercuric nitrate  trimeprazine
   fluphenazine  piperacetazine

6. Cataract (coralliform and aculeiform) usually autosomal dominant; sometimes recessive

**OIL DROPLET IN LENS**

1. Anterior displacement of lens
   *2. Galactosemia-transferase deficiency (von Reuss syndrome)
3. Lenticous


**LENTICONUS (CONICAL LENS SURFACE PROTUBERANCE) AND LENTIGLOBUS (GLOBULAR LENS SURFACE PROTUBERANCE)**

1. Anterior - rare and usually bilateral
   *A. Alport syndrome (hereditary nephritis)
   B. Spina bifida
   C. Waardenburg syndrome (embryonic fixation syndrome)
2. Posterior - more common and often unilateral
   A. Associated with persistent hyperplastic primary vitreous
   B. Associated with remnants of hyaloid artery
   C. Familial posterior lenticous and microcornea
   *D. Lowe syndrome (oculocerebrorenal syndrome)
   E. Trauma


**LENS ABSORPTION**

1. Congenital rubella syndrome (German measles)
2. Hallermann-Streiff syndrome (dyscephalic mandibulooculofacial syndrome)
3. Surgical trauma as discission
4. Trauma, blunt or penetrating
EXFOILIATION OF LENS CAPSULE

In this condition, superficial layers of the lens capsule split off and float in aqueous as a fine membrane.

*1. Senile exfoliation
2. Toxic exfoliation
   A. Atrophic eyes
   B. Prolonged iridocyclitis
   C. Lodgment of metallic foreign body, such as iron or copper
3. Traumatic
   A. Perforating injury
   B. Contusions with suspensory ligament separated from a dislocated lens
4. Heat exposure, such as that experienced by glassblowers


MICROPHAKIA OR SPHEROPHAKIA OR MICROSPHEROPHAKIA (SMALL LENS OR HIGHLY SPHERIC LENS)

1. Achard syndrome (Marfan syndrome with dysostosis mandibulofacialis)
2. Alport syndrome (hereditary nephritis)
3. Familial anomaly
4. Homocystinuria syndrome
5. Hyperlysinemia
6. Lenticular myopia as recessive inheritance trait
7. Little syndrome (hereditary osteoarthritis)
8. Lowe syndrome (renal rickets)
*9. Marchesani syndrome (brachymorphy with spherophakia) (Weill-Marchesani syndrome)
10. Marfan syndrome (dolicostenomelia-arachnodactyly hyperchondroplasia
dystrophica mesodermalis congenita)
11. Peter anomaly (anterior chamber cleavage syndrome)
12. Reticular dystrophy of the retinal pigment epithelium
13. Rubella syndrome (Gregg syndrome)
14. Waardenburg syndrome (embryonic fixation syndrome)


DISLOCATED LENS

1. Achard syndrome (Marfan with dysostosis)
2. Adenoma of the nonpigmented epithelium of the ciliary body
3. Apert syndrome (sphenoacrocraniodesyndactyly)
4. Ascarisis
5. Associated ocular findings
   * A. Aniridia
   B. Coloboma of iris and choroid
   * C. Congenital glaucoma
   D. Ectopia lentis et pupillae
   E. Focal dermal hypoplasia (Goltz syndrome)
   F. High myopia
   G. Isolated lens dislocation (up)
   H. Megalocornea
   I. Microcornea
   J. Microspherophakia with hernia
   K. Pseudoxanthoma
eus.
   L. Retinitis pigmentosa
6. Autosomal recessive or dominant abnormality without other defects, except usually ectopic pupils
7. Capsular exfoliation syndrome
8. Crouzon disease
9. Cryptophthalmia syndrome (cryptophthalmos-syndactyly syndrome)
10. Dwarfism, genetic type
11. Ehlers-Danlos syndrome (cutis hyperelastica)
12. Gillum-Anderson syndrome (blepharoptosis, myopia, ectopia lentis)
13. Gorlin-Goltz syndrome
14. Grönblad-Strandberg syndrome (pseudoxanthoma elasticum)
15. Hereditary ectodermal dysplasia syndrome
*16. Homocystinuria-usually downward displacement of lens
17. Hyperlysinemia
18. Kniest syndrome
19. Late-onset localized junctional epidermolysis bullosa and mental retardation
20. Mandibulofacial dysostosis (Franceschetti syndrome)
*22. Marfan syndrome (dolichostenomelia-arachnodactyly-hyperchondroplasia dystrophia mesodermalis congenita)-usually superior displacement of lens
23. Molybdenum cofactor deficiency (combined deficiency of sulfite oxidase and xanthine dehydrogenase)
24. Oculodental syndrome (Peters syndrome)
25. Pfaundler syndrome
26. Pierre Robin syndrome
27. Pseudoxfoliation (exfoliation syndrome)
28. Recession of anterior chamber angle
29. Refsum syndrome
30. Retinal disinsertion syndrome
31. Retinoblastoma
32. Rieger syndrome (dysgenesis mesostromalis)
33. Spherophakia (see p. 400)
34. Spontaneous (degenerative)
35. Sprengel deformity
36. Sturge-Weber syndrome
37. Sulfite oxidase deficiency
38. Surgical accidents (iatrogenic)
39. Syphilis (lues)
40. Trauma as in Frenkel syndrome (ocular contusion syndrome), beesting, and following YAG laser
41. Treacher Collins syndrome (mandibulofacial dysostosis)
42. Uveitis
43. Wildervanck syndrome (cervicooculoacoustic syndrome)


**Extracted Table Dislocated lens**

**APHAKIA (ABSENCE OF LENS IN USUAL POSITION BEHIND THE IRIS)**

1. Congenital absence of lens-rare
2. Dislocation of the lens into vitreous cavity, anterior chamber, or subconjunctival area
3. Following cataract extraction
4. Gradual absorption of the lens


**EQUATORIAL LENS PIGMENTATION**
1. Associated with myopia and retinal detachment
2. Congenital malformation
3. Pigmentary glaucoma
4. Uveitis


**UNILATERAL CATARACTS**

1. Argon laser treatment
2. Trauma
3. Complicated cataract
   A. Anterior segment involvement, such as that because of the following:
      1. Acute and chronic corneal ulcer
      2. Iridocyclitis
      3. Chronic anterior uveitis
      4. Acute or chronic glaucoma
   B. Posterior segment involvement, such as that because of the following:
      1. Chronic posterior uveal inflammation
      2. Long-standing retinal detachment
      3. High myopia
      4. Hereditary retinal lesions
      5. Persistent hyperplastic primary vitreous
4. Roy syndrome I (unilateral cataract associated with smoking)
5. Congenital posterior polar lens changes
6. Glaucomatocyclitic crisis (Posner-Schlossman syndrome)
7. Conditions that give bilateral manifestations with earlier onset in one eye


**LENTICULAR DISEASE ASSOCIATED WITH CORNEAL PROBLEMS**

1. Aberfeld syndrome (ocular and facial abnormalities syndrome)-cataracts, microcornea
2. Acrodermatitis chronic a atrophicans - keratomalacia, corneal opacification, cataracts
3. Addison syndrome (idiopathic hypoparathyroidism)- keratoconjunctivitis, corneal ulcers, keratitic moniliasis, cataracts
4. Alport syndrome (hereditary nephritis)-anterior lenticonus, posterior polymorphous corneal dystrophy
5. Amiodarone - corneal deposits, anterior subcapsular cataracts
6. Amyloidosis - amyloid deposits of cornea, corneal dystrophy, pseudopodia lentis
7. Anderson-Warburg syndrome (oligophrenia-microphthalmos syndrome)-corneal opacification and lenticular destruction with a mass visible behind the lens
8. Andogsky syndrome (atopic cataract syndrome)-atopic keratoconjunctivitis, keratoconus, uveitis, subcapsular cataract
9. Aniridia-microcornea and subluxated lenses
10. Anterior-chamber cleavage syndrome (Reese-Ellsworth syndrome)-corneal opacities, anterior pole cataract
11. Anterior segment ischemia syndrome-corneal edema, corneal ulceration, cataract
12. Apert syndrome (absent digits cranial defects syndrome)-exposure keratitis, cataracts, ectopia lentis
13. Arteriovenous fistula (arteriovenous aneurysm)-bullous keratopathy, cataract
14. Aspergillosis-corneal ulcer, keratitis, cataract
*15. Atopic dermatitis (atopic eczema, Besnier prurigo)-keratoconjunctivitis, keratoconus, cataract
16. Autosomal dominant-cataracts and microcornea
17. Avitaminosis C (scurvy)-keratitis, corneal ulcer, cataract
18. Chalcosis (intraocular copper containing foreign body) deposits in Descemet membrane and anterior lens capsule
19. Chickenpox (varicella)-corneal ulcer, corneal opacity, keratitis, cataract
20. Chlorpromazine - corneal and lens opacities
21. Cholera-keratomalacia, cataract
22. Chromosome partial deletion (short-arm) syndrome-cataracts, corneal opacities
23. Chrysiasis (gold)-corneal and lens deposits
24. Cockayne syndrome (Mickey Mouse syndrome)-cataracts, band keratopathy, corneal dystrophy
25. Congenital spherocytic anemia (congenital hemolytic jaundice)-congenital cataract, ring-shaped pigmentary corneal deposits
26. Crouzon syndrome (Parrot-head syndrome)-exposure keratitis, cataract, corneal dystrophy
27. Cryptophthalmia syndrome (cryptophthalmos-syndactyly syndrome)-cornea differentiated from sclera, lens absence to hypoplasia, dislocation, and calcification
28. Cytomegalic inclusion disease (cytomegalovirus)-cataract, corneal opacities
29. Darier-White syndrome (keratosis follicularis)-keratosis, corneal subepithelial infiltrations, corneal ulceration, cataract
30. Dermatitis herpetiformis (Duhring-Broca disease)-corneal vascularization, cataract
31. Dermochondral corneal dystrophy (of François)-cataract, corneal dystrophy
32. Diabetes mellitus-cataract, corneal edema secondary to rubeosis
33. Diphtheria-keratitis, corneal ulcer, cataract
34. Down syndrome (trisomy 21)-lens opacities, keratoconus
35. Ehler-Danlos syndrome (fibrodysplasia elastica generalisata)-microcornea, keratoconus, lens subluxation
36. Electrical injury-corneal perforation, necrosis of cornea, anterior or posterior subcapsular cataracts
37. Endothelial dystrophy and anterior polar cataract (Dohlman)
38. Exfoliation syndrome (capsular exfoliation syndrome)-cataract, dislocated lens, corneal dystrophy, lens capsule exfoliation
39. Fabry disease (glycosphingolipid lipidosis)-cataract, corneal dystrophy
40. Folling syndrome (phenylketonuria)-corneal opacities, cataracts
41. Fuchs syndrome (I) (heterochromic cyclitis syndrome)-secondary cataract, edematous corneal epithelium
42. Goldscheider syndrome (epidermolysis bullosa)-bullous keratitis, corneal subepithelial blisters to corneal perforation, cataract
43. Gorlin-Goltz syndrome (multiple basal cell nevi syndrome)-cataract, corneal leukoma
44. Grönblad-Strandberg syndrome (elastorrhexis)-keratoconus, cataract, subluxation of lens
45. Hallermann-Streiff syndrome (oculomandibulofacial dyscephaly)-cataracts, microcornea
46. Hanhart syndrome (recessive keratosis palmoplantaris)-dendritic corneal lesions, keratitis, corneal haze, corneal neovascularization, cataract
47. Heerfordt syndrome (uveoparotid fever)-band keratopathy, keratoconjunctivitis, cataract
48. Hereditary ectodermal dysplasia syndrome (Siemens syndrome)-keratosis, corneal erosions, corneal dystrophy, cataract, lens luxation
49. Herpes simplex-keratitis, corneal ulcer, cataracts
50. Herpes zoster-keratitis, corneal ulcer, cataract
51. Histiocytosis X (Hand-Schüller-Christian syndrome)-pannus, bullous keratopathy, corneal ulcer, cataract
52. Hodgkin disease-keratitis, cataract
53. Homocystinuria syndrome-dislocated lens, cataract, keratitis
54. Hutchinson-Gilford syndrome (progeria)-cataract, microcornea
55. Hydatid cyst (echinococcosis)-keratitis, corneal abscess, cataract
56. Hypervitaminosis D-band keratopathy, cataract
57. Hypoparathyroidism-keratitis, cataract
58. Hypophosphatasia (phosphoethanolaminuria)-cataract, corneal subepithelial calcifications
59. Influenza-keratitis, cataract
60. Jadassohn-type anetodermal-keratoconus, cataract
61. Jadassohn-Lewandowsky syndrome (pachyonychia congenita)-corneal dyskeratosis, cataract
*62. Juvenile rheumatoid arthritis (Still disease)-band keratopathy, cataract
63. Kussmaul disease (periarteritis nodosa)-corneal ulcer, cataract
64. Kyrle disease (hyperkeratosis penetrans)-subcapsular cataracts, subepithelial corneal opacities
65. Leber congenital amaurosis- cataracts, keratoconus
66. Leri syndrome (carpal tunnel syndrome)-corneal clouding, cataract
67. Listerelliosis (listeriosis)-keratitis, corneal abscess and ulcer, cataract
68. Little syndrome (nail-patella syndrome)-microcornea, keratoconus, cataract
*69. Lowe syndrome (oculocerebrorenal syndrome)-cloudy cornea, cataracts, megalocornea, corneal dystrophy
70. Malaria-keratitis, cataract
71. Marchesani syndrome (brachymorphy with spherophakia)-lenticular myopia, ectopia lentis, megalocornea, corneal opacity
72. Marfan syndrome (arachnodactyly-dystrophia-mesodermalis congenita)- lens dislocation, cataract, megalocornea, lenticular myopia
73. Matsoukas syndrome (oculocerebroarticuloskeletal syndromes)- cataract, corneal sclerosis
74. Meckel syndrome (dysencephalia splanchnocystic syndrome)- sclerocornea, microcornea, cataract
75. Micro syndrome (autosomal recessive microcephaly, microcornea, congenital cataract, mental retardation, optic atrophy, and hypogenitalism)
76. Morbilli (rubeola, measles)- corneal ulcer, cataract
77. Mucolipidosis IV (ML IV)- corneal clouding, cataract
78. Myotonic dystrophy (Curschmann-Steinert syndrome)- lens opacity, cornea-epithelial dystrophy
79. Nematode ophthalmia syndrome (toxocariasis)- cataract, larvae present in the cornea
80. Neurodermatitis (lichen simplex chronicus)- keratoconjunctivitis, atopic cataracts, keratoconus
81. Neurofibromatosis 1
82. Neurofibromatosis 2
83. Ocular toxoplasmosis (toxoplasmosis)- keratitis, cataract
84. Oculodental syndrome (Peter syndrome)- macropenea, opacities of the corneal margin, ectopic lentis, corneoscleral staphyloma
85. O'Donnell-Pappas syndrome- premenile cataract, peripheral corneal pannus
86. Paget syndrome (osteitis deformans)- corneal ring opacities, cataract
87. Passow syndrome (status dysraphicus syndrome)- neuroparalytic keratitis, zonular cataract
88. Pemphigus foliaceus (Cazenave disease)- pannus, corneal infiltration, cataract
89. Pseudoxyphopharathyroidism (Seabright-Bantan syndrome)- punctate cataracts, keratitis
90. Pseudohypoparathyroidism (Seabright-Bantan syndrome)- punctate cataracts, keratitis
91. Radiation corneal ulcer, punctate keratitis, cataracts, exfoliation of lens capsule
92. Refsum syndrome (phytanic acid oxidase deficiency)- corneal opacities, cataracts
93. Relapsing polycondritis- corneal ulcer, cataracts, keratoconjunctivitis sicca
94. Retinal disinsertion syndrome-lens subluxation, keratoconus
95. Retrolental fibroplasia (RLF)- cataracts, corneal opacification
96. Rieger syndrome (dysgenesis mesodermalis corneae et irides)- microcornea, corneal opacities in Descemet membrane, dislocated lens
97. Romberg syndrome (facial hemiatrophy)- neuroparalytic keratitis, cataracts
98. Rothmund syndrome (telangiectasia-pigmentation-cataract syndrome)- cataract, corneal lesions
99. Rubella syndrome (German measles)- corneal haziness, cataracts, microcornea
100. Sabin-Feldman syndrome-posterior lenticonus, microcornea
101. Sanfilippo-Good syndrome (mucopolysaccharidosis III)- deposits in cornea and lens
102. Schafer syndrome (keratosis palmoplantaris syndrome)- lesions in the lower portion of the cornea, cataract
103. Schaumann syndrome (sarcoidosis syndrome)- keratitis sicca, band-shaped keratitis, complicated cataract
105. Scheie syndrome (mucopolysaccharidosis IS)-diffuse haze to marked corneal clouding, cataracts
106. Siderosis (intraocular iron foreign body)-iron deposition in lens and cornea
107. Stannus cerebellar syndrome-corneal vascularization, corneal opacities, cataracts
108. Steroids-cataract, may worsen certain types of corneal infections
109. Stevens-Johnson syndrome (erythema multiforme exudativum)-keratitis, corneal ulcers, cataracts, pannus
110. Stickler syndrome (hereditary progressive arthro-ophthalmopathy)-keratopathy, cataracts
111. Thioridazine-corneal and lens opacities
112. Toxic lens syndrome-pigment precipitation on the surface of an intraocular lens, chronic uveitis
113. Trisomy syndrome-corneal and lens opacities
114. Turner syndrome (gonadal dysgenesis)-corneal neubulae, cataracts
115. Ultraviolet radiation-band keratopathy, keratitis, discoloring of lens
116. van Bogaert-Scherer-Epstein syndrome (familial hypercholesterolemia syndrome)-lipid keratopathy, cataract, juvenile corneal arcus
117. von Recklinghausen syndrome (neurofibromatosis)-nodular swelling of corneal nerves, cataracts
118. Waardenburg syndrome (interoculoiridodermatoauditove dysplasia)-microcornea, cornea plana, lenticous
119. Wagner syndrome (hereditary hyaloideoretinal degeneration and palatoschisis)-corneal degeneration, including band-shaped keratopathy, cataracts
120. Ward syndrome (nevus jaw cyst syndrome)-congenital cataracts, congenital corneal opacities
121. Wegener syndrome (Wegener granulomatosis)-corneal ulcer, corneal abscess, cataract
122. Weil disease (leptospirosis)-keratitis, cataract
123. Werner syndrome (progeria of adults)-juvenile cataracts, bullous keratitis, trophic corneal defects *124. Wilson disease (hepatolenticular degeneration)-sunflower cataract, Kayser-Fleischer ring
125. Yersinia-scorneal ulcer, cataract
126. Zellweger syndrome (cerebrohepatorenal syndrome of Zellweger)-corneal opacities, cataract


<table>
<thead>
<tr>
<th>DRUGS ASSOCIATED WITH CATARACTS</th>
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<tbody>
<tr>
<td>acetohexamide</td>
</tr>
<tr>
<td>Acetophenazine</td>
</tr>
<tr>
<td>(phenothiazine)</td>
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<tr>
<td>acetylcholine</td>
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<tr>
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<tr>
<td>azathioprine (?)</td>
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<td>(BCG) vaccine (?)</td>
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PUVA (psoralens and ultraviolet A) (?)
quinalcrine
silicone
sulfonamides (maternal ingestion)
sulindac (?)
tamoxifen
tetracaine
thiethylperazine
thiopropazate
thioproperazine
timolol (?)
tolazamide
tolbutamide
triamcinolone
trifluoperazine
triflupromazine
triflupromazine
trimeprazine
trioxxalen (?)
thioridazine
thiophixene
urokinase (?)
tolbutamide
vatin D2 (?)
triamcinolone
vatin D3 (?)
trifluoperazine
warfarin


SYNDROMES AND DISEASES ASSOCIATED WITH CATARACTS

1. Aberfeld syndrome (blepharophimosis associated with generalized myopathy)
2. Acrodermatitis chronic atrophicans
3. Addison syndrome (adrenal cortical insufficiency)
4. Albinism
5. Albright hereditary osteodystrophy (pseudohypoparathyroidism)
6. Alopecia areata
7. Alport syndrome (hereditary nephritis)
8. Alström disease (cataract and retinitis pigmentosa)
9. Andogsky syndrome (atopic cataract syndrome)
10. Anterior segment ischemia syndrome
11. Apert syndrome (acrocephalosyndactylism syndrome)
12. Apical malformations associated with cataracts
13. Arteriovenous fistula
14. Arthrogryposis multiplex congenita
15. Aspergillosis
16. Atopic dermatitis syndrome
17. Autosomal dominant foveal hypoplasia and presenile cataract syndrome (O'Donnell-Pappas syndrome)
18. Bassen-Kornzweig syndrome (abetalipoproteinemia)
20. Bonnevie-Ullrich syndrome (pterygolymphangiectasia)
21. Bourneville syndrome (tuberous sclerosis)
22. Buerger disease (thromboangiitis obliterans)
23. Caisson syndrome (bends)
24. Capsular exfoliation syndrome
25. Carotid artery syndrome
26. Cataract and hypertrophic neuropathy-autosomal recessive
27. Cataract with microcornea and coloboma of iris-autosomal dominant
28. Cataract, floriform-autosomal dominant
29. Cataract and cardiomyopathy-autosomal recessive
30. Cataract, congenital, or juvenile-autosomal recessive
31. Cataract, congenital total, with posterior sutural opacities in heterozygotes-X-linked
32. Cataract, congenital with absence deformity of leg-autosomal recessive
33. Cataract, congenital, with microcornea or slight microphthalmia-X-linked
34. Cataract, cortical, and congenital ichthyosis-autosomal recessive
35. Cataract, mental retardation, hypogonadism (Martsolf syndrome)
36. Cataract, microcephaly, arthrogryposis kyphosis syndrome (CAMAK syndrome)-autosomal recessive
37. Cataract microcephaly, failure to thrive, kyphoscoliosis syndrome (CAMFAK syndrome)-autosomal recessive
38. Cataract, nuclear and total nuclear-usually autosomal dominant rarely recessive
39. Cataract, zonular, and nystagmus-X-linked
40. Cat-eye syndrome (Schmid-Fraccaro syndrome)
41. Cerebral cholesterinosis (cerebrotendinous xanthomatosis)
42. Cerebellar ataxia, cataract, deafness, and dementia or psychosis
43. Cerebral palsy
44. Cerebrohepatorenal syndrome (Smith-Lemli-Opitz syndrome)
45. Cerebrotendinous xanthomatosis
46. Cholera
47. Chromosome 13q partial deletion (long-arm) syndrome
48. Chromosomal 3; 18 translocation
49. Chromosome deletion (short-arm) syndrome
50. Cockayne syndrome (dwarfism with retinal atrophy and deafness)
51. Cerebrooculofascioskeletal syndrome (COFS syndrome)
52. Congenital cataract and hypertrophic cardiomyopathy syndrome
53. Congenital cataract with oxycephaly (tower skull)
54. Congenital hemolytic icterus
55. Congenital ichthyosiform erythroderma
56. Congenital rubella syndrome (German measles)
57. Conradi syndrome (stippled epiphyses syndrome)
58. Comedo-cataract
59. Craniofacial dysostosis (Crouzon disease)
60. Cretinism (hypothyroidism)
61. Crome syndrome (congenital cataracts, epileptic fits, mental retardation, small stature)
62. Cushing syndrome
63. Cytomegalovirus
64. Darier-White syndrome (keratosis follicularis)
65. DeBarsy syndrome
66. Dermatitis herpetiformis
*67. Diabetes mellitus (Willis disease)
68. Diarrhea
69. Diphtheria
70. Ectodermal dysplasia
71. Edward syndrome
72. Electrical injury
73. Ellis-van Creveld syndrome (chondroectodermal dysplasia)
74. Engelmann syndrome (diaphyseal dysplasia)
75. Epidermal nevus syndrome (ichthyosis hystrix)
76. Fabry disease (diffuse angiokeratosis)
77. Familial congenital cataracts, microcornea, abnormal irides, nystagmus, and glaucoma syndrome
78. Familial congenital cataract, nonprogressive neurologic disorders, and mental deficiency syndrome
79. Familial histiocytic dermatooarthitis syndrome
80. Familial hypogonadism syndrome
81. Familial t(2;16) translocation
82. Fetal alcohol syndrome
83. Folling syndrome (phenylketonuria)
84. François dyscephalic syndrome (Hallermann-Streiff syndrome)
85. Frenkel syndrome (ocular contusion syndrome)
86. Fuchs syndrome (I) (heterochromic cyclitis syndrome)
87. Galactokinase deficiency-autosomal recessive
*88. Galactosemia-transferase deficiency
89. Goldenhar syndrome (oculoauriculo-vertebral dysplasia)
90. Goldscheider syndrome (epidermolysis bullosa)
91. Gorlin-Goltz syndrome (multiple basal cell nevi syndrome)
92. Grönhed-Strandberg syndrome (pseudoxanthoma elasticum)
93. Gyrate atrophy (ornithine ketoacid aminotransferase deficiency)
94. Hagberg-Santavuori syndrome (neuronal ceroid-lipofuscinosis)
95. Hallermann-Streiff syndrome (oculomandibulofacial dyscephaly)
96. Hand-Schüller-Christian syndrome (histiocytosis X)
97. Harada disease (uveitis-vitiligo-alopécia-poliosis syndrome)
98. Heerfordt syndrome (uveoparotid fever)
99. Hemifacial microsomia syndrome (François-Haustrate syndrome)
100. Herpes simplex virus
101. Hilding syndrome (destructive iridocyclitis and multiple joint dislocations)
102. Hodgkin disease
103. Homocystinuria
104. Hookworm disease
105. Hruby-Irvine-Gass syndrome (cystoid maculopathy following cataract extraction with vitreous loss)
106. Hutchinson-Gilford syndrome (progeria)
107. Hydatid cyst
108. Hypercalcemia (adult)
109. Hypercalcemia (infantile) with mental retardation (supravalvular aortic stenosis syndrome)
110. Hyperprolactinemia
111. Hypertrophic cardiomyopathy
112. Hypervitaminosis A
113. Hypervitaminosis D
114. Hypocalcemia
115. Hypoglycemia
116. Hypoparathyroidism
117. Hypophosphatasia (phosphoethanolaminuria)
118. Incontinentia pigmenti achromians
119. Infantile hypoglycemia (male)
120. Influenza
121. Infrared radiation
122. Intrauterine infections
   A. herpes virus
   B. mumps
   C. rubella
   D. toxoplasmosis
   E. vaccinia
123. Jacobsen-Brodwall syndrome
124. Jadassohn-Lewandowsky syndrome (pachyonychia congenita)
125. Karsch-Neugebauer syndrome (nystagmus-split hand syndrome)
126. Klippel-Trenaunay-Weber syndrome (angioosteohypertrophy syndrome)
127. Krause syndrome (congenital encephalophthalmic dysplasia)
128. Kussmaul disease (periarteritis nodosa)
129. Kyrle disease (hyperkeratosis penetrans)
130. Lanzieri syndrome (craniofacial malformations)
131. Laser treatment for retinopathy of prematurity
132. Laurence-Moon-Biedl syndrome (retinitis pigmentosa-polydactyly-adiposogenital)
133. Leber syndrome (optic atrophy-amaurosis-pituitary syndrome)
134. Leiomyoma
135. Leri syndrome (carpal tunnel syndrome)
136. Lightning
137. Listerellosis
*138. Lowe syndrome (oculocerebrorenal syndrome)
139. Majewski syndrome (short-rib polydactyly syndrome)
140. Malaria
141. Male Turner syndrome (Noonan syndrome)
142. Malignant hyperpyrexia syndrome
143. Mandibulofacial dysostosis (Franceschetti syndrome)
144. Mannosidosis
145. Maple-syrup urine disease (branched-chain ketoaciduria
146. Marfan syndrome (arachnodactyly dystrophia mesodermaliscongenita)
147. Marinesco-Sjögren syndrome (congenital cataract-oligophrenia syndrome)
148. Marshall syndrome (atypical ectodermal dysplasia)
149. Martsolf syndrome
150. Matsoukas syndrome (oculocerebroarticuloskeletal syndrome)
151. Meckel syndrome (dysencephalia splanchnocystic syndrome)
152. Microcephaly, microphthalmia, cataracts, and joint contractures syndrome
153. Microphthalmia-congenital anterior polar cataract syndrome-autosomal dominant
154. Micro syndrome
155. Miller syndrome (Wilms aniridia syndrome)
156. Monilethrix
157. Morgan syndrome (intracranial exostosis)
158. Morquio-Brailsford syndrome (mucopolysaccharidoses IV)
159. Multiple sulfatase deficiency
160. Myopic (high)
161. Myotonic dystrophy (Curschmann-Steinert syndrome)
162. Nail-patella syndrome (Little syndrome)
163. Nance-Horan syndrome (cataract-dental syndrome)
164. Neurodermatitis
165. Neurofibromatosis 1 (von Recklinghausen syndrome)
166. Neurofibromatosis 2 (central neurofibromatosis)
167. Nieden syndrome (telangiectasia-cataract syndrome)
168. Norrie disease
169. Oculootoororenoerythropoietic disease
170. Optic atrophy, cataract, and neurologic disorder-autosomal dominant
171. Osteogenesis imperfecta congenita, microcephaly, and cataracts-autosomal recessive
172. Osteopetrosis (Albers-Schönberg syndrome)
173. Oxyccephaly
174. Pachyonychia congenita syndrome
175. Paget syndrome (idiopathic hyperphosphatasemia)
176. Pallister-Killian syndrome
177. Partial trisomy 10q trisomy
178. Passow syndrome (syringomyelia)
179. Patau syndrome
180. Pellagra (avitaminosis B2)
181. Pemphigus foliaceus (Cazenave disease)
182. Pernicious anemia syndrome (vitamin B12 deficiency)
183. Pierre Robin syndrome (micrognathia-glossoptosis syndrome)
184. Prader-Labhart-Willi syndrome (hypogenital dystrophy with diabetic tendency)
185. Pseudoexfoliation syndrome
186. Pseudohypoparathyroidism
187. Radiation
188. Reese-Elsworth syndrome (anterior chamber cleavage syndrome)
189. Refsum syndrome (phytanic acid storage disease)
190. Renal failure (chronic)
191. Renal transplantation
192. Retinal ischemic infarction syndrome
193. Retinitis pigmentosa-deafness-ataxia syndrome
194. Rhizomelic chondrodysplasia punctata
195. Riboflavin deficiency syndrome (oculoorogenital syndrome)
196. Ring chromosome in the D group
197. Robert syndrome
198. Robert pseudothalidomide syndrome
199. Romberg syndrome (facial hemiatrophy)
200. Rothmund syndrome (infantile poikiloderma)
*201. Roy syndrome I (unilateral cataract associated with smoking)
202. Roy syndrome II - nuclear cataract associated with smoking
203. Rubeola (measles)
204. Rubinstein-Taybi syndrome (broad-thumbs syndrome)
205. Scaphocephaly syndrome (craniofacial dysostoses)
206. Schaefer syndrome (congenital dyskeratosis)
207. Schwartz syndrome
208. Scurvy (avitaminosis C)
209. Sickle cell disease (Herrick syndrome)
210. Siemen syndrome (congenital atrophy of the skin)
211. Sjögren syndrome (secretion inhibitor syndrome)
212. Sjögren-Larsson syndrome (oligophrenia ichthyosis)
213. Smith-Lemli-Opitz syndrome
214. Split-hand with congenital nystagmus, fundal changes, cataracts-autosomal dominant
215. Spondyloepiphyseal dysplasia (SED) dwarfism
216. Stannus cerebellar syndrome (vitamin B2 deficiency)
217. Stickler syndrome (hereditary progressive arthro-ophthalmopathy)
*218. Still disease (juvenile rheumatoid arthritis)
219. Thrombocytopenia-absent radius (TAR) syndrome
220. Toxocariasis (nematode ophthalmia syndrome)
221. Treacher Collins syndrome (mandibulofacial dysostosis)
222. Trichomegaly, spherocytosis, and cataract - autosomal dominant
223. Trichorrhexis nodosa (argininosuccinicaciduria)
224. Trisomy 13 syndrome (Patau syndrome)
225. Trisomy 16 syndrome (Edward syndrome)
226. Trisomy 20p syndrome
227. Trisomy 21 (Down syndrome)
228. Tuomaala-Haaninen syndrome
229. Turner syndrome (gonadal dysgenesis)
230. Tyrosinosis (Hanhart syndrome)
231. Usher syndrome (hereditary retinitis pigmentosa-deafness syndrome)
232. Uvea-touch syndrome
233. Van der Hoeve syndrome (brittle-bone disease)
234. Van Bogaert-Scherer-Epstein syndrome (primary hyperlipidemia)
235. Varicella infection
236. von Recklinghausen syndrome (neurofibromatosis)
237. Wagner syndrome (hypoideoretinal degeneration)
283. Warburg syndrome (hydrocephalus,agyria, and absent cortical laminar retinal dysplasia with or without encephalocele)
239. Ward syndrome (nevus-jaw cyst syndrome)
240. Wegener syndrome (Wegener granulomatosis)
241. Weil disease (leptospirosis)
242. Werner syndrome (scleropoilioderma)
243. Wilson disease (hepatolenticular degeneration)
244. Yersinia infection
245. Zellweger syndrome (cerebrohepatorenal syndrome)
246. 31 syndrome  
247. 4p syndrome  
248. 18p syndrome  
249. 18q syndrome


**LENTICULOCORNEAL ADHERENCE (LENS ADJACENT TO ENDOTHELIUM OF CORNEA)**

1. Acquired anterior corneal disease as ulcer with perforation or trauma  
2. Aniridia  
3. Peters anomaly (oculodental syndrome)  
4. Rieger anomaly (dysgenesis mesostromalis)


**SPASM OF ACCOMMODATION**

This condition involves increased tone of ciliary body with increased convexity of crystalline lens (see p. 619-622).

1. Alcoholism  
2. Cerebrovascular accident  
3. Contusion injury to the globe or head  
4. Cyclic oculomotor palsy or spasm  
5. Diabetes mellitus
6. Drugs, such as aceclidine, acetylcholine, carbachol, demecarium, DFP, digitalis, echothiophate, guanethidine, isofluorophate, methylene blue, morphine, neostigmine, opium, physostigmine, pilocarpine
7. Fatigue cramp of overworked ciliary muscle; most frequent with compound hyperopia and mixed astigmatism associated with anisometropia
8. Graves disease (hyperthyroidism, Basedow syndrome)
9. Infectious, such as diphtheria, helminthic infestations, or sinus disease
10. Irritative lesions of brain stem and oculomotor trunk, such as epidemic encephalitis, tabes, meningitis, influenza, scleritis, measles, or orbital inflammation
11. Middle cerebral artery occlusion
12. Ocular inflammation, such as ciliary muscle irritant
13. Pineal tumor
14. Reflex irritation, such as in trigeminal neuralgia
15. Sympathetic paralysis
16. Trauma


**PARESIS OF ACCOMMODATION**

This condition involves partial or total loss of physiologic ability to change the shape of the lens and thus the focus of the eye (see Mydriasis, p. 622-623); this ability is related to age (see Acquired Hyperopia, p. 344-348).

*1. Presbyopia—gradual decrease in amplitude of accommodation related to age
2. Accommodative insufficiency
   A. Asthenic individuals
   B. Illness or debilitation, including intestinal toxemia, tuberculosis, influenza, whooping cough, measles, and tonsillar and dental infections
   C. Anemia
   D. Overwork
   E. Whiplash injury
3. Ciliary body aplasia—with or without pupillary and iris abnormalities
4. Iridocyclitis—acute and chronic
5. Glaucoma with atrophy of ciliary body
6. Choroidal metastasis with suprachoroidal extension*
7. Trauma, such as tears in iris sphincter, tears at root of iris, or recession of the anterior chamber angle with posterior displacement of the ciliary attachment and ocular hypotension
8. Rupture of zonular fibers and partial subluxation of lens
9. Myotonic dystrophy (Curschmann-Steinert syndrome)
10. Drugs, including the following:

- acetazolamide
- acetophenazine
- adiphenine
- alcohol
- alprazolam
- ambutonium
- aminosalicylate (?)
- aminosalicylic acid (?)
- amitriptyline
- amodiaquine
- amoxapine
- amphetamine
- anisindione
- anisotropine
- antazoline
- atropine
- belladonna
- bendroflumethiazide
- benzathine penicillin G
- benzphetamine
- benzthiazide
- benztropine
- betamethasone
- betahistine
- biphenyl propane
- biperiden
- bromide
- butaperazine
- captopril (?)
- caramiphen
- carbachol
- carbamazepine
- carbinoxamine
- carbon dioxide
- carphenazine
- chloramphenicol
- chlordiazepoxide
- chlorisondamine
- chlorothiazide
- chlorphenoxamine
- chlorpromazine
- chlorthalidone
- cimetidine
- clomipramine
- clonazepam
- chlorzepate
- clomiphene
- clopxamine
- cocaine
- cortisone
- cyclopentolate
- cyclothiazide
- cymicine
- desipramine
- dexamethasone
- dextroamphetamine
- diacetylmorphine
- diazepam
- dibucaine
- dichlorphenamide
- dicyclomine
- diethazine
- diphenylanil
- diphenadione
- diphenhydramine
- diphenylpyraline
- emetine
- ergot
- ethopropazine
- fluorometholone
- fluphenazine
- glycopyrrolate
- hexamethonium
- hexocyclium
- homatropine
- hydrochlorothiazide
- hydrocortisone
- hydroflumethiazide
- hydromorphone
- hydroxyamphetamine
- imipramine
- indapamide
- iodide and iodine
- solutions and compounds
- isoniazid
- isopropamide
- maprotiline
- marijuana
- mecamylamine
- medrysone
- mepenzolamine
- meprobamate
- mescaline
- mesoridazine
- methacholine
- methamphetamine
- methanetheline
- methaqualone
- methazolamide
- methdilazin
- methixene
- methotrimoprazine
- methscopolamine
- methylclohexazine
- methylatropine nitrate
- methylene blue
- methyprylon
- methysergide
- metolazone
- mianserin
- midazolam
morphine  piroxicam (?)  temazepam
nalidixic acid  poldine  tetanus immune globulin
naproxen  polythiazide  tetanus toxoid
nitrazepam  potassium penicillin G  tetracaine
nortriptyline  potassium penicillin V  tetraethylammonium
opium  potassium phenethicillin  tetrahydrocannabinol
orphenadrine  pralidoxime  thiethylperazine
oxazepam  prazepam  thioproprazine
oxymorphone  prednisolone  thiopropazine
oxyphencyclidine  primidone  thioridazine
oxyphenonium  procaine penicillin G  thiothixene
pargyline  procarbazine  triazolam
pentazocine  prochlorperazine  trichlormethiazide
pentolinium  procyclidine  trichloroethylene
perazine  promazine  tridihexethyl
perhexazine  promethazine  trifluoperazine
phendimetrazine  propantheline  trifluperidol
phenindione  propiomazine  triflupromazine
phenmetrazine  propranolol  trihexyphenidyl
phentermine  protriptyline  trimethapram
pilocarpine  psilocybin  trimethaphan
pimozide  quinethazone  trimethidinium
pipenzolate  radioactive iodides  tripelequin
piperacetazine  rubella virus vaccine  tropicamide
piperazine  (live)  vinblastine
piperidolate  scopolamine  vincristine
piperocaine  streptomyacin

11. Neurogenic causes
   A. Infectious conditions
      (1) Epidemic encephalitis
      (2) Anterior poliomyelitis
      (3) Exanthemas and acute infections, such as scarlet fever, mumps,
           measles, influenza, typhoid fever, dengue fever, viral hepatitis, amebic
           dysentery, and malaria
      (4) Herpes zoster
      (5) Syphilis (lues)
      (6) Tuberculosis
      (7) Leprosy (Hansen disease)
      (8) Focal infections, such as from teeth or nasal sinuses
   B. Toxic conditions
      (1) Alcohol
      (2) Lead
      (3) Arsenic
      (4) Carbon monoxide
(5) Diphtheritic paralysis
(6) Botulism
(7) Extensive bum
(8) Snake venom

C. Degenerative conditions
(1) Congenital hereditary ophthalmoplegia
(2) Progressive congenital ophthalmoplegia
(3) Hereditary ataxia
(4) Myotonic dystrophy (Curschmann-Steinert syndrome)
(5) Myasthenia gravis

D. Metabolic conditions
(1) Acute hemorrhagic anterior polioencephalitis of Wernicke
(2) Diabetes mellitus
(3) Lactation
(4) Following pregnancy

E. Isolated internal ophthalmoplegia

F. Isolated failure of near reflex, such as with inverse Argyll-Robertson pupil

G. Lesions of parasympathetic nuclei in midbrain
(1) Encephalitis
(2) Pineal tumor
(3) Other signs of mesencephalic disease, including multiple sclerosis, infectious polyneuropathy, and vascular lesions
(4) Syphilis-bilateral

H. Trauma to head or neck
(1) Cerebral concussion
(2) Cervicocranial extension injuries


Pseudodetachment of vitreous 421
Anterior vitreous detachment 421
Posterior vitreous detachment 422
Vitreous hemorrhage 424
Diagnostic table 426
Vitreous opacities 429
Persistent hyperplastic primary vitreous 430
Beads in vitreous 430
Asteroid hyalosis versus synchysis scintillans 431
Complications following operative vitreous loss 431
Postoperative vitreous retraction 431
Vitreous cyst 432
Vitreous liquefaction 432

PSEUDODETACHMENT OF VITREOUS (CONDITIONS SIMULATING DETACHMENT OF VITREOUS)

1. Enormous cavity in the vitreous body with a relatively thin posterior wall
2. Membranous formations within the vitreous associated with uveitis and hemorrhage
3. Outline of the ascending portion of Cloquet canal just anterior to the disc


ANTERIOR VITREOUS DETACHMENT
In this condition, the anterior vitreous cortex may be separated from the posterior lens or posterior zonular fibers.

1. Retrolenticular-usually caused by vitreous shrinkage
   * A. Trauma (most common)
   B. Hemorrhage-usually secondary to trauma
   C. Senescence (rare)
   D. Inflammation
   E. Retinal detachment (see p. 487)
   *F. Iatrogenic after injection of vitreous substitutes (gas)
2. Retroocular
   A. Vitreous shrinkage (see p. 432)
   B. Ciliary body tumor
   C. Blood
   D. Exudate
3. Retrolenticular and retroocular combined occurs with rupture of the hyaloideocapsular ligament
POSTERIOR VITREOUS DETACHMENT

1. Complete posterior detachment
   A. Simple detachment-occurs in young persons
      (1) Exudate from chorioretinal focus
      (2) Hemorrhage between the vitreous and the retina
      (3) Retraction of the cortical vitreous caused by exudate within the vitreous
      (4) Vitreous hemorrhage in a young individual with vitreous shrinking due to thrombosis of central retinal vein, retinal neovascularization
   B. Complete posterior detachment with collapse
      (1) Senescent changes are primary cause
      (2) Uveitis
      (3) Trauma
      (4) Hemorrhage
      (5) Sodium hyaluronate
   C. Funnel-shaped posterior detachment
      (1) Perforating injuries of globe
      (2) Retinal neovascularization
      (3) Massive vitreous detachment
   D. Atypical complete posterior detachment-residual adherence of vitreous to a peripheral retinal area
      (1) Focus of chorioretinitis
      (2) Following cataract extraction with loss of vitreous
      (3) Following perforating wounds
      (4) Posterior uveitis with inflammatory cells

2. Partial posterior detachment (unusual)
   A. Superior detachment-primarily a senescent change; generally forerunner of posterior vitreous detachment with collapse
   B. Partial posterior detachment (not infrequent)
      (1) Preretinal hemorrhage
      (2) Retinal neovascularization
   C. Partial lateral or partial inferior detachment
      (1) Focus of choroiditis
      (2) Circumscribed retinal periphlebitis
      (3) Intraocular foreign body


**Extracted Table Posterior Vitreous Detachment**

**VITREOUS HEMORRHAGE**

1. Acquired lues (syphilis)
2. Arsenic toxicity
3. Ascariasis
4. Avulsed retinal vessel syndrome
5. Battered baby syndrome (Silverman syndrome)
6. Behçet syndrome (dermatostomatoophthalmic syndrome)
7. Blood disease-retinal hemorrhage breaking into vitreous
   A. Anemias
      (1) Aplastic anemia
      (2) Hemolytic anemia
      (3) Hypochromic anemia
      (4) Pernicious anemia
   B. Dysproteinemias - macroglobulins and cryoglobulins
   C. Hemophilia associated with trauma
   D. Leukemias
   E. Multiple myeloma (Kahler disease)
   F. Polycythemia vera (Vaquez disease)
   G. Thrombocytopenic purpura
8. Coats disease (retinal telangiectasia)
9. Collagen disease
   A. Dermatomyositis
   B. Disseminated lupus erythematosus (Kaposi-Libman-Sacks syndrome)
   C. Polyarteritis nodosa (Kussmaul disease)
   D. Scleroderma (progressive systemic sclerosis)
10. Complete posterior vitreous detachment with collapse (10%-15% at time of event)
11. Cysticercosis
12. Dengue fever
13. Diabetes mellitus-proliferative retinopathy
14. Dislocation of intraocular lenses
15. Disseminated intravascular coagulation
16. Drusen of optic disc
17. Eales disease
18. Exudative age-related macular degeneration
19. Familial exudative vitreoretinopathy
20. Ganciclovir Implant
21. Gronblad-Strandberg syndrome (systemic elastodystrophy)
22. Hemorrhages in the newborn
   A. Hemorrhagic disease of the newborn factor VII and prothrombin deficiency
   B. Persistent vessels of the hyaloid system
C. Retinal hemorrhage of newborn breaking through to vitreous cavity
23. Iatrogenic globe perforation associated with strabismus surgery
24. Indomethacin reaction
25. Influenza
26. Intraocular foreign body
27. Intraocular tumor
28. Hypertension (venous occlusive disease)
29. Juvenile retinoschisis
*30. Macroneurysm (retinal arterial)
31. Malaria
32. Malignant melanoma
33. Migration from anterior bleeding as from angle-supported, iris-supported, or posterior chamber lenses
*34. Neovascularization following vascular occlusion (primarily venous occlusive disease)
35. Neovascularization of cataract wound
36. Ocular ischemic syndrome with neovascularization of disc
37. Pars planitis
38. Persistent hyaloid artery
39. Persistent hyperplastic primary vitreous (PHPV)
*40. Posterior vitreous detachment (PVD)
41. Purtscher disease (traumatic retinal angiopathy)
42. Retinal angiomatosis (von Hippel disease)
*43. Retinal break or tear with or without retinal detachment and avulsed retinal vessels
*44. Retinal hemorrhage, including vein occlusion and sickle retinopathy, arterial macroaneurysm
45. Retinal tacks (intrusion)
46. Retinoblastoma
47. Retinopathy of prematurity-proliferative stage
48. Scleral buckle (intrusion)
49. Sleep apnea
50. Sickle cell disease (Herrick syndrome)-SA, SS, or SC
51. Surgical cataract complication with lenticular fragments dislocated into vitreous
52. Terson syndrome of associated vitreous and subarachnoid hemorrhage syndrome
53. Thalassemia (Cooley anemia)
54. Thromboangiitis obliterans (Buerger disease)
55. Tissue plasminogen activator (t-PA)
56. Tuberculous sclerosis
57. von Hippel-Lindau disease (angiomas retinae)
58. von Willebrand syndrome
59. Trauma
60. Traumatic asphyxia
61. Tuberculosis
62. Uveitis (associated with)
63. Varicella zoster


**Extracted Table: Vitreous hemorrhage**

**VITREOUS OPACITIES**

1. Opaque sheets anterior to the vitreous
   - A. Elschnig pearls after extracapsular cataract extraction or needling (posterior capsule opacification)
   - B. Normal posterior capsule-often following extracapsular cataract extraction or needling
   - C. Soemmerring ring following extracapsular cataract extraction or needling
   - D. Vitreous adhesions to iris, capsule, or intraocular lens (IOL) after cataract extraction with vitreous loss

2. Pseudoglioma-leukokoria (see p. 357)

3. Scattered opacities
   - A. Amyloid disease-rare (seen in older persons)
   - B. Ankylosing spondylitis
   - C. Coagula of the colloid basis of the gel
   - D. Crystalline deposits
     - (1) Asteroid hyalosis
     - (2) Synchysis scintillans
   - E. Endophthalmitis
   - F. Heterochromic uveitis-in persons 20 to 50 years of age; of all uveitis, iris atrophy, lens changes
   - G. Myeloma, multiple-rare: in persons 50 to 70 years old, associated with bone pain, anemia
   - H. Myopia, severe
   - I. Pigment cells-posttraumatic (hemorrhage), senile, or melanotic, associated with rhegmatogenous retinal detachment.
   - J. Protein coagulaplasmoid vitreous
     - (1) Choroidal tumors (very rare-reported in metastatic breast cancer once)
     - (2) Contusions
     - (3) Intermediate uveitis (pars planitis)
     - (4) Retinochoroiditis
K. Red blood cells (see vitreous hemorrhage, p. 424)
L. Snowball opacities-rare, associated with pars planitis or sarcoidosis, endophthalmitis (indolent)
M. Tissue cells-epithelial, histiocytic, glial
N. Toxoplasmosis-active
O. Tumor cells-retinoblastoma in older child, reticulum cell sarcoma (older persons)
P. Vitreous degeneration-Wagner disease, Ehlers-Danlos syndrome, and Marfan syndrome, senescent aging changes, myopia
Q. Whipple disease
R. White blood cells-inflammatory disease, vitreitis
S. Retinitis pigmentosa

4. Single opacities
   A. Anterior hyaloid remnant (Mittendorf dot)-25% normal eyes, dot on posterior lens surface
   B. Hyaloid remnants (uncommon)-persistent hyperplastic primary vitreous
   C. Foreign body-history of trauma or surgery
   D. Dislocated lens (see p. 401)
   E. Parasitic cysts
      (1) Hydatid disease (echinococciosis)-rare, children and young adults, tropical
      (2) Cysticercosis-rare
   F. Vitreous detachment-common in older or myopic persons


**PERSISTENT HYPERPLASTIC PRIMARY VITREOUS**

1. Cerebrooculodysplasia-muscular dystrophy
2. Fetal alcohol syndrome
3. Incontinentia pigmenti
4. Norrie disease
5. Septooptic pituitary
6. Sporadic unilateral and isolated finding
7. Trisomy 13
8. Warburg syndrome

**BEADS IN VITREOUS (SNOWBALLS IN VITREOUS)**

1. African eye-worm disease (loiasis)
2. Amyloidosis (Lubarsch-Pick syndrome)
3. Behçet syndrome (dermatostomatoophthalmic syndrome)
4. Birdshot retinochoroidopathy
5. Brucellosis (Bang disease)
6. Familial exudative vitreoretinopathy (Criswick-Schepens syndrome)
7. *Haemophilus influenzae*
8. Irvine syndrome
9. Jacobsen-Brodwall syndrome
10. Ocular toxocariasis
11. Ocular toxoplasmosis
12. Oculootooronoerythropoietic disease
13. Pars planitis
14. Retinoblastoma
15. Sarcoidosis
16. Severe uveitis
17. Sympathetic ophthalmia
18. Toxic lens syndrome
19. Typhus (Japanese river fever)


**COMPLICATIONS FOLLOWING OPERATIVE VITREOUS LOSS**

1. Inflammatory complications
   A. Irritable eye (chronically)
   B. Recurrent or persistent uveitis
   C. Vitreitis with vitreous opacities
2. Wound complications
   A. Epithelial invasion or downgrowth
   B. Fibrous ingrowth
   C. Fistula or gaping of wound (with or without vitreous wick syndrome)
   *D. Infection or endophthalmitis
   E. Excessive astigmatism
3. Corneal complications
   A. Corneal edema (vitreocorneal touch)
B. Bullous keratopathy
C. Corneal opacification

4. Secondary glaucoma
   A. Vitreous obstruction of anterior chamber angle
   B. Pupillary block (iridohyaloid adhesions, anterior hyaloid displacement, uveitis)
   C. Iris and vitreous adherence to wound (peripheral anterior synechiae)

5. Fibroblastic and traction phenomena
   A. Pupillary membrane
   B. Pupillary distortion- "peaked" or updrawn synechiae
   C. Cystoid macular edema (CME)
   D. Retinal detachment
   E. Optic neuritis or papilledema
   F. Vitreous hemorrhage
   G. Posterior vitreous detachment


**POSTOPERATIVE VITREOUS RETRACTION**

Usually, this condition is manifested by circular equatorial retinal fold or star-shaped retinal fold.

1. Accidental perforation of the sclera at operation, which may be associated with hemorrhage and loss of vitreous resulting in a pathologic formation of new epiretinal membrane or proliferative vitreoretinopathy
2. Giant retinal breaks allowing a large area of direct contact between the choroid and the vitreous
3. Perforating diathermy and excessively strong or repeated applications of superficial diathermy, which may cause vitreous hemorrhage or thermal injury to the vitreous; impairment of chorioretinal blood circulation may result in exudation and hemorrhage into the vitreous
4. Venous stasis caused by the compression of vortex veins by the indentation resulting from a buckling procedure


**VITREOUS CYST (CYSTIC STRUCTURE IN VITREOUS BODY)**

1. Congenital (developmental)-may be associated with hyaloid remnants
2. Acquired
   A. Infectious cyst
      (1) Coenurosis (*Coenurus cerebralis* larva of dog tapeworm)
      (2) Luetic retinochoroiditis
(3) Toxoplasmosis
B. Myopia
   C. Parasitic cysts
      (1) Cysticercosis-rare
      (2) Echinococcosis
      (3) Hydatid disease (echinococcosis)-rare, children and young adults, tropical
      (4) Nematode cyst (toxocariasis)
D. Pigmentary retinopathy
E. Retinal detachment
F. Trauma


VITREOUS LIQUEFACTION

1. Myopia
2. Peripheral uveitis
3. Retinitis pigmentosa
4. Spontaneous
5. Trauma
6. With aging
7. With vitreous traction such as Wagner disease


Anatomic classification of macular diseases  435
Bilateral macular lesions  439
Pseudomacular edema  439
Macular edema  439
Absence of foveal reflex  442
Macular pucker  442
Macular exudates and hemorrhages  443
Macular star or stellate retinopathy  443
  Diagnostic table  444
Retinociliary vein  446
Cherry-red spot in macula  446
Macular hemorrhage  447
  Diagnostic table  448
Parafoveal telangiectasia  449
Microhemorrhagic maculopathy  449
Macular cyst  449
Macular hole  449
Macular coloboma  450
Elevated macular lesion  451
Heterotopia of the macula  451
White or yellow flat macular lesion and pigmentary change  451
Pigmentary changes in macula  452
Bull's-eye macular lesion  454
Macular wisps and foveolar splinter  454
Macular hypoplasia  454
Premacular subhyaloid hemorrhage  455
Retinal vascular tortuosity  455
Venous beading  456
Ophthalmodynamometry  456
Pulsation of retinal arteriole  457
Retinal artery occlusion  457
  Diagnostic table  460
Localized arterial narrowing  462
Generalized arterial narrowing  462
Periarteritis retinalis segmentalis  464
Frosted branch angiitis  464
Sheathing of retinal veins  464
Absent venous pulsations  465
Dilated retinal veins  466
Tortuosity of retinal veins and hypoplasia of optic nerves  468
Central retinal vein occlusion  468
Talc retinopathy 523
Crystalline retinopathy 523
Pulfrich stereo-illusion phenomenon 523
Parafoveal telangiectasia 523
Hereditary pediatric retinal degenerations 524
Reticular pattern of dark lines in fundus 524
Retinal pigment epithelial tears 524
Retinal pigment epithelial folds 525
Mizuo phenomenon 525
White-dot fovea 525

ANATOMIC CLASSIFICATION OF MACULAR DISEASES

1. Vitreoretinal surface
   A. Preretinal hemorrhage and subinternal limiting membrane hemorrhage
   B. Vitreous traction on the macula
   C. Epiretinal membrane and macular pucker
2. Nerve fiber-ganglion cell layers
   A. Hereditary cerebromacular degeneration
      (1) Sphingolipidoses
         a. Tay-Sachs disease (GM$_2$-gangliosidosis type I)
         b. Sandhoff disease (GM MDSD$_2$-gangliosidosis type II)
         c. Niemann-Pick disease type A (essential lipid histiocytosis)
         d. Niemann-Pick disease type B (sea-blue histocyte syndrome)
         e. Lactoside ceramidosis
         f. Metachromatic leukodystrophy (arylsulfatase A deficiency)
         g. Gaucher disease (glucocerebroside storage disease)
         h. Farber lipogranulomatosis
         i. Generalized gangliosidosis (GM$_1$-gangliosidosis type I)
         j. Mucolipidosis I (lipomucopolysaccharidosis)
      (2) Goldberg disease (unclassified syndrome with features of mucopolysaccharidoses, sphingolipidoses, and mucolipidoses)
      (3) Ceroid lipofuscinosis
         a. Hagberg-Santevuori (infantile)
         b. Jansky-Bielschowsky disease (late infantile)
         c. Spielmeyer-Vogt Batten
   B. Vitreoretinal dystrophies
      (1) Macular degeneration in congenital hereditary x-linked retinoschisis
      (2) Goldmann-Favre syndrome (vitreotapetoretinal degeneration)-recessive
3. Nerve fiber, ganglion cell, inner plexiform, inner nuclear, outer plexiform layers
   A. Ischemia secondary to inadequate perfusion of retinal vessels
      (1) Branch artery occlusion
      (2) Branch vein occlusion
      (3) Diabetes mellitus
4. Outer plexiform layer
A. Cystoid macular degeneration (see p. 439)
   (1) With retinal vascular leakage
      a. Acute nongranulomatous iridocyclitis
      b. Acute cyclitis
      c. Hypertension
      d. Medication (epinephrine, nicotinic acid)
      e. Neoproliferative diabetic retinopathy
      f. Pars planitis
      g. Postoperative (Irvine-Gass syndrome)
      h. Radiation retinopathy
      i. Retinitis pigmentosa
      j. Sarcoidosis
      k. Vascular anomalies
   (2) Without obvious retinal vascular leakage
      a. Vitreous traction on the macula
      b. Serous detachment of sensory epithelium
      c. Serous detachment of pigment epithelium
      d. Hemorrhagic detachment of macula
      e. Choroidal tumors

B. Lipid deposits in macula secondary to vascular disease in retina
   (1) Stellate retinopathy (see p. 443)
      a. Hypertensive retinopathy
      b. Diabetic retinopathy
      c. Coats disease (retinal telangiectasia)
      d. Trauma-ocular or cerebral
      e. Retinal artery or vein occlusion (see p. 457 and 468)
      f. Retinal periphlebitis
      g. Juxtapapillary choroiditis
      h. Papilledema (see p. 593-601)
      i. Angiomatosis retinae
      j. Papillitis (see p. 578-585, 587-588)
      k. Acute, febrile illness, such as measles, influenza, meningitis,
         erysipelas, psittacosis, Behçet disease
      l. Chronic infections, such as tuberculosis or syphilis
      m. Coccidioidomycosis
      n. Parasitic infection, such as that due to teniae, Giardia,
         Ancylostoma
      o. Idiopathic
   (2) Circinate retinopathy
      a. Senile vascular disease
      b. Venous obstruction
      c. Diabetic retinopathy
      d. Coats disease (retinal telangiectasia)
      e. Retinal detachment
      f. Anemia
      g. Leukemia
h. Idiopathic (primary)
   i. Retinal arterial macro aneurysm

(3) Diabetic retinopathy

5. Outer nuclear layer or photoreceptor elements
   A. Congenital hereditary vision defects
      (1) Trichromatism (anomalous)
      (2) Dichromatism
      (3) Monochromatism
   B. Hereditary macular dystrophies
      (1) Progressive cone dystrophy
      (2) Inverse (macular) retinitis pigmentosa
   C. Olivopontocerebellar degeneration
   D. Light toxicity
      (1) Operating microscope bum
      (2) Solar bum

6. Pigment epithelium
   A. Hereditary macular dystrophies
      (1) Vitelliform dystrophy (Best disease)
      (2) Adult onset foveomacular vitelliform dystrophy (adult Best)
      (3) Fundus flavimaculatus
      (4) Fundus flavimaculatus with macular involvement (Stargardt disease)
      (5) Dominant drusen (Doyne honeycomb dystrophy)
      (6) Reticular pigment dystrophy (Sjögren)
      (7) Butterfly-shaped pigment dystrophy (Deutman)
      (8) Central areolar choroidal and pigment epithelial dystrophy
      (9) Sorsby fundus dystrophy
   B. Inflammatory lesions
      (1) Rubella syndrome (German measles)
      (2) Acute posterior multifocal placoid pigment epitheliopathy
   C. Toxic lesions
      (1) Chloroquine
      (2) Hydroxychloroquine
      (3) Phenothiazine
         a. Chlorpromazine
         b. Thoridazine
      (4) Sparsomycin
      (5) Ethambutol
      (6) Indomethacin
      (7) Quinine
      (8) Desferrioxamine
      (9) Penicillamine
   D. Drusen (senile, degenerative)
   E. Refsum syndrome (phytanic acid storage disease)
   F. Myotonic dystrophy syndrome

7. Bruch membrane
   A. Angioid streaks associated with
(1) Pseudoxanthoma elasticum (Grönblad-Strandberg syndrome)
(2) Senile elastosis of skin
(3) Osteitis deformans (Paget disease)
(4) Fibrodysplasia hyperelastica (Ehler-Danlos syndrome)
(5) Sickle cell anemia
(6) Acromegaly
(7) Beta-thalassemia
(8) Abetalipoproteinemia (Bassen-Kornzweig syndrome)

B. Lacquer cracks in pathologic myopia
C. Traumatic fracture of Bruch membrane

8. Pigment epithelium-Bruch membrane choriocapillaris
   A. Degenerative lesions
      (1) Disciform macular degeneration (senile, juvenile)
      (2) Age-related macular degeneration
      (3) Adult hereditary cerebromacular degeneration (Kufs?)
   B. Serous detachment of neuroepithelium or pigment epithelium associated with
      the following:
      (1) Central serous chorioretinopathy
      (2) Hemangioma of choroid
      (3) Malignant melanoma
      (4) Pit of the optic disk
      (5) Hypotony (see p. 325-326)
      (6) Leukemic infiltrates of choroid
      (7) Terminal illness
      (8) Trauma
      (9) Uveitis
      (10) Optic neuritis (see p. 578-585, 587-588)
      (11) Papilledema (see p. 593-601)
      (12) Acute hypertension
      (13) Vitreous traction
      (14) Angioid streaks (see p. 526-529)
      (15) Vogt-Koyanagi-Harada syndrome
      (16) Toxocara canis
      (17) Myopic choroidal degeneration
      (18) Metastatic carcinoma
      (19) Choroidal nevus
      (20) Collagen vascular disease
      (21) Hemorrhagic or organized disciform detachment

9. Choroid
   A. Degenerative lesions
      (1) Central areolar choroidal atrophy
      (2) Myopic choroidal atrophy
      (3) Helicoid peripapillary chorioretinal atrophy (?)
   B. Inflammatory lesions
      (1) Histoplasmosis
   C. Vascular occlusive lesions
10. Miscellaneous
   A. Retinal inflammations (multilayer alterations that may involve the macula)
      (1) *Toxoplasma gondii*
      (2) *T. canis*
      (3) Septic emboli
      (4) Cytomegalovirus retinitis
      (5) *Candida* organisms
      (6) Bacteria
   B. Congenital anomalies of the macula
      (1) Aplasia
      (2) Hypoplasia
      (3) Heterotopia
      (4) Colobomas (see p. 450)
      (5) Aberrant macular vessels


**BILATERAL MACULAR LESIONS**

1. Development defects (colobomas)
2. Drugs, including the following:
   - allopurinol (?)
   - amodiaquine
   - broxyquinoline
   - chloroquine
   - chlorpromazine
   - clonidine (?)
   - griseofulvin (?)
   - hydroxychloroquine
   - ibuprofen (?)
   - iodochlorhydroxyquin
   - iodoquinol
   - quinine
   - thioridazine
3. Infectious entities
   - A. Herpes simplex
   - *B. Cytomegalic retinitis*
   - C. Candidiasis and nocardiosis
   - D. *T. canis* (visceral larva migrans syndrome)
   - E. Congenital syphilis
   - *F. Tuberculosis*
   - *G. Ocular histoplasmosis*
   - H. Congenital toxoplasmosis
4. Intrauterine inflammations
5. Noninfectious entities
   - A. Best disease
   - B. Stargardt disease
   - *C. Exudative age-related macular degeneration*
6. Presumed inflammatory origin
PSEUDOMACULAR EDEMA

1. Exudative senile maculopathy-serous or hemorrhagic detachment of the macular retina in persons 50 years of age or older, including "giant cyst of macula"
2. Serous detachment of retinal pigment epithelium
3. Central serous retinopathy caused by drugs, including the following:
   - adrenal cortex injection
   - aldosterone
   - betamethasone
   - cortisone
   - desoxycorticosterone
   - dexamethasone
   - fludrocortisone
   - hydrocortisone
   - indomethacin
   - methylprednisolone
   - oral contraceptives

MACULAR EDEMA

This condition involves the loss of foveal depression with ophthalmoscope and outline of multiple cystoid. Spaces can be retroilluminated with slit lamp; often a yellow exudate lies deep within or beneath retina in foveal area.

1. Acquired parafoveal telangiectasis
2. Amebiasis (amebic dysentery)
3. Bang disease (brucellosis)
4. Behçet syndrome (dermatostomatopathmalic syndrome)
5. Carotid artery obstruction
6. Central angiospastic retinopathy
7. Coats disease
8. Choroidal tumors
9. Crohn disease
10. Cytomegalovirus retinitis
11. Diabetic retinopathy
12. Dominant inheritance macular dystrophy
13. Drugs, including the following:
   - acetazolamide
   - acetophenazine
   - adrenal cortex injection
   - aldosterone
   - allopurinol (?)
   - aluminum nicotinate
   - amithiozone
   - amodiaquine
   - aspirin
14. Electrical injuries to the retina
15. Epikeratophakia complication
16. Fabry disease (ceramide trihexoside lipidosis)
17. Felty syndrome (rheumatoid arthritis with hypersplenism)
18. Following corneal-relaxing incisions
19. Goldmann-Favre disease (hyaloideoretinal degeneration)
20. Gyrate atrophy
21. Hallermann-Streiff syndrome (dyscephalic mandibulooculofacial syndrome)
22. Hemangiomas of choroid
23. Hemangioma of choroid
24. Hunter syndrome (mucopolysaccharidoses II, or MPS II)
25. Hurler syndrome (MPS I-H)
26. Hypertensive retinopathy
27. Hypotony (postoperative)
*28. Irvine-Gass syndrome
29. Large central foveal cyst
30. Leukemia
31. Meningococcemia (*Neisseria meningitidis*)
32. Macular dystrophy-dominant
33. Nematode, intraretinal
34. Nylon suture toxicity
35. Optic nerve pit
36. Pars planitis (peripheral uveitis)
37. Photocoagulation
38. Porphyria cutanea tarda
39. Posterior capsule rupture
*40. Preretinal fibrosis (macular pucker)
41. Punctata albescens retinopathy
42. Radiation retinopathy
43. Retinitis pigmentosa
44. Retinohypophysary syndrome (Lijo-Pavia-Lis syndrome)
45. Scheie syndrome
46. Scleral buckle
47. Silverman syndrome (battered baby syndrome)
48. Subacute sclerosing panencephalitis (Dawson disease)
49. Toxoplasmonic chorioretinitis
50. Trauma to globe (commotio retinae)
51. Ultraviolet light from sun, operating microscope, or other bright light sources
52. Uveitis (anterior or posterior)
*53. Vein occlusion, including branch vein occlusion (see p. 468)
54. von Hippel-Lindau syndrome (retinocerebral angiomatosis)
55. Yttrium-aluminum-garnet (YAG) laser posterior capsulotomy


**ABSENCE OF FOVEAL REFLEX**

The absence of foveal reflex is caused by drugs, including amodiaquine, chloroquine, diiodohydroxyquin, hydroxychloroquine, iodochlorhydroxyquin, or quinine.


**MACULAR PUCKER**

Macular pucker involves tiny folds that often are arranged in a stellate manner around macula and usually are associated with a preretinal membrane (preretinal macular fibrosis, preretinal vitreous membrane, surface wrinkling retinopathy, cellophane maculopathy).

1. Associated with proliferative retinopathy  
   *A. Diabetes retinopathy  
   B. Eales disease  
   *C. Hypertension  
   D. Sickle cell disease  
   E. Vein occlusion*
2. Central serous chorioretinopathy  
   A. Psychopharmacologic medication use  
   B. Corticosteroid use  
   C. Hypertension
3. Congenital
4. Following photocoagulation or cryoretinopexy
5. Following traumatic posterior vitreous separation, such as blunt trauma to the eye and whiplash injury (craniocervical syndrome)
6. Loss of formed vitreous at operation
7. Idiopathic (probably related to spontaneous posterior vitreous detachment)
8. Macular detachment
9. Multiple retinal operations
10. Penetrating or blunt injuries
11. Posterior uveitis
* 12. Proliferative vitreoretinopathy following vitreoretinal surgery
13. Retinal detachment
14. Trauma (blunt)
15. Vitreous hemorrhage


**MACULAR EXUDATES AND HEMORRHAGES**

1. Retinal macroaneurysms
2. Hemorrhagic age-related macular degeneration (ARMD)
3. Diabetic macular edema
4. Retinal telangiectasis
5. With dense subretinal and subretinal pigment epithelium (sub-RPE [retinal pigment epithelium]) hemorrhage simulate the appearance of retinal macroaneurysms.


**MACULAR STAR OR STELLATE RETINOPATHY (EXUDATES IN A STAR FORMATION RADIATING AROUND MACULA IN THE NERVE FIBER LAYER)**

1. Acute febrile illness, such as measles, influenza, meningitis, erysipelas, psittacosis, Behçet disease (dermatostomatopathalmic syndrome)
2. Cat-scratch fever
3. Chronic infections, such as tuberculosis or syphilis
4. Coccidioidomycosis
5. Gansslen syndrome (familial hemolytic icterus)
*6. Hypertension
7. Idiopathic
8. Juxtapapillary choroiditis (Jensen disease)
*9. Neuroretinitis
10. Obstruction of the artery or vein supplying the macular area (see p. 457 and 468)
11. Ocular or cerebral trauma
12. Parasitic infection, such as that due to teniae, *Giardia, Ancylostoma*
*13. Papilledema (see p. 563-601)
15. Retinal periphlebitis


**Extracted Table Steellate Retinopathy**

**RETINOCILIARY VEIN-DISAPPEARS FROM THE RETINA AT DISC MARGIN WITHOUT CONNECTION TO CENTRAL RETINAL VEIN**

1. Acquired  
   A. Arachnoid cyst of the optic nerve  
   B. Central retinal vein occlusion (see p. 468)  
   C. Chronic atrophic papilledema from causes including craniopharyngioma (see p. 595-597)  
   D. Glioma of the optic disc

2. Congenital


**CHERRY-RED SPOT IN MACULA (RULE OUT MACULAR HEMORRHAGE)**

1. Cardiac myxomas  
2. Cryoglobulinemia  
3. Dapsone poisoning  
4. Hallervorden-Spatz disease (pigmentary degeneration of globus pallidus)  
5. Hollenhorst syndrome (chorioretinal infarction syndrome)  
6. Hurler syndrome (MPS I-H)  
*7. Hypertension (severe)  
8. Intralesional chalazion corticosteroid injection  
9. Leber congenital amaurosis  
10. Macular hemorrhage  
*11. Macular hole with surrounding retinal detachment  
12. ML I (lipomucopolysaccharidosis)  
13. Myotonic dystrophy syndrome (Curschmann-Steinert syndrome)  
14. Multiple sulfatase deficiency  
*15. Occlusion of central retinal artery (see p. 457)  
16. Quinine toxicity  
17. Sphingolipidoses  
   A. Cherry-red spot myoclonus
B. Farber syndrome (Farber lipogranulomatosis)
C. Gangliosidosis GM₁-type (juvenile gangliosidosis)
D. Gaucher disease (glucocerebroside storage disease)
E. Goldberg syndrome
F. Infantile metachromatic leukodystrophy (van Bogaert-Nyssen disease)
G. Niemann-Pick disease type A
H. Niemann-Pick disease type B
I. Sandhoff disease (gangliosidosis GM₂-type 2)
*J. Tay-Sachs disease (gangliosidosis GM-type 1)

18. Steroid injection intranasally
*19. Temporal arteritis (giant cell arteritis)
20. Traumatic retinal edema (commotio retinae; Berlin edema)
21. Vogt-Spielmeyer cerebral degeneration (Batten-Mayou syndrome)


**MACULAR HEMORRHAGE**

1. Choroidal neovascular membranes
   *A. Age-related macular degeneration
   B. Angioid streaks
   C. Histoplasmosis
   D. Idiopathic
   E. Pathologic myopic
   F. Posterior uveitis
2. Infectious retinitis
   *A. Cytomegalovirus retinitis
   B. Subacute bacterial endocarditis
3. Retinal vascular disease
   A. Radiation retinopathy
   B. Retinal arterial macro aneurysm
   *C. Vein occlusion
4. Systemic diseases
   A. Blood dyscrasias
      (1) Anemia
(2) Leukemia
(3) Polycythemia vera
(4) Sickle cell disease
(5) Thrombocytopenia
(6) Waldenström macroglobulinemia

B. Cardiovascular shock (especially gastrointestinal hemorrhage)
* C. Diabetes mellitus
* D. Human immunodeficiency virus (HIV)-related retinopathy
* E. Hypertension

F. Toxemia of pregnancy

5. Trauma
   A. Choroidal rupture
   B. Purtscher retinopathy
   C. Shaken-baby syndrome
   D. Terson syndrome
   E. Valsalva retinopathy
   F. Vitreous detachment


**Extracted Table Macular Hemorrhage**

**PARAFOVEAL TELANGIECTASIA**

This condition is a retinal microvascular anomaly involving the parafoveal capillary network as well as the immediately adjacent vascular bed and is best demonstrated by fluorescein angiography.

1. Carotid artery obstruction
* 2. Diabetes mellitus, usually bilateral
* 3. Idiopathic
4. Localized form of Coats disease, usually unilateral
* 5. Small branch vein occlusion
6. Small retinal capillary hemangioma, usually unilateral
7. Roentgenogram, irradiation


MICROHEMORRHAGIC MACULOPATHY - SMALL MONOCULAR MACULAR HEMORRHAGE THAT IS PUNCTATE, ROUND OR BILOBED

1. Increased venous stasis (Valsalva stress)
2. Impaired blood platelet aggregation
3. Medications that impair platelet function including aspirin, ibuprofen (Motrin), pentazocine, propranolol hydrochloride and oral contraceptives.


MACULAR CYST

Macular cyst must be differentiated from macular hole with Hruby lens or contact lens and slit lamp.

1. Amebiasis
2. Cysticercosis-subretinal cyst
3. Cystic degeneration-common following trauma, uveitis, and vascular disease
4. Hamman-Rich syndrome (alveolar capillary block syndrome)
5. Histoplasmosis
6. Hydatid disease (echinococcosis)
7. Parasitic and mycotic cysts


MACULAR HOLE

Macular hole must be differentiated from macular cyst with Hruby lens or contact lens and slit lamp)

* 1. Idiopathic (most common, may be bilateral)
2. From the following:
   A. Edema (see p. 439)
      (1) Inflammatory
      (2) Toxic
      (3) Vascular
      (4) Following papilledema
   B. High myopia
   C. Ischemic, such as with retinal detachment or choroidal tumor-the macula is separated from choriocapillaris
   D. Degenerative conditions of the retina and retinal dystrophy
   E. Trauma
   F. Radiation injury
G. Glaucoma
H. Posterior senile retinoschisis
I. High tension electric shock
J. Central serous chorioretinopathy
K. Optic disc coloboma
L. Posterior retinal detachment associated with optic pits
M. Industrial laser burns
N. Lightning-induced
O. Posterior microphthalmos
P. Septic embolization
Q. Subhyaloid hemorrhage
R. Topical pilocarpine use
S. YAG laser

3. Dawson disease (subacute sclerosing panencephalitis)
4. Foveomacular retinitis-usually young males
*5. Pseudohole due to epiretinal membrane (may differentiated from true hole by fluorescein angiography)
6. Sickle cell disease


**MACULAR COLOBOMA**

This condition involves bilaterally symmetric, circumscribed, excavated defects in choroid and retina in the region of macula associated with reduced vision

1. Autosomal dominant
2. Autosomal recessive inheritance with skeletal anomalies
3. Conditions that exhibit choroidal coloboma (see p. 555-556)
4. Down syndrome
5. Hypercalciuria, myopia, and macular coloboma
6. Isolated
7. Macular coloboma with brachydactyly
8. Sorsby syndrome I


**ELEVATED MACULAR LESION**

1. Angiospastic retinopathy
2. Central serous detachment of retina
3. Chorioretinitis especially histoplasmosis and toxoplasmosis
4. Choroidal hemangioma
5. Dawson disease (subacute sclerosing panencephalitis)
6. Malignant melanoma
7. Varix of the vortex ampulla
8. Sickle cell disease


**HETEROTOPIA OF THE MACULA**

This condition involves an abnormal location of the macula in relation to the optic disc; the eye with the ectopic macula tends to deviate in the same direction as macular displacement; visual fields show displacement of blind spot and cover-uncover test shows no shift of fixation.

1. Chorioretinitis
2. Congenital
   *3. Retinopathy of prematurity
4. Inflammatory
   *5. Proliferative diabetic retinopathy


## WHITE OR YELLOW FLAT MACULAR LESION AND PIGMENTARY CHANGE

1. Posttraumatic - pigmentary disturbance; cysts or hole at macula
2. Postinflammatory - chorioretinal atrophy with pigment clumping at center and periphery of lesion
3. Coloboma of macula-atrophic area at macula often associated with coloboma of disc; sclera may be ectatic (see p. 450)
4. Radiation injuries-common after solar eclipse; punched-out appearance
5. Fuchs dark spot-pigmented spot associated with other signs of degenerative myopia
6. Drugs, including the following:
   - adrenal cortex injection
   - aldosterone
   - allopurinol (?)
   - amodiaquine
   - betamethasone (?)
   - chloroquine
   - cortisone (?)
   - desoxycorticosterone (?)
   - dexamethasone (?)
   - diiodohydroxyquin
   - fludrocortisone
   - fluprednisolone (?)
   - griseofulvin
   - hydrocortisone (?)
   - hydroxychloroquine
   - indomethacin (?)
   - iodochlorhydroxyquin
   - methylprednisolone
   - oral contraceptives
   - paramethasone (?)
   - prednisolone (?)
   - prednisone (?)
   - quinine
   - triamcinolone
7. Stellate retinopathy - star-shaped exudates (see p. 443)
8. Hard exudates and circinate retinopathy (see p. 495)
9. Drusen – common, discrete yellow spots beneath the retina
10. Doyne honeycomb choroiditis-rare; honeycomb pattern of yellow patches at posterior pole; degenerative changes at macula
11. Heredomacular dystrophies
   - A. Best disease (vitelliruptive macular dystrophy) up to 18 years of age; egg-yolk lesion at macula, later absorbed to leave atrophic scar
   - B. Fundus flavimaculatus - yellow patches at posterior pole; degenerative changes at macula
   - C. Stargardt disease (juvenile macular degeneration) to 10 years of age; variable appearance in different families; bilateral lesions showing some degree of symmetry
   - D. Behr disease (optic atrophy-ataxia syndrome) – adults, similar to Stargardt type
   - E. Presenile and senile-pigmentary changes followed by atrophy, bilateral and symmetric
13. Central choroidal sclerosis - rare, atrophic retina with sclerosed choroidal vessels showing clearly
14. Central areolar choroidal atrophy-rare, exude and edema followed by sharply defined atrophic area with white strands of choroidal vessels
15. Pseudoinflammatory macular dystrophy-rare, initially edema and exudates followed by scarring with pigmentary disturbance and atrophic patches
16. Gaucher disease (glucocerebroside storage disease)-rare, ring-shaped macular lesions, lipid deposits in cornea and conjunctiva
17. Diffuse leukoencephalopathy - rare, white deposits in periphery and macular area
18. Sjögren-Larsson syndrome (oligophrenia-ichthyosis-spastic diplegia syndrome)
19. Angioid streaks (see p. 526-529)
20. Multiple evanescent white-dot syndrome (MEWDS) usually unilateral, predominantly healthy women, vitreitis
21. Acute multifocal placoid pigment epitheliopathy – rare, map-like pigmented disturbance of posterior pole or more widespread over posterior fundus


**Pigmentary Changes in Macula**

1. Hereditary macular degeneration without cerebral or other disease
   A. Best disease (vitelliform macular dystrophy)
   B. Stargardt disease (juvenile flavimaculatus)
   C. Behr syndrome (optic atrophy-ataxia syndrome)
*2. Retinitis pigmentosa*
3. Secondary pigmentary retinopathy following trauma or inflammation (see p. 497)
*4. Age-related macular degeneration*
5. Metabolic disease associated with pigmentary retinopathy
   A. Abetalipoproteinemia (Bassen-Kornzweig syndrome)
   B. Alpha- lipoprotein deficiency (Tangier syndrome)
   C. Cereoid lipofuscinosis
      (1) Batten-Mayou syndrome
      (2) Dollinger-Bielschowsky syndrome, late infantile (Bielschowsky-Jansky disease)
      (3) Infantile type of neuronal ceroid lipofuscinosis
   D. Hepatic disease
   E. Refsum disease (phytanic acid storage disease)
   F. Tay-Sachs syndrome (gangliosidosis GM2-type I)
   G. Vitamin A
   H. MPS
      (1) Hunter syndrome (MPS II)
      (2) Hurler syndrome (MPS I-H)
      (3) Sanfilippo-Good syndrome (MPS III)
      (4) Scheie syndrome (MPS I-S)
*6. Drugs, including the following:
   acetophenazine  amodiaquine  benzatropine (?)
   amiodarone (?)  azathioprine  biperiden (?)
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<td>7. Inflammation</td>
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<td>8. Multifocal necrotizing encephalopathy</td>
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<td>9. Dawson disease (subacute sclerosing panencephalitis)</td>
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**BULL'S-EYE MACULAR LESION-CIRCULAR AREA OF RETINAL PIGMENT EPITHELIUM ATROPHY SURROUNDING A SPARED FOVEA**
1. Autosomal-dominant, benign, concentric annular macular dystrophy
2. Ceroid lipofuscinosis
*3. Chloroquine or hydroxychloroquine retinopathy
4. Cone dystrophy
5. Hereditary ataxia
6. NARP syndrome
7. Spielmeyer-Vogt syndrome (Batten-Mayou syndrome)
*8. Stargardt disease (or fundus flavimaculatus)
9. Trauma
10. Unknown


**MACULAR WISPS AND FOVEOLAR SPLINTER**

These are noted in focal illumination with Goldmann contact lenses but are invisible ophthalmoscopically.

1. Direct and indirect ocular concussion
2. Following absorption of small prefoveal hemorrhage
3. Foveomacular retinitis
4. Juvenile macular degeneration
5. Old, healed chorioretinitis
6. Retinitis pigmentosa
7. Spontaneous senile posterior vitreous detachment
8. Whiplash injury


**MACULAR HYPOPLASIA (INCOMPLETE MACULAR DEVELOPMENT MANIFESTED BY DECREASED VISION)**

* 1. Albinism
*2. Associated with autosomal-dominant aniridia
3. Associated with microcornea and corectopia
4. Associated with myelinated nerve fibers
5. Forsius-Eriksson syndrome (Aland disease)
*6. Goldenhar-Gorlin syndrome (oculoauriculovertebral dysplasia)
7. Krause syndrome (encephaloophthalmic dysplasia)
8. Ring chromosome
9. Syndrome of foveal hypoplasia and presenile cataract (O'Donnell-Pappas syndrome) - autosomal dominant
10. Tuomaala-Haapanen syndrome
11. Waardenburg syndrome (interoculoriridodermatoauditve dysplasia)


PREMACULAR SUB HYALOID HEMORRHAGE

1. Branch retinal vein occlusion
2. Blood dyscrasia
3. Diabetic retinopathy
4. Retinal macroaneurysm
5. Terson syndrome
6. Valsalva retinopathy


RETINAL VASCULAR TORTUOSITY

1. Acute malnutrition
2. Aortic coarctation
3. Bazzana syndrome (angiospastic ophthalmoauricular syndrome)
4. Choked disc (see p. 593)
5. Chronic respiratory insufficiency, such as in cystic fibrosis and familial dysautonomia (Riley-Day syndrome)
*6. Coats disease (retinal telangiectasia)
7. Congenital
8. Cri-du-chat syndrome (cat-cry syndrome)
9. Cryoglobulinemia
10. Down syndrome (trisomy 21)
11. Eales disease (periphlebitis)
12. Engelmann syndrome (diaphyseal dysplasia)
13. Fabry disease (diffuse angiokeratosis)
*14. Glaucoma, open angle
15. Granulocytic sarcoma of orbit
16. Hereditary hemorrhagic telangiectasia (Osler disease)-tortuosity and varicosity
17. Hypertension
18. Kenny syndrome (dwarfism, thickened long bone cortex, transient hypocalcemia)
19. Leukemia
20. Lymphogranuloma venereum (Nicolas-Favre disease)
21. Maroteaux-Lamy syndrome (mucopolysaccharidoses type VI)
22. Macroglobulinemia
23. Mosse syndrome (polycythemia-hepatic cirrhosis syndrome)
24. Myopia
25. Normal variation with fullness
26. Polycythemia with vessel fullness
27. Retinopathy of prematurity
28. Racemose hemangioma of retina, angiomatosis retinae without obvious tumor formation, or von Hippel-Lindau syndrome (retinocerebral angiomatosis)
29. Reimann syndrome (hyperviscosity syndrome)
*30. Sickle cell disease
31. Visceral larva migrans (nematode ophthalmia syndrome)


**VENOUS BEADING**

*1. Diabetes mellitus
2. Loaiasis (*Loa loa*)
3. Macroglobulinemia (Waldenstrom syndrome)


**OPHTHALMODYNAMOMETRY**

When blood pressure of the retinal artery is measured, a difference between the eyes of about 15% of diastolic readings is considered significant.

1. False-positive or variable readings
   A. Abnormally high or low intraocular pressure or asymmetry between the two eyes
B. Cardiac abnormalities, such as atrial fibrillation, heart block, or extrasystoles
C. Marked asymmetry of retinal vessels in the two eyes
D. Measurements of ophthalmic artery pressure lower than 20 g on the instrument
E. Poor patient cooperation
F. Variation in systemic blood pressure between readings

2. High ophthalmodynamometry values
   A. Basilar-vertebral occlusion
   B. Bilateral distal internal carotid occlusion-unusual
   C. Progressive intracranial arterial occlusion syndrome

3. Low ophthalmodynamometry values
   A. Both sides reduced with orthostatic hypotension
   *B. Reduced on side of an occluded internal carotid artery


**PULSATION OF RETINAL ARTERIOLE (HIGH PULSE PRESSURE)**

1. Aortic regurgitation
2. Hyperthyroidism
   *3. Intraocular blood pressure higher than diastolic blood pressure but lower than systolic blood pressure


**RETINAL ARTERY OCCLUSION**

This condition involves a sudden, painless visual loss; on ophthalmoscopic examination, a diffuse retinal pallor and a cherry-red spot in macula are noted.

1. Embolism-cardiac or pulmonary sources
   A. Air emboli following trauma or surgery
   B. Amniotic fluid embolization
   C. Cardiac myxoma
   D. Corticosteroid emboli
   E. Espildora-Luque syndrome (ophthalmic Sylvian syndrome)
   F. Fat emboli following long-bone fractures
   G. Iatrogenic trauma induced by angiography
   *H. In older patients due to atheroma of carotid artery
   I. In young persons due to postrheumatic vegetations (rheumatic fever), cardiac catheterization, or valvotomy
   J. Leuodoemboli - vasculitis, Purtscher retinopathy, septic endocarditis
   K. Moyamoya disease (multiple progressive intracranial arterial occlusion)
   L. Nicolau syndrome (emboli of medication inadvertently introduced into artery)
   M. Synthetic material used in cardiac and vascular procedures
   N. Talc emboli-long-term intravenous drug abusers
O. Tumors - atrial myxoma, mitral valve papillary fibroelastoma
P. With cerebral infarction after periocular subcutaneous cosmetic silicone injection
*2. Atherosclerosis of common carotid artery (ophthalmodynamometry employed for diagnosis)
3. Ischemia
   A. Carotid occlusion or dissection
   B. Essential hypotension
   C. Following orbital floor fractures or repair
   D. Following surgery for retinal detachment
   E. Generalized shock
   F. Heart failure (rare)
   G. Kahler disease (multiple myeloma)
   H. Knee-chest position
   I. Massive hemorrhage, such as that occurring in hematemesis. Gastrointestinal bleeding, or surgical procedures
   J. Migraine
   K. Mosse syndrome (polycythemia-hepatic cirrhosis syndrome)
   L. Orbital hemorrhage following retrobulbar injection
   M. After surgery for scoliosis
   N. Too rapid lowering of blood pressure in hypertensive subjects
4. Inflammation
   A. Abdominal typhus (typhoid fever)
   B. African eye-worm disease (loiasis)
   C. Arteriole vasculitis, such as periarteritis nodosa (Kussmaul disease)
   D. Bacterial endocarditis
   E. Behçet disease (dermatostomatoophthalmic syndrome)
   F. Diphtheria
   G. Familial factor V Leiden polymorphism and positive rheumatoid factor
   H. Giant cell arteritis
   I. Herpes zoster
   J. Metastatic bacterial endophthalmitis
   K. Mucormycosis (phycomycosis)
   L. Pancreatitis
   M. Recurrent toxoplastic retinochoroiditis
   N. Rocky Mountain spotted fever (spotted fever)
   O. Rubeola (measles)
   P. Subacute bacterial endocarditis
   Q. Systemic lupus erythematosus
   R. Takayasu disease (pulseless disease)
   *S. Temporal arteritis
   T. Toxoplasma retinochoroiditis
   U. Varicella (chickenpox)
5. Blood disease
   A. After platelet transfusion
B. Following ocular trauma with secondary glaucoma in youths with sickle-trait hemoglobinopathy
C. Polycythemia vera (Vaquez-Osler syndrome)
D. Sickle cell disease

6. Syphilis (acquired lues)
7. Associated factors
8. Diathermy of persistent hyaloid
   A. Drusen of optic nerve (see p. 559-560)
   B. Giant cell arteritis
   C. Papilledema (see p. 593-601)
   D. Subdural cerebral hemorrhage
   E. Arteriosclerosis of central retinal artery
   F. Chronic simple glaucoma

9. After dye, yellow photocoagulation
10. Complication of retrobulbar block
11. Degos syndrome (malignant atrophic papulosis)
12. Disseminated lupus erythematosus
13. Fabry-Anderson syndrome (glycosphingolipid lipidosis)
14. Goldenhar-Gorlin syndrome (oculoauriculovertebral dysplasia)
15. Homocystinuria syndrome
16. Hyperhomocystinemia
17. Lyme disease
18. Neoplastic angioendotheliomatosis
19. Polymyalgia rheumatica
20. Protein S deficiency
21. Relapsing polychondritis
22. Sneddon syndrome (livedo reticularis, neurologic abnormalities, and labile hypertension)
23. Use of tranexamic acid therapy


Extracted Table Retinal artery occlusion

LOCALIZED ARTERIAL NARROWING

1. Retinal atrophy following:
   A. Degeneration
   B. Inflammation
   C. Trauma
   D. Treatment with diathermy, light, or cryopexy
2. Any vascular retinopathy


GENERALIZED ARTERIAL NARROWING

1. Local causes
   A. Apparent narrowing
      (1) High hypermetropia-common, small disc, narrow vessels, sometimes pseudopapilledema (see p. 601)
      (2) Congenital microphthalmos - rare, hypermetropia, often cataract (see p. 252-253)
      (3) Aphakia - cataract operation, dislocated lens (see p. 405)
      (4) Hollenhorst syndrome (chorioretinal infarction syndrome)
      (5) Wagner syndrome (hyaloideoretinal degeneration)
   B. Trauma
      (1) Avulsion of optic nerve-rare, secondary optic atrophy
      (2) Fracture involving bony optic canal-rare, secondary optic atrophy
      (3) Following retroocular injection-rare secondary optic atrophy
      (4) Orbital hemorrhage following retroocular injection or orbital operation- rare, secondary optic atrophy
      (5) Carotid ligation for carotid-cavernous fistula, rare, secondary optic atrophy
      (6) Following angiography-rare, secondary optic atrophy
      (7) Siderosis bulbi-metallic intraocular foreign body
   C. Infection and edema
      (1) Orbital cellulitis - exophthalmos, restricted ocular movements
      (2) Following thyrotropic exophthalmos-ocular muscle paresis, lid retraction
D. Degenerations, such as progressive cone-rod degeneration
E. Primary tapetoretinal degenerations, such as retinitis pigmentosa; Hallgren syndrome (retinitis pigmentosa-deafness-ataxia syndrome)

2. Systemic disease

A. Arteriosclerosis
   (1) In involutionary sclerosis-population older than 50 years of age, generalized arteriolar narrowing, diminished light reflexes
   (2) In arteriosclerotic disease
      a. Arteriosclerotic central artery occlusion-common arteriovenous crossing signs, focal arteriolar constriction
      b. Embolus from atheromatous plaque, common, sudden onset, visible white embolus

B. Hypertensive conditions
   *(1) Essential hypertension-retinal hemorrhages, cotton-wool spots, arteriovenous crossing signs
   *(2) Malignant hypertension-retinal hemorrhages, cotton-wool spots, edema of disc
   (3) Toxemia of pregnancy-rare, sometimes hemorrhages, cotton-wool spots, edema of disc, serous detachment
   (4) Coarctation of aorta-rare, hypertensive changes vary greatly in degree
   (5) Pheochromocytoma-rare, hypertensive changes vary greatly in degree
   (6) Adrenal tumor, hyperaldosteronism (adrenal medulla tumor syndrome) - rare, hypertensive changes vary greatly in degree
   (7) Cushing tumor (adrenocortical hyperfunction)-rare, hypertensive changes vary greatly in degree
   (8) Motor-neuron disease of cervicothoracic cord hypertension; may occur after prolonged artificial pulmonary ventilation

C. Other forms of vascular disease
   (1) Retinal ischemia -hypotension following severe or recurrent bleeding, unilateral blindness in patients
   *(2) Temporal arteritis (cranial arteritis, giant -cell arteritis)-common, 50 years or older; mean age at onset, 55 years; sudden blindness at onset
   (3) Polyarteritis nodosa (Kussmaul disease)-multiple signs involving choroid, retina, cornea, episclera, and ocular muscles
   (4) Proliferative diabetic retinopathy-arterial narrowing occurs in 17% of patients with proliferative diabetic retinopathy, mainly in cicatricial stage
   (5) Cardiac arrest-thread-like arterioles, segmentation of blood column, generalized retinal pallor, pallor of disc, sometimes macular cherry-red spot
   (6) Raynaud disease (idiopathic paroxysmal digital cyanosis)-young adults, more common in women

D. Renal disease
   (1) Acute glomerulonephritis-preceding illnesses, including scarlet fever, streptococcal tonsillitis, otitis media, erysipelas (St. Anthony fire), subacute bacterial endocarditis, polyarteritis nodosa (Kussmaul disease)
(2) Chronic glomerulonephritis—often asymptomatic and found on routine examination
(3) Pyelonephritis and pyelitis—most common causes of renal failure

E. Diseases of the central nervous system
(1) Migraine
*(2) Syphilitic neuroretinitis
(3) Viral neuroretinitis (rare complication)
(4) Tay-Sachs disease (amaurotic familial idiocy)
(5) Jansky-Bielschowsky disease (amaurotic familial idiocy, late form)
(6) Myotonic dystrophy syndrome (Curschmann-Steinert syndrome)
(7) Retinohypophysis syndrome (Lijo Pavia-Lis syndrome)
(8) Zellweger syndrome (cerebrohepatorenal syndrome of Zellweger)

F. Toxic causes
(1) Chloroquine, hydroxychloroquine, quinacrine, amodiaquine
(2) Lead
(3) Quinine - rare, may follow large dose (abortifacient) or normal dose in sensitive subjects

G. Other causes
(1) Caisson syndrome (bends)
(2) Chédiak-Higashi syndrome (oculocutaneous albinism with recurrent infections; autosomal recessive)
(3) Hunter syndrome (MPS II)
(4) Sanfilippo-Good syndrome (MPS III)


PERIARTERITIS RETINALIS SEGMENTALIS

White or yellow plaques are arranged in segments encircling arteries like a cuff and are localized to one or more arterial branches.

*1. Arteriosclerosis secondary to vein obstruction
2. Herpes zoster
3. Hypercholesterolemia
4. Lupus erythematosus (disseminated lupus erythematosus)
5. Metastatic uveitis
6. Periarteritis nodosa (necrotizing angiitis)
*7. Sarcoidosis syndrome
8. Syphilis (acquired lues)
*9. Temporal arteritis (giant cell arteritis)
* 10. Tuberculous retinitis
11. Uveitis, idiopathic
FROSTED-BRANCH ANGITIS

In this condition, unusual thick sheathing surrounds all the retinal veins and less often the arteries, making them look like frosted tree branches.

1. Herpes simplex viruses types 1 and 2
2. Acute lymphoblastic leukemia
3. Large cell lymphoma
4. Crohn disease
5. Systemic lupus erythematosus
6. Acquired immune deficiency syndrome


SHEATHING OF RETINAL VEINS

In this condition, white or gray envelopes are around veins; retinal or vitreous hemorrhage and exudates may be present.

1. Disc only-developmental
2. Disc and retina-papillitis or papilledema
3. Peripheral sheathing
   *A. Acute retinal necrosis
   B. Amebiasis
   C. Behçet disease (dermatostomatoophthalmic syndrome)
   D. Brucellosis-rare, tortuosity and sheathing of veins, vitreous haze, retinal hemorrhages
   E. Candidiasis
   F. Coccidioidomycosis
   G. Eales disease (periphlebitis)
   H. Diabetes mellitus
   I. Filariasis-hemorrhages and exudates
   J. Hypertension
   K. Infectious mononucleosis-peripheral or central perivascular involvement, venous engorgement and sheathing associated with retinal hemorrhages
   L. Lupus erythematosus
   M. Non-Hodgkin lymphoma
   N. Onchocerciasis syndrome (river blindness)
O. Rickettsial infections - peripheral or central perivascular involvement, venous engorgement and sheathing associated with retinal hemorrhages
*P. Sarcoidosis
Q. Septicemia and bacteremia - rare, venous engorgement, usually with multiple hemorrhages and focal sheathing
R. Sickle cell disease
S. Syphilis (secondary) (acquired lues)
T. Tuberculin or bacille Calmette-Guerin (BCG) vaccination - rare, sectorial, or generalized changes
U. Viral infections, including the following:
    (1) Cytomegalovirus retinitis
    (2) Herpes simplex (likely responsible for acute retinal necrosis)
    (3) Herpes zoster ophthalmicus
    (4) Influenza
    (5) Rift Valley fever
4. Peripheral sheathing without secondary retinopathy - multiple sclerosis
5. Wide and usually dense sheathing of dilated and tortuous veins, suggestive of myelogenous leukemia


**ABSENT VENOUS PULSATIONS (SPONTANEOUS VENOUS PULSATIONS ABSENT AT VENULES ON THE DISC)**

1. Normal individuals
*2. Impending central vein occlusion (see p. 468)
*3. Papilledema (see p. 593-601)


**DILATED RETINAL VEINS**

Normally, the arteriole-venule ratio is 2:3; with an increase in this ratio, the retinal veins may be dilated.

1. Congenital
A. Congenital tortuosity of retinal vessels - rare, sometimes associated with coarctation of aorta

B. Fabry disease (hereditary dystrophic lipidosis)

C. Hemangioma

D. Longfellow-Graether syndrome

E. Ocular fundi in newborns

F. Racemose (arteriovenous) aneurysm - rare, arteriovenous anastomoses localized to sector of retina

*G. Retinopathy of prematurity with plus disease

H. von Hippel-Lindau disease (angiomatosis) - familial in 20% of cases, bilateral in 50%

2. Trauma and inflammation

A. Anterior uveitis - dilatation of veins, often slight hyperemia of disc

B. Carotid-cavernous fistula - fracture of base of skull, progressive exophthalmos, bruit

C. Cavernous sinus thrombosis - rare, proptosis and orbital edema

*D. Impending obstruction of the central retinal vein

E. Periphlebitis - sheathing of vessels

3. Cardiovascular disease - dilatation may be present but rarely dominates the fundus picture

A. Arteriosclerosis

B. Involutionary sclerosis (later stages)

C. Secondary to defective arterial flow, such as in the following:
   (1) Aortic arch syndrome (pulseless disease)
   (2) Cardiac insufficiency
   (3) Congenital heart disease
   (4) Iatrogenic (lowering of blood pressure)
   (5) Severe blood loss
   *(6) Stenosis or occlusion of common carotid
   *(7) Temporal arteritis
   (8) Venous stasis (hypotensive retinopathy of Kearns and Hollenhorst)

D. Heritable thrombophilia and hypofibrinolysis

4. Respiratory disease - venous dilatation may occur with purplish hue of whole retina; obstruction of venous return from the head, such as in the following:

A. Congenital septal defect (Fallot tetralogy)

B. Emphysema

C. Hamman-Rich syndrome (diffuse pulmonary fibrosis syndrome)

D. Heart failure of any type

E. Kartagener syndrome (sinusitis-bronchiectasis-situs inversus syndrome)

F. Mechanical compression of chest

G. Mediastinal tumor obstructing superior vena cava

5. Diseases of the central nervous system

*A. Carotid-cavernous fistula - fractured base of skull; rupture of berry aneurysm, arteriosclerosis

B. Hemangioma of posterior fossa - rare, papilledema, often grossly dilated veins

C. Optic nerve lesion - rare, secondary to orbital space-occupying lesion
D. Papilledema (see p. 593-601)
E. Retrolenticular syndrome (Dejerine-Roussy syndrome)
F. Subarachnoid hemorrhage-head injury; subhyaloid hemorrhages near disc, dilated veins, sometimes papilledema

6. Blood diseases
   A. Aplastic anemia-hemorrhage is the most striking sign, often spreading around the disc
   B. Cryoglobulinemia-rare, may occur with multiple myeloma, veins dilated, tortuous, and sometimes beaded
   C. Gansslen syndrome (familial hemolytic icterus)
   D. Lymphatic leukemia
   E. Macrocytic anemia of all types-common, retinopathy absent unless hemoglobin below 50%; pale fundus, superficial hemorrhages, cotton-wool spots
      (1) Pernicious anemia
      (2) Steatorrhea
      (3) Celiac disease
      (4) Carcinoma of stomach
   F. Macroglobulinemia-rare; veins dilated tortuous and sometimes beaded, hemorrhages and occasionally microaneurysms
   G. Monocytic leukemia
   H. Myelogenous leukemia
   I. Multiple myeloma (Kahler disease)
   J. Polycythemia rubra vera (primary; Vaquez disease)-common in males; hemorrhages; papilledema may be marked and venous thrombosis may occur
   K. Secondary polycythemia-common; hemorrhages, papilledema and venous thrombosis may occur
   L. Sickle cell disease-dilatation of peripheral veins with retinal, subhyaloid, and vitreous hemorrhages
   M. Thrombocytopenic purpura-retinal and subhyaloid hemorrhages near disc, moderate venous dilatation

7. Acute febrile illnesses-rare, occasional dilatation of retinal veins with a few hemorrhages and mild edema of disc
   A. Infectious mononucleosis
   B. Influenza
   C. Rickettsial infections
   D. Septicemia

8. Metabolic diseases
   A. Cystic fibrosis syndrome (fibrocystic disease of pancreas)-venous congestion often swelling of disc
   B. Plasma lecithin
   C. Diabetic retinopathy-larger veins affected, often beaded

9. Collagen diseases
   A. Polyarteritis nodosa-among other fundus lesions, dilated veins may occur
   B. Sclerosis, progressive systemic (scleroderma)
   C. Systemic lupus erythematosus-cotton-wool spots, occasional hemorrhages, and moderate dilatation of veins
10. Toxic conditions, such as methyl alcohol ingestion


**TORTUOSITY OF RETINAL VEINS AND HYPOPLASIA OF OPTIC NERVES**

1. Endocrinopathy, especially pituitary deficiency  
2. Alcohol abuse; fetal alcohol syndrome  
3. Migraine disturbances  
4. Agenesis of the corpus callosum associated with mutations  
5. Preterm birth also associated with retinal artery/visceral tortuosity


**CENTRAL RETINAL VEIN OCCLUSION**  
This condition is characterized by massive hemorrhage into the posterior portion of the eye and dilated retinal veins.

1. External compression of the vein  
   A. Atherosclerosis of central retinal artery  
   B. Connective tissue strand within the floor of the physiologic excavation  
   C. Multiple crossings of the same artery and vein or congenital venous loops or twists in the retinal surface  
   D. Pseudotumor cerebri  
2. Degenerative or inflammatory venous disease, causing detachment, proliferation, and hydrops  
   A. Acquired immunodeficiency syndrome (AIDS; HIV retinopathy)  
   B. Arterial hypertension  
   C. Arteriovenous malformations of retina  
   D. Cardiac decompensation  
   E. Closed-head trauma  
   *F. Diabetes mellitus (Willis disease)  
   G. Ipsilateral internal carotid artery stenosis  
   H. Lyme disease  
   I. Optic disc drusen  
   J. Optic nerve inflammation  
   K. Sarcoidosis  
   L. Serpiginous choroiditis  
   M. Systemic granulomatous disease, particularly tuberculosis
3. Thrombosis from venous stagnation
   A. Spasm of corresponding retinal arterioles
   B. Blood dyscrasias
      (1) Cryoglobulinemia
      (2) Emphysema with secondary erythrocytosis
      (3) Deficiencies in endemic pathway (factor V R506Q mutation)
      (4) Homocystinemia
      (5) Increased platelet aggregation
      (6) Leukemias
      (7) Lymphoma
      *(8) Multiple myeloma
      *(9) Polycythemia vera
      (10) Sickle cell disease
   C. Increased viscosity of the blood
      (1) Cystic fibrosis of pancreas
      (2) Following peritoneal dialysis
      (3) Hyperproteimemia
      *(4) Macroglobulinemia
      (5) Use of tranexamic acid
   D. Sudden reduction of systemic blood pressure because of cardiac
decompensation, surgical or traumatic shock, or therapy for arterial hypertension
   *E. Glaucoma (preexisting)
   F. Increased risk of thrombosis
      (1) Hereditary
         a. Antithrombin III deficiency
         b. Protein C deficiency or protein S deficiency
         c. Rare disorders of fibrinogen and fibrinolysis
            (i) Certain dysfibrinogenemias
            (ii) Abnormal plasminogen
      (2) Acquired
         a. Carcinoma
         b. Hematologic proliferative disorders
            (i) Acute promyelocytic leukemia
            (ii) Myeloproliferative disorders (polycythemia, essential
                 thrombocytemia)
         c. Behçet syndrome
         d. Lupus anticoagulant
         e. Nephrosis
         f. Complications of therapy
            (i) Oral contraceptives
            (ii) Infusion of prothrombin complex concentrates
            (iii) Heparin-induced thrombocytopenia
   G. Carotid-cavernous sinus fistula
   *H. Syphilis
   I. With immunoglobulin G (IgG) lambda monoclonal gammopathy
   J. Coil embolization of carotid-ophthalmic aneurysms
K. Oral contraceptive


### Extracted Table Central Retinal Vein Occlusion

**DILATED RETINAL VEINS AND RETINAL HEMORRHAGES**

1. Carotid-cavernous fistula  
2. Cavernous sinus fistula syndrome (carotid artery syndrome)  
3. Cavernous sinus thrombosis (hypophyseal-sphenoidal syndrome)  
4. Central retinal vein occlusion (see p. 468)  
5. Cervical tuberculosis  
6. Choroidal melanoma remote to the neovascularization  
7. Congenital tortuosity and dilatation of the retinal vessels  
8. Cryoglobulinemia  
9. Diabetes mellitus  
10. Intravitreal myiasis  
11. Leukemia  
12. Lymphomas  
13. Macroglobulinemia (Waldenstrom syndrome)  
14. Multiple myeloma (myelomatosis)  
15. Ophthalmic vein thrombosis  
16. Pappataci fever (phlebotomus fever)  
17. Paraproteinemias and dysproteinemias  
18. Polycythemia vera  
19. Retinal arteritis  
20. Sickle cell disease
**RETNAL HEMORRHAGES**

Retinal hemorrhages include bleeding that may be intraretinal or preretinal hemorrhages into the vitreous or subretinal hemorrhages.

1. Congestion of the head and neck, such as in newborns, in hanging, or during choking
2. Trauma, including electrical injury, hypothermal injury, and child abuse
3. Vascular obstruction, such as cardiorespiratory obesity syndrome, cystic fibrosis syndrome, negative acceleration syndrome, (hydrostatic pressure syndrome), ophthalmoplegic migraine syndrome, papilledema (see p. 593-601), subarachnoid hemorrhages, superior vena cava syndrome, Symonds syndrome (benign intracranial hypertension), thrombocytopenia, thrombosis, and Wernicke syndrome (avitaminosis B₁)
4. Inflammatory conditions, such as Criswick-Schepens syndrome (familial exudative vitreoretinopathy), Loffler syndrome (eosinophilic pneumonitis), perivasculitis, and subacute bacterial endocarditis
5. Acute febrile and infectious illnesses, including amebiasis ankylostomiasis, aspergillosis, bacterial endocarditis, coccidiodomycosis, cryptococcosis (torulosis), cysticercosis, dengue fever, hydatid cyst (echinococcosis), hydrophobia (rabies), infectious mononucleosis, influenza, Japanese River fever (typhus), lymphogranuloma venereum (Nicolas-Favre disease), metastatic bacterial endophthalmitis, nematode ophthalmia syndrome (toxocariasis), pertussis (whooping cough), Q fever, relapsing fever, trichinelllosis, Weil disease (leptospirosis), and yersiniosis
6. Vascular disease, such as arteriosclerosis, atherosclerosis, arteriovenous fistula, disseminated intravascular coagulation, hypertension, Paget syndrome (hypertensive diencephalic syndrome), progressive systemic sclerosis, pulmonary insufficiency, the retinopathies, particularly diabetic and hypertensive, and when the circulation through the eye is diminished in hypotensive retinopathy, such as in carotid vascular insufficiency syndrome or pulseless disease (Takayasu syndrome), suprarenal-sympathetic syndrome, temporal arteritis syndrome (cranial arteritis syndrome), and in conditions of extreme cachexia
7. Anemia that may be secondary to drugs, including the following:

  - acebutolol
  - acenocoumarin
  - acetaminophen
  - acetonilid
  - acetazolamide
  - acetoheaxamide
  - acetophenazine
  - actinomycin C
  - acyclovir
  - allobarbital
  - allopurinol
  - alprazolam
  - aminopterin
  - aminosalicylic acid (?)
  - amithiozone
  - amitriptylne
  - amobarbital
  - amodiaquine
  - amoxicillin
  - amphotericin B
  - ampicillin
  - anisindione
  - antazoline
  - antimony lithium
  - thiomalate
  - antimony potassium tartrate
antimony sodium
  tartrate
antimony sodium
  thioglycollate
antipyrine
aprobarbital
aminosalicylate (?)
atenolol
auranofin
aurothioglucone
aurothioglycanide
azatadine
azathioprine
barbital
BCG vaccine
bendroflumethiazide
benzathine penicillin G
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dicloxacillin
dicumarol
diethazine
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dimethindene
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diphenylhydantoin
diphenylpyraline
diphtheria and tetanus toxoids and pertussis vaccine (adsorbed)
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doxylamine
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droperidol
enalapril
erygocalciferol
erythromycin
ethacrynic acid
ethopropazine
ethosuximide
ethotoin
ethoxzolamide
ethyl biscoumacetate
fenfluramine
fenoprofen
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quinine  tetracycline  trifluperidol
ranitidine  thiabendazole  triflupromazine
rifampin  thiamylal  trimiprazine
rubella and mumps virus  thioethylperazine  trimethadione
vaccine (live)  thioguanine  tripelemannine
rubella virus vaccine  thiopental  triprolidine
(live)  thiopropazate  uracil mustard
secobarbital  thiopropazine  urethan
semustine  thioridazine  vancomycin
sodium  thiouracil  verapamil
antimonylgluconate  thiourea  vidarabine
stibogluconate  tolbutamide  vinblastine
stibophen  tolvaptan  vincristine
streptomycin  tolbutamide  vitamin A
sulfonamides  tranylcypromine  vitamin D
suramin  triazolam  vitamin D2
talbutal  triazolam  vitamin D3
temazepam  trichlormethiazide  warfarin

8. Vascularized neoplasms, including hereditary hemorrhagic telangiectasia (Rendu-Osler-Weber disease), and periocular and ocular metastatic tumors
9. Drugs, including the following:
acetylcholine  cobalt (?)  isosorbide
cacid bismuth sodium  cocaine  ketoprofen
tartrate  cortisone  lincomycin
adrenal cortex injection  deserpidine  mannnitol
aldosterone  desoxycorticosterone  medrysone
allopurinol (?)  dexamethasone  meprednisone
alsoroxylon  dibucaine  methaqualone
aspirin  dyclonine  methylphenidate
benoxinate  epinephrine  methylprednisolone
betamethasone  ethambutol  mithramycin
bismuth carbonate  fludrocortisone  mitotane
bismuth oxychloride  fluorometholone  oxyphenbutazone
bismuth salicylate  fluorouracil  paramethasone
bismuth sodium tartrate  fluprednisolone  penicillamine
bismuth sodium  glycerin  phenacetin
thioglycollate  heparin  phenylbutazone
bismuth sodium  hexachlorophene  piperocaine
triglycollamate  hydrocortisone  plicamycin
bismuth subcarbonate  indomethacin  pralidoxime
bismuth subsalicylate  iodide and iodine  prednisolone
butacaine  solution compounds  prednisone
proparacaine  sulfadimethoxine  sulfaﬁzole
radioactive iodides  sulfamerazine  sulﬁsazole
rauwolfia serpentina  sulfameter  sulindac
rescinnamine  sulfamethazine  syrosingopine
reserpine  sulfamethizole  tamoxifen
sodium chloride  sulfamethoxazole  tetracaine
sodium salicylate  sulfamethoxypyridazine  triamcinolone
sulfacetamide  sulfanilamide  trichloroethylene
sulfachlorpyridazine  sulfaphenazole  urea
sulfacytine  sulfapyridine  urokinase (?)
sulfadiazine  sulfasalazine  vitamin A

10. Hematopoietic system, such as the anemias, Bing-Neel syndrome (association of
macroglubulinemia and central nervous system symptoms), Fanconi syndrome (amino
diabetes), Gansslen syndrome (familial hemolytic icterus), Henoch-Schönlein purpura,
Herrick syndrome (sickle cell disease), Jacobsen-Brodwell syndrome, leukemias,
hemophilia, polycythemia, purpuras, oculootoooroerythropoietic disease, Plummer-
Vinson syndrome (sideropenic dysphagia syndrome), Reimann syndrome (hyperviscosity
syndrome), Waldenstrom syndrome (macroglubulinemia syndrome), Wiskott-Aldrich
syndrome (purpura), also following blood transfusion with incompatibility of blood
groups
11. Acosta syndrome (mountain climber syndrome)
12. Amyloidosis
13. Behçet syndrome (dermatostomatoophthalamic syndrome)
14. Bloch-Sulzberger syndrome (incontinentia pigmenti)
15. Bourneville syndrome (tuberous sclerosis)
16. Epidural steroid injection, gas myelography, or epiduroscopy
17. Following labor induced by oxytocin or dinoprostone in newborns
18. Following use of YAG laser
19. Histiocytosis X (Hand-Schüller-Christian syndrome)
20. HIV-related retinopathy
21. Hodgkin disease
22. Juvenile diabetes-dwarfism-obesity syndrome
23. Macular degeneration, age related (exudative type)
24. Morning-glory syndrome (hereditary central glial anomaly of the optic disc)
25. Mycosis fungoides syndrome (Sézary syndrome)
26. Neuroblastoma
27. Optic nerve drusen (see p. 559-560)
28. Paget syndrome (osteitis deformans)
29. Plasma lecithin (cholesterol acyltransferase deﬁciency)
30. Polymyalgia rheumatica
31. Polymyositis dermatomyositis (Wagner-Unverricht syndrome)
32. Porphyria cutanea tarda
33. Purtscher retinopathy
34. Radiation retinopathy
35. Sarcoidosis syndrome (Schaumann syndrome)
36. Schamberg disease (self-limiting cutaneous vasculitis)
37. Terson syndrome


**LARGE HEMORRHAGES IN THE FUNDUS OF AN INFANT OR YOUNG CHILD (SUGGESTIVE OF INCREASED INTRACRANIAL PRESSURE AND PARALYSIS OF CRANIAL NERVES)**

1. Hygroma
   *2. Shaken-baby syndrome*
3. Subarachnoid hemorrhage
4. Subdural hematoma


**RETINOVITREAL HEMORRHAGE IN A YOUNG ADULT**

1. Incontinentia pigmenti (Bloch-Sulzberger syndrome)
2. Congenital x-linked (juvenile) retinoschisis
   *3. Diabetes mellitus*
   *4. Sickle cell anemia*
5. Trauma
6. von Hippel-Lindau syndrome
RETINAL HEMORRHAGE WITH CENTRAL WHITE SPOT (ROTH SPOT)

1. Collagen disease
2. Cyanosis retinae-carcinoma of the lung
   *3. Diabetes mellitus
4. Following heart surgery
5. Following uncomplicated pediatric cataract extraction
   *6. Hematopoietic system
      A. Anemias
      B. Leukemia
      C. Multiple myeloma (Kahler disease)
7. Intracranial hemorrhage (infants)
8. Septic retinitis
   *A. *Candida albicans* infection
   B. Kala azar
   C. Phlebitis
   D. Rheumatic mitral and aortic valvulitis
   E. Rocky mountain spotted fever
   *F. Subacute bacterial endocarditis
   G. Syphilitic aortitis
   H. Viral pneumonia
9. Vascular disease

Extracted Table Retinal hemorrhage with Central White Spot

Mets MB, Del Monte M. Hemorrhagic retinopathy following uncomplicated pediatric cataract extraction. *Arch Ophthalmol* 1986; 104:975.


MICROANEURYSMS OF RETINA (PUNCTATE RED SPOTS SCATTERED OVER REGION OF POSTERIOR POLE)

1. Aging
2. Aplastic anemia-punctate hemorrhage
3. Associated with cotton-wool spots (see p. 491)
4. Bonnet-Dechaume-Blanc syndrome (cerebroretinal arteriovenous aneurysm syndrome)
5. Choroiditis
6. Chronic uveitis
7. Coats disease (retinal telangiectasia)
*8. Diabetes mellitus
9. Disseminated lupus erythematosus (Kaposi-Libman-Sacks syndrome)
10. Eales disease (periphlebitis)
11. Fabry disease (diffuse angiokeratosis)
*12. Hypertension
13. Hypotensive retinopathy, such as pulseless disease (aortic arch syndrome)
14. Kahler disease (myelomatosis)
15. Leukemias-punctate hemorrhages
16. Loa loa infection
17. Macroglobulinemia (Waldenstrom syndrome)
18. Mauriac syndrome (juvenile diabetes-dwarfism-obesity syndrome)
19. Ocular ischemic syndrome (carotid occlusive disease)
20. Osler hemorrhagic telangiectasia (hereditary hemorrhagic telangiectasis)
21. Pelizaeus-Merzbacher syndrome (aplasia axialis extracorticalis congenita)
22. Reimann syndrome (hyperviscosity syndrome)
*23. Retinoblastoma
24. Sickle cell hemoglobin C disease
25. Skin divers
26. Subacute bacterial endocarditis
27. Venous occlusion-occlusion of central retinal vein or one of its branches (see p. 468)


**Extracted Table Microaneurysms of Retina**

**RETINAL ARTERIOVENOUS SHUNT AT THE ARTERIOVENOUS CROSSING**

1. Diabetic retinopathy
2. Leber's miliary aneurysm
3. Retinal vein occlusion
4. Takayasu disease


MACROANEURYSMS OF RETINAL ARTERIES

These macroaneurysms are found within the first three orders of bifurcation of arterioles; they are frequently associated with localized hemorrhage and exudation.

1. Congenital
*2. Generalized arteriosclerosis
*3. Hypertension
4. Idiopathic
5. Following open heart surgery


RETINAL NEOVASCULARIZATION (GROWTH OF ABNORMAL NEW BLOOD VESSELS INTO THE VITREOUS)

1. Anemia
2. Behçet syndrome (dermatostomatoophthalmic syndrome)
*3. Central retinal vein occlusion (see p. 468)
*4. Diabetes mellitus
5. Eales disease (periphlebitis)
6. Ehlers-Danlos syndrome (fibrodysplasia elastica generalisata)
7. Hypertension (malignant and essential)
8. Leukemia
9. Lupus erythematosus
10. Macroglobulinemia (Waldenstrom syndrome)
11. Retinal detachment with hemorrhage
*12. Sickle cell disease
13. Syphilis (acquired lues)
14. Trauma
15. von Hippel-Lindau syndrome (retinocerebral angiomatosis)
16. Werlhof disease (hemophilia and thrombocytopenic purpura)


PREDISPOSITION TO RHEGMATOGENOUS RETINAL DETACHMENT
1. Aphakia (see p. 405)
2. Branch retinal vein occlusion
3. High myopia
4. Chorioretinitis
5. Peripheral retinal degeneration
   A. Vitreous base excavation
   *B. Retinal hole
   C. Retinoschisis
   D. Cystic retinal tuft
   E. Zonular traction tuft
   F. Meridional folds
   G. Partial-thickness retinal tear
   *H. Full-thickness retinal tear
   *I. Lattice degeneration
   J. Vitreous base avulsion
6. Trauma-blunt and perforating, including operation for strabismus and block excision of epithelial implantation cysts and tumors of the anterior uvea
7. Angiomatosis retinae


**RETINAL DETACHMENT (LOCATION AND MORPHOLOGIC CLASSIFICATION)**

1. Equator
   A. Myopic type-equatorial horseshoe tear
   B. Equatorial type associated with lattice degeneration
2. Ora serrata
   A. Aphakic, with multiple small breaks often in nasal periphery
   B. Dialysis in young, lower temporal quadrant, often bilateral
   C. Giant dialysis, often bilateral
3. Posterior pole
   A. Macular breaks, rare
   B. Other breaks at posterior pole, from cellular proliferation in inner retinal surface


**SYNDROMES AND DISEASES ASSOCIATED WITH RETINAL DETACHMENT**

1. Exudative
   A. Systemic disease
      (1) Abdominal typhus
      (2) Aspergillosis
      (3) Atopic dermatitis
      (4) Blood diseases
         a. Dysproteinemias
         b. Leukemia
         c. Sickle cell disease
      (5) Boutonneuse fever (rickettsia)
      (6) Candidiasis
      (7) Coenurosis
      (8) Cryoglobulinemia
      (9) Cryptococcosis
      (10) Cysticercosis
      (11) Disseminated intravascular coagulation
      (12) Extreme venous congestion, such as occurs during choking
      (13) Goldsheider syndrome (epidermolysis bullosa)
      (14) Goodpasture syndrome (chronic relapsing pulmonary hemosiderosis)
      (15) Grönblad-Strandberg syndrome (systemic elastodystrophy)
      (16) Histiocytosis X (Hand-Schüller-Christian syndrome)
      (17) Homocystinuria syndrome
      (18) Hurler syndrome (MPS I-H)
      (19) Hydatid cyst
      (20) Hypertension-grade IV
      (21) Krause syndrome (congenital encephalo-ophthalmic dysplasia)
      *(22) Lupus erythematosus
      (23) Lymphoma
      (24) Polyarteritis nodosa (Kussmaul disease)
      (25) Reese syndrome (D trisomy)
      (26) Regional enteritis
      (27) Relapsing polychondritis
      (28) Renal disease, including chronic glomerulonephritis or uremia
      (29) Rheumatoid arthritis
      (30) Rheumatic fever
(31) Rift Valley fever
(32) Sturge-Weber syndrome (meningocutaneous syndrome)
(33) Syphilis
(34) Temporal arteritis syndrome (cranial arteritis syndrome)
(35) Toxemia of pregnancy
(36) Vogt-Koyanagi-Harada syndrome

B. Ocular disease

(1) Acute retinal necrosis
(2) Choroidal or retinal tumor
   a. Hemangioma
   b. Melanoma
   c. Metastasis-including that from breast, lung, and stomach
   d. Retinoblastoma
(3) Colobomas of the optic nerve
(4) Dominant myopia and retinal detachment
(5) Familial exudative vitreoretinopathy
(6) Harada disease and Vogt-Koyanagi syndrome
(7) Lymphoid hyperplasia of the uveal tract
(8) Morning-glory syndrome (hereditary central glial anomaly of the optic disk)
(9) Nanophthalmos
(10) Norrie disease (atrophia oculi congenita)-x-linked
(11) Optic nerve pit
(12) Postinflammation of the orbit or sinuses or cyclitis
(13) Retina, congenital nonattachment and falciform folds-autosomal recessive
(14) Schwartz syndrome (glaucoma associated with retinal detachment)
(15) Scleritis (especially posterior scleritis)
(16) Sympathetic ophthalmia
(17) *Toxocara* infection
(18) Uveal effusion syndrome

C. Associated with retinal or choroidal vascular disease

(1) Coats disease (retinal telangiectasia)
   a. In juvenile
   b. In adult
(3) Central serous choroidopathy
(4) Detached pigment epithelium
(5) Eales disease (periphlebitis)
(6) Excessive panphotocoagulation
(7) Exudative age-related macular degeneration
(8) Hollenhorst syndrome (chorioretinal infarction syndrome)
(9) Incontinentia pigmenti
(10) Osteoporosis-pseudoglioma syndrome
(11) Post irradiation
(12) Scleral buckling
(13) Subpigment epithelium hemorrhage
(14) von Hippel Lindau disease (retinocerebral angiomatosis)

D. Drugs, including the following:

aceclidine  
adrenal cortex injection  
aldosterone  
beclomethasone  
be tamethasone  
carbachol  
chymotrypsin (?)  
cortisone  
chymotrypsin (?)  
cortisone  
demecarium (?)  
dexamethasone  
diisopropyl flurophosphate (DFP)  

echothiophate  
fludrocortisone  
fluorometholone  
fluprednisolone  
ganciclovir  
hydrocortisone  
isoflurophate (?)  
medrysone  
methyl prednisolone  
methylphenidate  
neostigmine (?)  
oxygen  
oxypenbutazone  
penicillamine  
phenylbutazone  
physostigmine (?)  
pilocarpine  
prednisolone  
prednisone  
sane paramethasone  
triamcinolone

2. Traction

*A. Pull of adherent and degenerated vitreous

B. Organized vitreous band

(1) After vitreous hemorrhage
   a. Spontaneous
   b. Traumatic

(2) Hypertensive retinopathy

(3) Posthemorrhagic proliferative retinopathy

(4) Sickle cell retinopathy

C. Postneovascularization of vitreous

*(1) Diabetic retinopathy, proliferative

(2) Eales disease (periphlebitis)

(3) Ehlers-Danlos syndrome (fibrodysplasia elastica-generalisata)

(4) Fibrinoid syndrome

*(5) Retinopathy of prematurity

(6) Severe uveitis

D. Congenital deformities, such as retinal dysplasia, coloboma, persistence of fetal vascular system, and pit of optic nerve

E. 18Q syndrome

F. Penetrating injury

*G. Proliferative vitreoretinopathy

H. Puckering syndrome

I. Retinal disinsertion syndrome

J. Retinopathy of prematurity

K. Warburg syndrome

3. Rhegmatogenous

A. Accommodation spasm, including strong miotics

B. Alport syndrome (neuropathy and deafness)

C. Apert syndrome (acrocephalosyndactylism syndrome)

D. Equatorial or anterior choroiditis

E. FOAR syndrome
F. Following YAG laser capsulotomy  
G. Hereditary ocular vitreoretinal degeneration and skeletal abnormality (cleft palate)  
H. Juxtapapillary microholes  
I. Knobloch syndrome (retinal detachment and occipital encephalocele)-autosomal recessive  
J. Marchesani syndrome  
K. Marfan syndrome (arachnodactyly dystrophia mesodermalis congenita)  
L. Marshall (D) syndrome  
M. Meckel syndrome  
N. Myopia, including staphyloma-autosomal dominant or recessive  
O. Retinal degeneration at periphery  
  (1) Presenile or myopic type  
  (2) Lattice and paving-stone types-autosomal dominant  
P. Retinal detachment-autosomal dominant or x-linked  
Q. Retinal vein occlusion  
R. Retinoschisis-adult or juvenile  
S. Smith-Magenis syndrome  
T. Spondyloepiphyseal dysplasia, congenital  
U. Stickler syndrome (hereditary progressive arthro-opthalmopathy)  
V. Trauma  
  (1) Direct injury-perforating wound and foreign body  
  (2) Indirect injury including block excision of epithelial implantation cysts and tumors of the anterior uvea.  
  (3) Post cataract operation  
  a. Sunset syndrome  
  b. Vitreous tug syndrome  
  (4) Battered-baby syndrome (Silverman syndrome)  
*W. Viral retinitis  
  (1) Acute retinal necrosis  
  (2) Cytomegalovirus retinitis  
X. Vitreous degeneration  
Y. Wagner syndrome (hyaloideoretinal degeneration) 


**RETINAL FOLDS**

1. Proliferative retinal folds-inner layer outstrips outer layer
2. Traction folds
   A. Associated with remnants of hyaloid artery
   B. Secondary to vitreous traction
   *C. Retinopathy of prematurity (cicatricial form)*
3. Falciform retinal fold (congenital retinal septum)
   A. Familial exudative vitreoretinopathy
   B. Isolated
   C. Trisomy syndrome
   D. Warburg syndrome
4. Chronic uveitis
5. Parasite
6. Occult intraocular foreign body
*7. Shaken baby syndrome*
8. Terson syndrome


**COTTON-WOOL SPOTS**

These spots are soft exudates (fluffy, white, focal infarcts in the nerve fiber layer).

1. Acute pancreatitis
2. Amniotic fluid embolization
3. Anemic conditions
   A. Cirrhosis of the liver
   B. Following cardiac surgery
   C. Gastric ulcer syndrome
   D. Hypotensive retinopathy
   E. Ligation of the carotid artery
F. Severe primary and secondary anemias  
G. Severe systemic blood loss  

4. Blood disease  
   A. Aplastic anemia  
   B. Dysproteinemia  
   C. Leukemia  
   D. Multiple myeloma (myelomatosis)  
   E. Pernicious anemia (vitamin B\textsubscript{12} deficiency)  
   F. Waldenström syndrome (macroglobulinemia syndrome)  

5. Carbon monoxide poisoning  
6. Carcinomatous cachexia  
7. Collagen diseases  
   A. Dermatomyositis (polymyositis dermatomyositis)  
   B. Diffuse scleroderma  
   C. Disseminated lupus erythematosus (systemic lupus erythematosus)  
   D. Polyarteritis nodosa (necrotizing angitis)  
   E. Rheumatoid arthritis with scleromalacia perforans or polymyalgia rheumatica  

*8. Diabetic retinopathy  
9. Hodgkin disease  
10. Infective conditions  
   *A. HIV  
   B. Pneumonia  
   C. Rheumatic fever  
   D. Rift Valley fever  
   E. Rocky mountain spotted fever (spotted fever)  
   F. Roth septic retinitis  
   G. Subacute bacterial endocarditis  

11. Microemboli following cardiac operation  
12. Primary amyloidosis (idiopathic amyloidosis)  
13. Primary open-angle glaucoma  
14. Protein C and protein S deficiency  
15. Purtscher retinopathy (fat embolism syndrome)  
16. Renal disease  
17. Serum disease  
18. Suprarenal-sympathetic syndrome (pheochromocytoma syndrome)  
19. Takayasu syndrome (aortic arch syndrome)  
20. Toxemic retinopathy of pregnancy  
*21. Untreated malignant hypertension  


**Extracted Table Cotton-wool spots**

**HARD EXUDATES (YELLOWISH WHITE DISCRETE MASSES DEEP IN THE RETINA)**

1. Circinate retinopathy  
2. Coats disease (retinal telangiectasia)  
*3. Diabetes mellitus  
*4. Exudative age-related macular degeneration  
*5. Hypertensive disease  
6. Radiation induced  
7. Retinal arterial macroaneurysm


**Extracted Table Hard Exudates**

**RETINAL EXUDATE AND HEMORRHAGE**

1. Capillary telangiectasis of retina (Reese)  
2. Coats disease (retinal telangiectasia)  
*3. Diabetes mellitus  
4. Eales disease (periphlebitis)  
5. Multiple retinal aneurysms (Leber syndrome)  
6. Racemose hemangioma of the retina  
7. von Hippel-Lindau, with absence of visible angioma (retinocerebral angiomatosis)


**RETINITIS OR PSEUDORETINITIS PIGMENTOSA**
Pigment may be bone corpuscular dots or heaped-up masses; salt and pepper fundus

1. Retinitis pigmentosa
   A. Abetalipoproteinemia (Bassen-Kornzweig syndrome)
   B. Alström disease (cataract and retinitis pigmentosa)
   C. Cockayne syndrome (dwarfism with retinal atrophy and deafness)
   D. Dialinas-Amalric syndrome (deaf-mutism-retinal degeneration syndrome)
   E. Hallgren syndrome (retinitis pigmentosa-deafness-ataxia syndrome)
   F. Hypotrichosis, syndactyly and retinitis pigmentosa-autosomal recessive
   G. Hunter syndrome (MPS II)
   H. Hurler syndrome (MPS I)
   I. Infantile phytanic acid storage disease
   J. Jeune syndrome
   *K. Kearns-Sayre syndrome (ophthalmoplegic retinal degeneration syndrome)
   L. Laurence-Moon-Bardet-Biedl syndrome (retinitis pigmentosa-polydactyly-adiposogenital syndrome)
   M. Metaphyseal chondrodysplasia with retinitis pigmentosa-autosomal recessive
   N. Microcephaly with chorioretinopathy
   O. Multiple sulfatase deficiency
   P. Muscular atrophy, ataxia, retinitis pigmentosa, diabetes mellitus-autosomal dominant
   Q. Olivopontocerebellar atrophy, type III
   R. NARP syndrome
   S. Pallidal degeneration, progressive with retinitis pigmentosa-autosomal recessive
   *T. Refsum syndrome (phytanic acid storage disease)
   U. Retinitis pigmentosa alone (usually autosomal recessive but may be autosomal dominant or sex linked)
   V. Retinitis pigmentosa associated with myopia, keratoconus, or glaucoma
   W. Retinitis pigmentosa, congenital deafness-sex linked
   X. Retinitis pigmentosa inversa (predominant-pigmentation around the disc and macula) and deafness-autosomal recessive
   Y. Retinitis pigmentosa, nerve deafness, mental retardation, and hypogonadism - autosomal recessive
   Z. Retinitis pigmentosa, PPRPE type (with preserved para-arteriole retinal pigment epithelium) - autosomal recessive
   AA. Retinitis pigmentosa, spastic quadriplegia, and mental retardation-autosomal recessive
   BB. Rud syndrome
   CC. Sanfilippo disease (MPS III)
   DD. Scheie disease (MPS IS)
   EE. Spielmeyer-Vogt syndrome (cerebroretinal degeneration)
   FF. Usher syndrome (hereditary retinitis pigmentosa-deafness syndrome)

2. Senile changes-degenerative pigmentation

3. Vascular lesion, such as occlusion of arteriole
4. Inflammatory
   A. Behçet disease (oculobuccogenital syndrome)
   B. Chickenpox virus
   C. Cytomegalic inclusion disease
   D. Dawson disease (inclusion-body encephalitis)
   E. Fetal varicella effects
   F. Focal dermal hypoplasia (Goltz syndrome)
   G. Harada disease (Vogt-Koyanagi-Harada syndrome)
   H. Hypomelanosis of Ito
   I. Influenza virus
   J. Nematode endophthalmitis (visceral larva migrans syndrome)
   K. Onchocerciasis (river blindness)
   L. Polyarteritis nodosa (Kussmaul disease)
   M. Rubella (German measles)
   N. Rubeola (measles)
   *O. Syphilis
   *P. Toxoplasmosis
   Q. Typhoid fever (enteric fever)
   R. Vaccinia

5. Toxic
   A. Accidental intraocular injection of depot corticosteroids
   B. Chloroquine and atabrine
   C. Diaminodiphenoxyalkanes—possible drug for treatment of schistosomiasis
   D. Indomethacin
   E. Phenothiazine
      (1) Chlorpromazine
      (2) Thioridazine (Mellaril)
   F. Pregl solution (Septojod, formerly used for treatment of puerperal sepsis)
   G. Quinine
   H. Sparsomycin

6. Acute lymphocytic leukemia
7. Alagille syndrome
8. Alport syndrome
9. Bardet-Biedl syndrome
10. Battens disease
11. Cryogenic "pigmentary fallout"—following use of cryosurgery for retinal detachment
12. Cystinosis syndrome (Lignac-Fanconi syndrome)
13. External ophthalmoplegias
    *14. Gardner syndrome (congenital hypertrophy of the retinal pigment epithelium and familial intestinal polyposis)
15. Hagberg-Santavuori syndrome (neuronal ceroid lipofuscinosis)
16. Hallervorden-Spatz syndrome (pigmentary degeneration of globus pallidus)
17. Hereditary ataxias (Friedrich and Marie)
18. Leber congenital amaurosis
19. Lens dislocated into vitreous
20. MERRF syndrome
21. Mucolipidoses IV (ML IV)
22. Myotonic dystrophy syndrome (Curschmann-Steinert syndrome)
23. Neuronal ceroid lipofuscinosis
   A. Infantile form
   B. Late infantile form (Jansky-Bielschowsky)
   C. Adult form (Kufs syndrome)
24. Pelizaeus-Merzbacher syndrome (aplasia axialis extracorticalis congenita)
25. Pellagra (ariboflavinosis)
26. Progressive cone-rod degeneration
27. Renal disorders, including familial juvenile nephronophthisis (medullary cystic
disease)
28. Rud syndrome (hypophyseal deficiency)
29. Sjögren-Larsson syndrome (oligophrenia-ichthyosis-spastic diplegia syndrome)
30. Tapetal-like reflex syndrome
31. Trauma, including blunt, penetrating, obstetric, and radiotherapy, Frenkel syndrome
   (ocular contusion syndrome)
32. Waardenburg syndrome (interoculoridodermatoauditive dysplasia)
33. Zellweger syndrome and pseudo-Zellweger syndrome


Bloome MA, Garcia CA. *A manual of retinal and choroidal dystrophies*. East Norwalk,


Roy FH. *Ocular syndromes and systemic diseases*, 3rd ed. Philadelphia: Lippincott
Williams & Wilkins, 2002.

**Extracted Table Retinal bone corpuscular dots**

**LESIONS CONFUSED WITH RETINOBLASTOMA**

1. Anomalous optic disc
2. Anteriorly dislocated lens with secondary glaucoma
*3. Coats disease (retinal telangiectasia)
4. Coloboma of choroid and optic disc (see p. 555-556)
5. Congenital corneal opacity
6. Congenital rubella syndrome (Gregg syndrome)
7. Cysts in a remnant of the hyaloid artery
8. Developmental retinal cyst
9. Glioma of the retina
10. Hematoma under retinal pigment epithelium
11. High myopia with advanced chorioretinal degeneration
12. Juvenile (x-linked) retinoschisis
13. Juvenile xanthogranuloma (nevoxanthoendothelioma)
* 14. Larval granulomatosis (T. canis)
15. Medullation of nerve fiber layer
16. Metastatic endophthalmitis
17. Norrie disease (atrophia oculi congenital
18. Oligodendroglioma of the retina
19. Organization of intraocular hemorrhage
*20. Persistent hyperplastic primary vitreous
21. Retinal detachment due to choroidal or vitreous hemorrhage
22. Retinal dysplasia (massive retinal fibrosis)
*23. Retinopathy of prematurity
24. Retrolental membrane associated with Bloch-Sulzberger syndrome (incontinentia pigmienti)
25. Rhegmatogenous and falciform retinal detachment
26. Secondary glaucoma
27. Sex-linked microphthalmia
28. Tapetoretinal degeneration
29. Trisomy 13 (Patau syndrome)
*30. Toxoplasmosis (ocular toxoplasmosis)
31. Traumatic chorioretinitis
*32. Tumors other than retinoblastoma
33. Uveitis in secondary retinal detachment
34. "White-with-pressure" sign


**Extracted Table Single dark fundus lesion**

**SINGLE WHITE LESION OF RETINA**

*1. Amelanotic melanoma
2. Astrocytoma of tuberous sclerosis
3. Degeneration of retinal pigment epithelium
4. Diktyoma
5. Glioma of optic nerve
6. Hamartomas of the optic disc of retinitis pigmentosa
7. Metastatic or direct extension of a tumor
8. Neurofibroma of von Recklinghausen syndrome
*9. Retinoblastoma
*10. T. canis
PALE FUNDUS LESIONS

1. Generalized pallor
   A. Albinism - photophobia; defective vision; absence of pigment in iris, retina. And choroid
   B. Chediak-Higashi syndrome (oculocutaneous albinism with recurrent infections)
   C. Waardenburg syndrome (embryonic fixation syndrome)
   D. Choroideremia - rare; night blindness; contraction of visual fields; degeneration of pigment epithelium in periphery with exposure of choroidal vessels
   E. Myopia - thinning of retina and choroidal crescent at disc
   F. Retinal ischemia
      (1) Occlusion of retinal arteries (see p. 457)
      (2) Spasm of retinal arteries - angiospasm: quinine, lead poisoning, migraine, or Raynaud disease
      (3) Anemia
   *G. Vascular retinopathies - hypertension, edema, hemorrhages, swelling of disc
   H. Leukemia
I. The lipidoses
   (1) Congenital, rare
   (2) Infantile (Tay-Sachs disease)
   (3) Late infantile - (Jansky-Bielschowsky syndrome) - 2 to 4 years of age
   (4) Juvenile - (Spielmeyer-Vogt syndrome) - 5 to 8 years of age; optic atrophy
   (5) Adult - (Kufs disease) - 15 to 25 years of age; eyes may be normal or show some pigmented macular changes
   J. Gaucher disease (glucocerebrosidase storage disease)
   K. Hereditary dystrophic lipidosis (Fabry disease)
   L. Hyperlipemia
      (1) Diabetes - rare, yellowish retinal and choroidal vessels
      (2) Essential hyperlipemic xanthomatosis - rare, yellowish retinal and choroidal vessels
   M. Oguchi disease

2. Localized pale areas
   A. Medullated nerve fibers (see p. 507)
   B. Retinopathy of prematurity
   C. Localized retinal edema
      (1) Inflammation
      (2) Trauma
      (3) Vascular lesions
   D. Retinal detachment and schisis (see p. 487)
*E. Retinoblastoma
F. Coats disease (retinal telangiectasia)
G. Coloboma (see p. 450-451)
H. Normal fundus features - pale streaks mark site of ciliary nerves
I. Atrophic areas - diathermy, light coagulation, or cryosurgery
J. Scattered retinal exudates
   (1) Preretinal - severe posterior uveitis; discrete white spots, often most marked along vessels adjacent to a patch of choroiditis
   (2) Retinal
      a. Purtscher compression syndrome - cotton-wool spots
      b. Fat emboli
      c. Hemangiomatosis - yellow exudates
      *d. Hypertensive retinopathy - cotton-wool and hard exudates
      e. Toxemia of pregnancy
      f. Hypotensive retinopathy
      g. Pulseless disease (Takayasu syndrome)
      h. Arterial occlusion
      i. Blood loss - cotton-wool spots
      j. Anemia (all types)
      k. Leukemia
      l. Purpura
      m. Macroglobulinemia (Waldenström syndrome)
      n. Hodgkin disease - soft exudates
      *o. Diabetes - cotton-wool and hard exudates
      p. Hypercholesterolemia - lipid deposits
      q. Systemic lupus erythematosus (disseminated lupus erythematosus)
      r. Dermatomyositis - cotton-wool spots
      s. Polyarteritis nodosa (Kussmaul disease)
      t. Scleroderma (progressive systemic sclerosis)
      u. Vitamin A deficiency - small white spots along course of retinal vessels
      v. Retinal capillariosis - yellowish white spots in substance of retina
      w. Leber congenital retinal aplasia - bilateral blindness, multiple white specks
      x. Female carrier of retinitis pigmentosa - brilliant silvery reflex with shining yellow spots deep to retinal vessels

3. Dystrophic conditions
   A. Gyrate atrophy - rare, irregular atrophic areas with visual defects and night blindness
   B. Choroidal sclerosis - rare, diffuse peripapillary or central choroidal atrophy with larger choroidal vessels prominent
   C. Infarction or occlusion of ciliary arteries - rare, embolism (air, fat), injury, atrophic area with prominent choroidal vessels
   D. Pseudoinflammatory macular dystrophy - rare, fourth to sixth decades, central edema, hemorrhage and exudate, bilateral and symmetric
E. Helicoid peripapillary chorioretinal atrophy-rare, congenital and adult forms, star-shaped atrophic areas radiating from disc

F. Retinitis punctata albscens-rare, onset in second and third decades, multiple discrete whitish dots which may appear crystalline, night blindness and field defects in progressive type

G. Fundus flavimaculatus-rare, onset in second and third decades, yellow flecks deep in the retina

H. Geographic choroiditis-rare, map-like pigmentary disturbance at posterior pole or more widespread over posterior fundus

I. Doyne honeycomb dystrophy-rare; middle age and older; drusen at posterior pole, with pigmented or cystoid macular changes

J. Progressive bifocal chorioretinal atrophy-atrophy temporal to disc, extending later; night blindness in late stage


**MEDULLATED NERVE FIBERS**

In this condition, an opaque white patch is usually adjacent to and may cover the disc; it is localized to one sector of the disc and peripapillary or arcuate with a peripheral, feathered edge.

*1. Isolated finding
2. Autosomal-recessive or -dominant inheritance
3. Associated with the following:
   A. Aplasia of macula
   B. Coloboma of optic nerve or choroid (see p. 555-556)
   C. Conus of disc
   D. Cranial dysostosis (oxycephaly, dolichocephaly, brachycephaly, and craniofacial dysostosis)
   E. Hyaloid remnants
   F. Macular colobomas (see p. 450)
   G. Myopia
   *H. Neurofibromatosis


**PIGMENTED FUNDUS LESIONS**
1. Diffuse pigmentation
   A. Negroid fundus-accentuation of fundus pigmentation
   B. Melanosis bulbi-rare, pigmentation of external eye and fundus
   C. Nevus of Ota
   D. Waardenburg syndrome (embryonic fixation)
2. Single pigmented lesions
   A. Flat lesions
      (1) Benign melanoma-bluish, gray, or black lesion
      (2) Pigmented scar-patch of dense pigment, usually atrophic area in center
      (3) Fuchs dark spot-dark spot in macular region
      *(4) Macular degeneration (exudate, age-related)
   B. Raised lesions
      (1) Simple detachment (see p. 487)
         a. Macular, such as in central serous retinopathy
         b. Associated with uveitis, such as that associated with Vogt-Koyanagi-Harada syndrome
         c. Hemorrhagic macrocyst
      *(2) Malignant melanoma-raised pigmented lesion with secondary detachment, abnormal vessels
      (3) Choroidal hemorrhage-trauma, spontaneous in patients with vascular disease, high myopia
      *(4) Exudative macular lesion - common, old age, subretinal exudate
      (5) Hemangioma of choroid - rare, raised grayish tumor near disc. Secondary detachment later
      *(6) Metastatic tumor - flat tumor with little pigment, primary in breast, or lung
      (7) Chorioretinitis
      (8) Foreign body
      (9) Coats disease (retinal telangiectasia)
3. Multiple pigmented lesions
   A. Scattered focal lesions
      *(1) Congenital melanosis - cat's-paw patches of pigment in one sector of fundus (may be part of Gardner syndrome)
      (2) Postinflammatory - flat pigment with areas of atrophy
      (3) Hypertensive retinopathy-hypertensive vascular changes with scattered pigmentation
      (4) Siegrist streaks-rare, chain of pigment spots along sclerosed choroidal vessel
      (5) Paravenous retinochoroidal atrophy - paravenous pigmentation with chorioretinal atrophy
      (6) Incontinentia pigmenti (Bloch-Sulzberger syndrome)
      (7) Chorioretinal scars from cryosurgery
   B. Widely disseminated pigmentary changes
      (1) Genetic conditions
         a. Typical retinitis pigmentosa-attenuation of retinal vessels, optic atrophy (myopia, posterior polar cataract, keratoconus)
b. Atypical retinitis pigmentosa - rare, little or no pigment, pigment in clumps
c. Retinitis pigmentosa syndromes
   (i) Cockayne syndrome (dwarfism with retinal atrophy and deafness)
   (ii) Hallgren syndrome (retinitis pigmentosa-deafness-ataxia syndrome)
   (iii) Kearns syndrome (ophthalmoplegic retinal degeneration syndrome)
   (iv) Laurence-Moon-Biedl syndrome (retinitis pigmentosa polydactyly-adiposogenital syndrome)
   (v) Leber congenital retinal aplasia syndrome
   (vi) Lignac-Fanconi syndrome (cystinosis syndrome)
   (vii) Myotonic dystrophy syndrome (dystrophia myotonica syndrome)
   (viii) Pelizaeus-Merzbacher syndrome (aplasia axialis extracorticalis congenital)
(2) Infectious conditions - secondary retinitis pigmentosa
   *a. Syphilis (congenital) - pepper-and-salt pigmentation, interstitial keratitis
   b. Syphilitic neuroretinitis - rare, retinitis pigmentosa
   c. Rubella - cataract, secondary retinitis pigmentosa (nonprogressive)
   d. Vaccinia - rare, retinitis pigmentosa, history of vaccination
(3) Metabolic Disturbances
   a. Refsum syndrome (phytanic acid storage disease)
   b. Bassen-Kornzweig syndrome (familial hypolipoproteinemia)
(4) Toxic conditions, such as chloroquine, phenothiazine derivatives; usually central pigmentation; cornea and lens change
4. Ciliary body and choroid
   A. Tumors
      (1) Hemangioma
      (2) Malignant melanoma
      *(3) Metastatic carcinoma, such as that from the lungs, breast, testis, kidney, prostate gland, bladder
      (4) Nevus
      (5) Neurilemmoma
      (6) Neurofibroma
   B. Detachment - serous or hemorrhagic
   *C. Lymphoma and leukemias
   D. Peripheral giant cysts
5. Vitreous body
   A. Hemorrhage
   B. Abscess
6. Staphyloma of sclera
CHOLESTEROL EMBOLI OF RETINA (HOLLENHORST PLAQUES)

Bright-yellow plaques are often observed at bifurcation of arterioles, indicative of generalized atherosclerosis, and should signal the ophthalmologist to measure retinal artery pressures and refer the patient for general medical evaluation.

1. Abdominal aortic aneurysms
   *2. Aortic stenosis
3. Arteriography showing occlusions in one or more cervical arteries
4. Atrial fibrillation
5. Bleeding duodenal or gastric ulcer
   *6. Bruits in one or both carotid arteries
   *7. Calcification of internal carotids (Doppler ultrasonography)
8. Congestive heart failure
9. Coronary heart disease with myocardial infarct or angina
   *10. Diabetes mellitus
11. New or old strokes or transient attacks of cerebral ischemia
12. Peripheral atherosclerosis obliterans, popliteal or femoral aneurysms
13. Renal artery occlusions
   *14. Retinal arterial occlusions
15. Torsion and calcification of aorta (roentgenogram)
16. Vocal cord paralysis (aortic arch aneurysm)


RETINAL MICROEMBOLI

1. Platelet fibrin-mural or "tail" thrombus in carotid occlusion
   *2. Cholesterol-lipid-eroding atheroma in carotid bifurcation
3. Calcific or fibrinoid
   A. Calcific valvular disease dislodged spontaneously following cardiac catheterization, or angiography, or prolapse of mitral valve
   B. Rheumatic heart disease
   C. Myocardial disease
   D. Septic emboli
4. Foreign bodies
   A. Silicone or cloth particles covered cardiac valves
B. Talc or cornstarch emboli from drug addicts
C. Mercury
D. Secondary to retrobulbar or intranasal methyl prednisolone acetate

5. Tumors
   A. Cardiac myxomas
   B. Metastatic tumors including malignant melanomas and breast carcinomas

6. Fat emboli from fracture of the long bones
7. Air emboli from crushing injuries of the chest


LIPEMIA RETINALIS (ARTERIOLES AND VENULES SIMILAR IN COLOR AND APPEARING ORANGE-YELLOW TO WHITE)

1. Primary hyperlipoproteinaemia
   A. Type I-familial fat-induced hyperlipoproteinaemia (hyperchylomicronemia)
   B. Type III-familial hyperbetalipoproteinaemia and hyperprebetalipoproteinaemia (carbohydrate-induced hyperlipemia)
   C. Type IV -familial hyperprebetalipoproteinaemia (carbohydrate-induced hyperlipemia)
   D. Type V-familial hyperchylomicronemia with hyperprebetalipoproteinaemia (mixed hyperlipemia)
*2. Diabetes mellitus with hyperlipemia

3. Secondary hyperlipoproteinaemia
   A. Biliary obstruction
   B. Chronic pancreatitis
   C. Chronic renal failure
   D. Coats disease in adults (retinal telangiectasia)
   E. Glycogen storage disease
   F. Hypergammaglobulinemia
   G. Hypothyroidism (cretinism)
   H. Idiopathic hypercalcaemia
   I. Insulin-deficient diabetes mellitus (Willis disease)
*J. Malignant neoplasms
   K. Nephrotic syndrome (lipoid nephrosis)
   L. Progressive lipodystrophy


HEMORRHAGIC OR SEROUS EXUDATES BENEATH PIGMENT EPITHELIUM

1. Angioid streaks (see p. 526-529)
2. Best macular degeneration (vitelliruptive macular dystrophy)
3. Coats disease (retinal telangiectasia)
4. Doyne honeycomb macular degeneration
*5. Histoplasmosis (histoplasmosis choroiditis)
*6. Macular drusen in age-related macular degeneration
7. Myopia
8. Solid neoplasms
9. Trauma


RETINAL VASCULAR TUMORS AND ANGIOMATOSIS RETINAE SYNDROMES

1. Associated with pheochromocytoma
2. Blue rubber bleb nevus syndrome (Bean syndrome)
3. Bonnet-Dechaune-Blanc syndrome (neuroretinal angiomatosis syndrome)
4. Cavernous retinal hemangioma - intraretinal angiomas
*5. Coats disease (retinal telangiectasia)
6. Gorlin syndrome
7. Racemose angioma - with arteriovenous anomalies of central nervous system (Wyburn-Mason syndrome)
8. Retinal telangiectasis (Leber military aneurysms) - telangiectasia retinae of Reese
9. Sturge-Weber syndrome (meningocutaneous syndrome)
*10. von Hippel-Lindau syndrome (retinocerebral angiomatosis retinae)


Extracted Table Retinal vascular tumors
Extracted Traumatic retinopathies

RETINAL "SEA-FANS"
These are vasoproliferative lesions with a characteristic fan-shaped appearance, also called a "parachute" lesion.

1. Aortic arch syndrome (pulseless disease)
2. Carotid-cavernous fistula (carotid artery syndrome)
3. Central and branch retinal vein occlusion (see p. 468)
4. Chronic myelocytic leukemia
5. Diabetes mellitus
6. Eales disease (periphlebitis)
7. Facioscapulohumeral muscular dystrophy (FSH [facio scapulo numeral] syndrome)
8. Incontinentia pigmerti I (Block-Sulzberger syndrome)
9. Familial exudative vitreoretinopathy (Criswick-Schepens syndrome)
10. Hemoglobin C trait
11. Leukemia
12. Long-standing retinal detachment
13. Lupus erythematosus
14. Macroglobulinemia
15. Multiple sclerosis
16. Polycythemia vera (erythremia)
17. Retinopathy of prematurity
18. Sarcoidosis syndrome
19. Sickle cell disease
20. Talc and cornstarch emboli
21. Uveitis, including pars planitis


**RETINAL VESSELS DISPLACED TEMPORALLY**

1. Familial vitreoretinopathy
2. Hamartomas
3. Inflammation
4. Myopia with lattice-like retinal degeneration
5. Retinopathy of prematurity
6. Sickle cell disease (drepanocytic anemia)
7. Trauma
RETINAL VESSELS DISPLACED NASALLY

1. Axial myopia  
   *2. Glaucoma  
3. Inflammation  
4. Trauma

PERIPHERAL FUNDUS LESIONS

1. Pale raised lesions  
   A. Vitreous opacities-white fluffy or discrete opacities, associated with pars planitis or sarcoid uveitis  
   B. Retinopathy of prematurity-retinal edema and dense white lesions with neovascularization  
   C. Toxocariasis (nematode ophthalmia syndrome)-vitreous opacities with peripheral granuloma  
   D. Leprosy (Hansen disease)-peripheral exudates with anterior uveal involvement  
   E. Vitreoretinal dystrophies-bands in vitreous with retinoschisis or retinal detachment  
   F. Angiomatosis-retinal tumor with enlarged, feeding vessels  
   *G. Retinoblastoma-raised, creamy-white, fluffy lesion without inflammatory signs

2. Flat lesions  
   A. Coloboma-pale area with pigmented edge in region of fetal cleft  
   B. Chorioretinitis  
      (1) Disseminated, congenital syphilis-pepper-and-salt or larger confluent lesions  
      (2) Toxoplasmosis-pigmented scars of old lesions  
      (3) Cytomegalic inclusion disease-localized chorioretinitis or general peripheral infiltration  
      *(4) Histoplasmosis-peripheral punched-out lesions with or without pigmentation  
   C. Peripheral degenerations
(1) Senile changes-of eyes older than years of age, depigmented areas with pigmented margins (cobblestone degeneration).
(2) Secondary pigmentary degeneration-peripheral pigmentary changes similar to senile type or to retinitis pigmentosa
(3) Cystoid degeneration-multiple cystic spaces and thin areas in peripheral retina
*(4) Lattice degeneration-lace work of white lines with depigmented and pigmented patches
(5) Cystinosis (cystine storage aminoaciduria dwarfism syndrome)-granular rings of pigment in periphery of fundus, similar to cobblestone degeneration

D. Equatorial linear pigment disturbance
   (1) Ophthalmomyiasis internal
   (2) Histoplasmosis syndrome

E. Retinitis
   *(1) Acute retinal necrosis
   *(2) Cytomegalovirus retinitis

3. Dark raised lesions
   A. Choroidal detachment (see p. 532-535)
      (1) Spontaneous-slowly progressive detachment, no inflammatory signs
      *(2) Postoperative-intraocular operation; particularly for cataract and glaucoma; shallow anterior chamber; leaking wound
      (3) Exudative
         a. Inflammatory-shallow anterior chamber, myopia, and peripheral detachment
         b. Vascular-nephritis, hypertension, toxemia of pregnancy, polyarteritis nodosa, leukemia
            *(c. Tumors-intraocular tumors; tumors of orbit and lacrimal gland
            d. Traumatic-contusion injuries, perforating wounds, hypotony, anterior chamber may be shallow or deep if perforation occurs posteriorly
   B. Exudative retinal detachment (see p. 487)
      *(1) Secondary to general disease with retinopathy-hypertension, toxemia of pregnancy, leukemia, dysproteinemia, polyarteritis nodosa, rickettsial arteritis, venous congestion, talc and cornstarch emboli
      (2) Secondary to local disease of the eye-inflammatory signs with exudative detachment. Harada disease, sympathetic ophthalmitis, scleritis, tenonitis, choroidal tumor, and ophthalmomyiasis
   C. Simple detachment-myopia in two thirds of patients, trauma, may follow cataract extraction or discission for congenital cataract
   D. Cysts
      (1) Ciliary body-larger cysts usually push iris forward; rarely, cyst extends backward to be seen ophthalmoscopically
      (2) Pars plana-may enlarge and appear as a multilocular reddish-brown cyst
   E. Scleral indentation-retinal detachment operation
F. Neoplasms of ciliary body
   (1) Benign epithelioma - brown spot to 2 mm in diameter on surface of ciliary body
   *(2) Other tumors - diktyoma, leiomyoma, reticuloses, neurofibroma, malignant melanoma, rare, usually present as a mass protruding through the root of iris; may cause glaucoma; dark bulge seen ophthalmoscopically; lens changes adjacent to tumor

G. Neoplasms of choroid
   (1) Congenital melanosis-cat's-paw patches of pigment in one sector of fundus
   (2) Choroidal nevus - flat, bluish gray or black lesion
   *(3) Malignant melanoma-raised, pigmented lesion with secondary detachment
   (4) Secondary metastatic-rare, primary lesion in breast, lung, and so on

4. Vascular lesions
   A. Periphlebitis (Eales disease) common; young adults; sheathing of peripheral veins; hemorrhages in new vessels and later retinal detachment (see p. 487)
   *B. Perivasculitis secondary to uveitis - perivascular infiltration, particularly in pars planitis, sarcoidosis, Behçet disease, and toxoplasmosis
   C. Systemic diseases
      (1) Rickettsia-engorgement of veins, retinal edema, hemorrhages, and exudates
      (2) Multiple sclerosis (disseminated sclerosis) - sheathing of veins (see p. 468)
      (3) Polyarteritis nodosa (necrotizing angiitis) - hemorrhages, exudates, and serous detachment of retina (see p. 488)
      (4) Tuberculin or BCG inoculation-rare, sheathing of peripheral veins with hemorrhages
      *(5) Sickle cell retinopathy (Herrick syndrome) - dilatation of peripheral veins, hemorrhages, and connective tissue sheets in periphery; new vessel formation


**RETINAL DISEASE ASSOCIATED WITH CORNEAL PROBLEMS**

1. Abdominal typhus (enteric fever)-corneal ulcer, retinal detachment, central retinal artery emboli
2. Acanthamoeba-keratitis, pannus, corneal ring abscess, retinal perivasculitis
3. African eyeworm disease-keratitis, central retinal artery occlusion, macular hemorrhages
4. Amyloidosis - amyloid corneal deposits, corneal dystrophy, retinal hemorrhages
5. Anderson-Warburg syndrome (oligophrenia-microphthalmos syndrome)-corneal opacification, malformed retina with retina pseudotumors
6. Angioedema (hives)-central serous retinopathy, corneal edema
*7. Anterior segment ischemia syndrome-corneal edema midperiphery retinal hemorrhages
8. Apert syndrome (acrocephalosyndactyly)-exposure keratitis, retinal detachment
9. Arteriovenous fistula - bullous keratopathy, retinal hemorrhages
10. Aspergillosis-corneal ulcer, keratitis, retinal hemorrhages, retinal detachment
11. Atopic dermatitis-keratoconjunctivitis, keratoconjunctivitis sicca, corneal ulcer, central retinal vein occlusion, retinal detachment
12. Avitaminosis C-retinal hemorrhages, keratitis, corneal ulcer
13. Bacillus cereus-ring abscess of cornea, necrosis of retina
14. Bang disease (brucellosis)-keratitis, choroidal retinitis, macular edema
15. Behçet syndrome (dermatostomata-ophtalmic syndrome)-keratitis, posterior corneal abscess, retinal vascular changes
16. Biéty disease (Bietti marginal crystalline dystrophy)-marginal corneal dystrophy, retinitis punctata albscens
17. Candidiasis-keratitis, corneal ulcer, retinal atrophy, retinal detachment
*18. Carotid artery syndrome-corneal ulcer, loss of corneal sensation, retinal edema, engorgement of retinal veins
19. Chickenpox (varicella)-corneal ulcer, corneal opacity, retinitis, hemorrhagic retinopathy
20. Chloroquine-corneal epithelial pigmentation, macular lesions
21. Chronic granulomatous disease of childhood-keratitis, destructive chorioretinal lesions
22. Cockayne syndrome (dwarfism with retinal atrophy and deafness)-pigmentary degeneration, band keratopathy, corneal dystrophy
23. Crohn disease (granulomatous ileocolitis)-marginal corneal ulcers, keratitis, macular edema, macular hemorrhages
24. Cryoglobulinemia-deep corneal opacities, venous stasis
25. Cystinosis (aminoaciduria)-crystals in cornea and pigment in retina
26. Dengue fever-keratitis, corneal ulcer, retinal hemorrhages
27. Diffuse keratoses syndrome-corneal nodular thickening in the stroma worse in fall, retinal phlebitis
28. Diphtheria-keratitis, corneal ulcer, central artery occlusion
29. Disseminated lupus erythematosus (Kaposi-Libman-Sacks syndrome)-keratitis, keratoconjunctivitis sicca, corneal ulcer, central retinal vein occlusion, retinal detachment
30. Ehlers-Danlos syndrome (cutis hyperelastica)-keratoconjunctivitis sicca, corneal ulcer, central retinal vein occlusion, retinal detachment
31. Electrical injury-corneal perforation, retinal edema, retinal hemorrhages, pigmentary degeneration, retinal holes, dilatation of retinal veins
32. Fabry disease (diffuse angiokeratosis)-whorl-like changes in cornea, central retinal artery occlusion, tortuosity of retinal vessels
33. Goldscheider syndrome (epidermolyis bullosa)-bullous keratitis with opacities, retinal detachment
34. Gronblad-Strandberg syndrome (systemic elastodystrophy) - angiod streaks of the retina, macular hemorrhages, retinal detachment, keratoconus
35. Hamman-Rich syndrome (alveolar capillary block syndrome) - keratomalacia ischemic retinopathy, cystic macular changes
36. Heerfordt syndrome (uveoparotid fever) - band keratopathy, retinal vasculitis
37. Hennebert syndrome (luetic otitic nystagmus syndrome) - interstitial keratitis, disseminated syphilitic chorioretinitis
38. Histiocytosis X (Hand-Schüller-Christian syndrome) - retinal hemorrhage, retinal detachment, bullous keratopathy, corneal ulcer, pannus
39. Hodgkin disease - keratitis, retinal hemorrhages
40. Hollenhorst syndrome (chorioretinal infarction syndrome) - hazy cornea, serous retinal detachment, pigmentary retinopathy
41. Hunter syndrome (MPS II) - splitting or absence of peripheral Bowman membrane, stromal haze, pigmentary retinal degeneration, narrowed retinal vessels
42. Hurler-Scheie syndrome (MPS IH-S) - corneal clouding, pigmentary retinopathy
43. Hurler syndrome (gargoylism) - diffuse corneal haziness, retinal pigmentary changes, megalocornea, retinal detachment
44. Hydatid cyst (echinococcosis) - keratitis, abscess of cornea, retinal detachment, retinal hemorrhages
45. Hyperlipoproteinemia - arcus juvenilis, lipemia retinalis, xanthomata of retina
46. Hyperparathyroidism - band keratopathy, vascular engorgement of retina
47. Hypovitaminosis A - keratomalacia with perforation, corneal opacity, retinal degeneration
48. Idiopathic hypercalcemia (blue-diaper syndrome) - band keratopathy, optic atrophy, papilledema
49. Indomethacin - corneal deposits, reduced retinal sensitivity
50. Influenza - keratitis, retinal hemorrhage
51. Japanese River fever (typhus) - keratitis, retinal hemorrhages
52. Juvenile rheumatoid arthritis (Still disease) - band keratopathy, macular edema
53. Kahler disease (multiple myeloma) - crystalline deposits of cornea, central retinal artery occlusion, retinal microaneurysms
54. Kussmaul disease (periarteritis nodosa) - retinal detachment, pseudoretinitis pigmentosa, corneal ulcer
55. Leber tapetoretinal dystrophy syndrome (retinal aplasia) - keratoconus, salt-and-pepper or "bone corpuscle" pigmentation, yellowish brown or gray macular lesions
56. Lubarsch-Pick syndrome (primary amyloidosis) - amyloid corneal deposits, retinal hemorrhages
*57. Lymphogranuloma venereum disease (Nicolas-Favre disease) - keratitis, pannus, corneal ulcer, keratoconus, tortuosity of retinal vessels, retinal hemorrhages
58. Marfan syndrome (arachnodactyly dystrophia mesodermalis congenita) - keratoconus, retinitis pigmentosa
59. Meckel syndrome (dysencephalia splanchnocystic syndrome) - sclerocornea, microcornea, retinal dysplasia
60. Meningococcemia - keratitis, retinal endophlebitis
61. Mikulicz-Radeski syndrome (dacryosialoadenopathy) - keratoconjunctivitis, retinal candlewax spots
62. ML IV (mucolipidosis IV)- corneal clouding, corneal opacities, retinal atrophy
63. Morbilli (measles-rubeola)- keratitis, corneal ulcer, pigmentary retinopathy, central retinal artery occlusion
64. Mucormycosis (phycomycosis)- corneal ulcer, striate keratopathy, retinitis, central retinal artery thrombosis
65. Mycosis fungoides syndrome (malignant cutaneous reticulosis syndrome) - keratoconjunctivitis, retinal edema, retinal hemorrhage
66. Myotonic dystrophy syndrome- corneal epithelial dystrophy, loss of corneal sensitivity, tapetoretinal degeneration, macular red spot, macular degeneration, chorioretinitis
*67. Neurofibromatosis (von Recklinghausen syndrome)- nodular swelling nerves, hamartoma of retina
68. Norrie disease (atrophia oculi congenita)- malformation of sensory cells of retina, corneal nebulae
69. Oculodental syndrome (Peters syndrome)- corneoscleral staphyloma, megalocornea, corneal marginal opacities, macular pigmentation
70. Onchoerciasis syndrome- punctate keratitis, sclerosing keratitis, chorioretinitis, retinal degeneration
71. Paget syndrome (osteitis deformans)- corneal ring opacities, retinal hemorrhages, pigmentary retinopathy, macular changes resembling Kuhnt-Junius degeneration
72. Phenothiazine-epithelial and endothelial pigment, retinal pigmentation
73. Pierre Robin syndrome (micrognathia-glossoptosis)- retinal disinsertion, megalocornea
74. Plasma lecithin (cholesterol acyltransferase deficiency)- corneal stromal opacities, retinal hemorrhages
75. Porphyria cutanea tarda - keratitis, retinal hemorrhages, cotton-wool spots, macular edema
76. Postvaccinal ocular syndrome- corneal vesicles, and marginal ulcers, chorioretinitis, central serous retinopathy, central retinal vein thrombosis
77. Progressive systemic sclerosis- marginal corneal ulcers with cicatrization, cotton-wool spots, retinal hemorrhages
78. Radiation-corneal ulcer, punctate keratitis, keratoconjunctivitis sicca, retinal hemorrhage, macular degeneration, macular holes with vascularization
*79. Refsum syndrome (phytanic acid oxidase deficiency)- band keratopathy, retinitis pigmentosa
80. Relapsing fever- interstitial keratitis, retinal hemorrhage
81. Relapsing polychondritis- corneal ulcer, retinal detachment, retinal artery thrombosis, keratoconjunctivitis sicca
82. Renal failure- cotton-wool spots, band keratopathy
83. Rendu-Osler syndrome (hereditary hemorrhagic telangiectasis) - intermittent filamentary keratitis, small retinal angiomas, retinal hemorrhages
84. Retinal disinsertion syndrome- bilateral keratoconus, retinal detachment
*85. Retinoblastoma-corneal neovascularization, retinal tumor
86. Rothmund syndrome (telangiectasia-pigmentation cataract syndrome)- corneal lesions, retinal hyperpigmentation
87. Rubella syndrome (Gregg syndrome)- microcornea, pigmentary retinal changes
88. Sabin-Feldman syndrome-microcornea, chorioretinitis or atrophic degenerative chorioretinal changes
89. Sanfillipo-Good syndrome (mucopolysaccharidosis III)-slight narrowing of retinal vessels, acid mucopolysaccharide deposits in cornea.
*90. Schaumann syndrome (sarcoidosis syndrome)-mutton fat keratitic precipitates, keratitis sicca, band-shaped keratitis, inflammatory retinal exudates
91. Scheie syndrome (MPS I-S)-diffuse to marked corneal clouding, tapetoretinal degeneration
92. Schwartz syndrome (glaucoma associated with retinal detachment)-retinal detachment, microcornea
93. Shy-Gonatas syndrome (orthostatic hypotension syndrome)-keratopathy, corneal ulcer, lattice-like white opacities in the area of Bowman membrane, retinal pigmented degeneration
94. Smallpox-keratitis, congenital corneal clouding, chorioretinitis
95. Stannus cerebellar syndrome (riboflavin deficiency)-corneal vascularization, superficial diffuse keratitis, corneal opacities, brownish retinal patches
96. Stickler syndrome (hereditary progressive arthroophthalmopathy)-keratopathy, chorioretinal degeneration, total retinal detachment
97. Sturge-Weber syndrome (neurooculocutaneous angiomatosis)-retinal detachment, increased corneal diameter with cloudiness
*98. Syphilis (acquired lues)-keratitis, retinal hemorrhages, retinal proliferation
*99. Temporal arteritis syndrome (Hutchinson-Horton-Magath-Brown syndrome)-retinal detachments, narrowing of retinal vessels, central retinal artery occlusion, corneal hypesthesia
100. Trisomy 13 (Patau syndrome)-malformed cornea, retinal dysplasia
*101. Tuberculosis-keratitis, pannus, corneal ulcer, retinitis
102. Ullrich syndrome (dyscraniopygophalangy)-cloudy cornea, corneal ulcers, chorioretinal coloboma
103. Ultraviolet radiation-photokeratitis, band keratopathy, herpes simplex keratitis, recurrent corneal erosions, retinal degeneration
104. Vaccinia-keratitis, pannus, corneal perforation, central serous retinopathy, pseudoretinitis pigmentosa
105. van Bogaert-Scherer-Epstein (primary hyperlipidemia)-arcus juvenilis of the cornea, lipid keratopathy, retinopathy with yellowish deposits
106. Vitreous tug syndrome-vitreous strands attached to corneal wound or scar, circumscribed retinal edema, posterior retinal detachment
107. von Gierke disease (glycogen storage disease type I)-corneal clouding, discrete nonelevated, yellow flecks in macula
108. Waardenburg syndrome (embryonic fixation syndrome)-microcornea, cornea plana, hypopigmentation and hypoplasia of retina
109. Wagner syndrome (hyaloideoretinal degeneration)-corneal degeneration, band-shaped keratopathy, hyaloideoretinal degeneration, narrowing of retinal vessels, retinal detachment, avascular preretinal membranes
110. Waldenström syndrome (macroglobulinemia syndrome)-crystalline corneal deposits, keratoconjunctivitis sicca, retinal venous thrombosis, retinal microaneurysms, cotton-wool spots
111. Weil disease (leptospirosis)-keratitis, retinitis
112. Werner syndrome (progeria of adults)-bullous keratitis, paramacular retinal degeneration
113. Wiskott-Aldrich syndrome (sex-linked draining ears, eczematoid dermatitis, bloody diarrhea)-corneal ulcers, retinal hemorrhages
114. Yersiniosis-corneal ulcer, retinal hemorrhages
115. Zellweger syndrome (cerebrohepatorenal syndrome)-corneal opacities, narrowing of retinal vessels, retinal holes without detachment, tapetoretinal degeneration
116. Zieve syndrome (hyperlipemia hemolytic anemia-icterus syndrome)-cloudy cornea, corneal ulcers, retinal lipemia


**RETINAL LESIONS ASSOCIATED WITH DEAFNESS**

1. Alport syndrome (hereditary familial congenital hemorrhagic nephritis)
2. Alström disease-retinitis pigmentosa
3. Choroideremia, obesity, and congenital deafness
4. Cockayne syndrome (dwarfism with retinal atrophy and deafness)
5. Dialinas-Amalric syndrome (deaf-mutism-retinal degeneration syndrome)
7. Hallgren syndrome (retinitis pigmentosa-deafness-ataxia syndrome)
8. Hunter syndrome (MPS II)
9. Laurence-Moon-Bardet-Biedl syndrome (retinitis pigmentosa-polydactyly adiposogenital syndrome)
10. Norrie disease-mental retardation, X-linked retinal malformation, and hearing loss
*11. Refsum syndrome (phytanic acid storage disease)
12. Retinal vessel changes, muscular dystrophy, mental retardation, and hearing loss
13. Rubella (German measles)-cardiac disorders, cataract, salt-and-pepper pigmentation
14. Sanfilippo syndrome (MPS III autosomal recessive)
*15. Syphilis-acquired or congenital
16. Usher syndrome (hereditary retinitis pigmentosa-deafness syndrome)
17. Waardenburg syndrome (embryonic fixation syndrome)


**SUBRETINAL FIBROSIS**

1. Central serous chorioretinopathy
2. Long-standing rhegmatogenous retinal detachment
3. Proliferative vitreoretinopathy
4. Vogt-Koyanagi-Harada disease


**EPIRETINAL MEMBRANES-MEMBRANES THAT GROW ON THE INNER SURFACE OF THE RETINA**

*1. After retinal photocoagulation, cryotherapy, or reattachment of retina
2. Following blunt or penetrating injuries
*3. Idiopathic
4. Nonproliferative retinal vascular disorders
*5. Proliferative retinopathies
6. Rhegmatogenous retinal detachment (see p. 468)
7. Sickle cell disease, including sickle cell C, sickle cell S, and sickle cell B with thalassemia
8. Vitreous hemorrhage (see p. 424)


**LINEAR STREAKS PATTERN IN FUNDUS**

*1. Angioid streaks (see p. 526-529)
2. Bird-shot retinochoroidopathy
3. Choroidal rupture
4. Demarcation lines
*5. Presumed ocular histoplasmosis syndrome-peripheral, parallel to equator
6. Migrating parasites
   A. Botfly larvae
   B. Trematodes
*7. Retinal and choroidal detachment
*8. Snail-track configuration of lattice degeneration

**YELLOW-ORANGE LESIONS OF SUBRETINAL FUNDUS**

1. Acute inflammatory lesions of pigment epithelium, choriocapillaris, and choroid
2. Detachment of retinal pigment epithelium
3. Isolated pocket of subretinal fluid
4. Subretinal fluid following scleral buckling procedure


**TALC RETINOPATHY-DRUG ADDICTS WHO INJECT DRUGS INTRAVENOUSLY**

1. Optic disc neovascularization (see p. 562-563)
2. Peripheral retinal neovascularization (see p. 514)
3. Vitreous hemorrhage (see p. 424)


**CRYSTALLINE RETINOPATHY**

1. Bietti crystalline dystrophy (Bietti disease)
2. Chronic retinal detachment
3. Cystinosis (cystine storage-aminoaciduria-dwarfism syndrome)
4. Gyrate atrophy with hyperornithemia (ornithine ketoacid aminotransferase deficiency)
5. Hyperoxaluria (oxalosis)
6. Nitrofurantoin therapy
7. Retinal pathology
8. Sjögren-Larson syndrome (oligophrenia-ichthyosis-spastic diplegia syndrome)
9. Talc emboli
10. Tamoxifen retinopathy


**PULFRICH STEREO-ILLUSION PHENOMENON**

This condition involves central serous elevation of the macula with abnormal latency of the visual-evoked potential.

* 1. Optic nerve disease - demyelinating optic neuropathy
  2. Media opacity
  3. Anisocoria
  4. Macular disease


**PARAFOVEAL TELANGIECTASIA**

This type of retinal microvascular anomaly involves the parafoveal capillary network as well as immediately adjacent vascular bed and is best demonstrated by fluorescein angiography.

* 1. Carotid artery obstruction
* 2. Diabetes mellitus usually bilateral
* 3. Idiopathic
  4. Localized form of Coats disease, usually unilateral
  5. Small-branch venular occlusion
  6. Small retinal capillary hemangioma, usually unilateral
  7. Roentgenogram, irradiation


**HEREDITARY PEDIATRIC RETINAL DEGENERATIONS**

1. Acquired
   A. Juvenile retinitis pigmentosa
   B. Early onset retinitis pigmentosa
      (1) Autosomal dominant
      (2) Autosomal recessive
      (3) X-linked recessive

2. Congenital
A. Complicated Leber congenital amaurosis
   (1) Multiple neurologic abnormalities
   (2) Others
   (3) Saldino-Mainzer syndrome
   (4) Senior-Loken syndrome (tubulointerstitial nephropathy syndrome)
   (5) Zellweger syndrome (cerebrohepatorenal syndrome of Zellweger)

B. Uncomplicated Leber congenital amaurosis


**RETICULAR PATTERN OF DARK LINES IN FUNDUS**

1. Granular pigmentary pattern of the peripheral fundus
2. Multiple drusen of peripheral fundus
3. Reticular degeneration of the pigment epithelium (peripheral)
4. Reticular pattern dystrophy of posterior fundus (Sjögren reticular dystrophy, Mesker macroreticular dystrophy, pattern dystrophy of the retinal pigment epithelium, Doyne honeycomb reticular degeneration)
5. Tapetochoroidal hypopigmentation


**RETINAL PIGMENT EPITHELIAL TEARS**

This condition involves a flat, uniform, crescent-shaped area of exposed choroid of pigment epithelial elevation.

1. Acute retinal necrosis
   *2. After laser photocoagulation
   3. Along margin of retinal detachment
   *4. Associated with pigment epithelial detachments
   5. Spontaneous
   6. Trauma


**RETINAL PIGMENT EPITHELIAL FOLDS**

1. Choroidal folds (see p, 530)
2. Pigment epithelial detachment
3. Retinal pigment epithelial tears (rips)
4. Retinal striae
*5. Subretinal neovascularization


**MIZUO PHENOMENON**

This condition involves a change of color of the fundus from red in the dark-adapted state to golden immediately or shortly after the onset of light.

1. Oguchi disease
2. X-liked juvenile retinoschisis
3. X-linked recessive cone dystrophy


**WHITE-DOT FOVEA**

This is a ring-like lesion in the macula with numerous confluent white dots arranged in a ring around the foveal margin,

1. Crystalline retinopathies
2. Epiretinal membrane with pseudohole
3. Gunn dots
4. Macular halo syndrome
5. Macular hole
6. Niemann-Pick disease
7. Vitreomacular fraction syndrome
ANGIOID STREAKS

Angioid streaks are ruptures of Bruch membrane characterized ophthalmoscopically by brownish lines surrounding the disc and radiating toward the periphery.

1. AC hemoglobinopathy
2. Acanthocytosis (abetalipoproteinemia, Bassen-Kornzweeg syndrome)
3. Acromegaly
4. Acquired hemolytic anemia
5. Beta thalassemia minor
6. Calciosis
7. Chronic congenital idiopathic hyperphosphatasemia
8. Chronic familial hyperphosphatemia
9. Cardiovascular disease with hypertension
10. Cooley anemia
11. Diffuse lipomatosis
12. Dwarfism
13. Epilepsy
14. Facial angiomatosis
15. Fibrodyplasia hyperelastica (Ehlers-Danlos syndrome)
16. François dyscephalic syndrome (Hallermann-Streiff syndrome)
17. Hemochromatosis
18. Hereditary spherocytosis
19. Hypercalcinosis
20. Idiopathic thrombocytic purpura
21. Lead poisoning
22. Myopia
23. Neurofibromatosis
24. Ocular melanocytosis
25. Optic disc drusen
26. Osteitis deformans (Paget disease)
27. Pituitary tumor
28. Previous choroidal detachment
*29. Pseudoxanthoma elasticum (Grönblad-Strandberg syndrome)
30. Senile (actinic) elastosis of the skin
31. Sickle cell disease (Herrick syndrome)
32. Sturge-Weber syndrome
33. Trauma
34. Tuberous sclerosis
35. Thrombocytopenic purpura


Extracted Table Angioid Streaks

<table>
<thead>
<tr>
<th>CHOROIDAL FOLDS</th>
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<tbody>
<tr>
<td>Choroidal folds are folds of the posterior pole, at the level of the choroid, with Hruby lens and pattern of alternating light lines on fluorescein angiography.</td>
</tr>
</tbody>
</table>

1. Choroidal tumor, such as a melanoma
2. Disciform degeneration
3. Exophthalmos
4. Graves disease (Basedow syndrome)
5. High hyperopia
6. Idiopathic-no underlying pathologic state
7. Infection of paranasal sinuses
8. Long-standing orbital inflammation
9. Massive cranioorbital hemangiopericytoma
*10. Ocular hypotony (see p. 325)
11. Orbital mass
12. Papilledema (see p. 593)
13. Posteriorly located choroidal detachment
14. Postoperative condition, such as scleral buckle
15. Primary retinal detachment
16. Subretinal neovascularization
17. Uveitis


**LESIONS CONFUSED WITH MALIGNANT MELANOMA**

1. Ciliary body and choroid
   A. Angioid streaks (see p. 526)
   B. Choroiditis
   C. Coats disease
   D. Detachment
   E. Leukemia and lymphoma
   F. Limited choroidal hemorrhage
   G. Lymphoid hyperplasia
   H. Nodular hyperplasia
   I. Sclerouveitis
   J. Tumors
      (1) Hemangioma
      (2) Melanocytoma
      (3) Meningioma
      (4) Metastatic carcinoma, including lung
      (5) Neurilemmoma
      (6) Neurofibroma
      (7) Neuroendocrine tumor
      (8) Nevus
      (9) Retinal oligodendrogliaoma
   K. Uveal effusion

2. Optic-nerve head
   A. Congenital crater
   B. Melanocytoma
3. Retina
   A. Chorioretinitis
   B. Ciliary body and choroid
   C. Disciform macular degeneration
   D. Foreign body
   E. Hemorrhagic macrocyst of retina
   F. Lesions of pigment epithelium
   G. Retinal detachment
      (1) Macular
      (2) More extensive
   H. Retinoschisis
4. Scleral thickening as amyloidosis
5. Vitreous body
   A. Abscess
   B. Hemorrhages


**CHOROIDAL HEMORRHAGE**

1. Acute choroiditis
2. After glaucoma filtering procedure (especially with Sturge-Weber syndrome)
3. Choroidal vascular aneurysm
4. Choroidal vascular sclerosis, such as senile macular degeneration with hemorrhage (disciform degeneration of the macula)
5. General diseases
   A. Arteriosclerosis
   B. Blood dyscrasias
      (1) Leukemia
      (2) Pernicious anemia
      (3) Purpura
      (4) Thrombocytopenia
   C. Diabetes mellitus (Willis disease)
   D. Ehlers-Danlos syndrome (fibrodysplasia elastica generalisata)
E. Paget disease (osteitis deformans)
F. Valsalva maneuver

6. Myopia-accompanied by choroidal atrophy
7. Papilledema-rare


**CHOROIDAL DETACHMENT**

Choroidal detachment can be differentiated from retinal detachment and tumor by its solid appearance, smooth surface, and appearance of normal retinal vessels with color unchanged and good transillumination.

*1. Acute ocular hypotony (see p. 325)*
   A. Myopia
   B. Operative or perforating wounds, including those required for surgical treatment of cataract, glaucoma, grafting of cornea, and retinal detachment
   C. Severe uveitis with severe visual loss, intense ocular pain, unusually low tension, and extremely deep anterior chamber in women
   D. Yttrium-aluminum-garnet (YAG) laser cyclophotocoagulation

2. Inflammatory disease
   A. Acute sinusitis
   B. Chronic cyclitis
   C. Harada disease (Vogt-Koyanagi-Harada syndrome)
   D. Orbital abscess
   E. Orbital pseudotumor
   F. Scleritis and tenonitis
   G. Sympathetic ophthalmia

3. Neoplastic disease
   A. Intraocular tumor, such as metastatic or malignant melanoma
   B. Leukemia
   C. Orbital tumor

4. Spontaneous detachment associated with uveal effusion, such as nonrhegmatogenous retinal detachment, shifting subretinal fluid, and peripheral annular choroidal detachment affecting males almost exclusively

5. Trauma
   A. Complication of scleral buckling retinal detachment surgery
   B. Contusion of globe without perforation
C. Following perforation injury, including that because of perforating corneal ulcer
D. Phthisical eye with traction of organized inflammatory tissue

6. Vascular disease
   A. Diabetes mellitus (Willis disease)
   B. Disseminated intravascular coagulation
   C. Hypertension
   D. Leukemia
   E. Multiple myeloma (Kahler disease)
   F. Nephritis
   G. Oral acetazolamide
   H. Periarteritis nodosa (Kussmaul disease)
   I. Syphilitic vascular disease
   J. Toxemia of pregnancy


**Extracted Table Choroidal detachment**

<table>
<thead>
<tr>
<th>CONDITIONS SIMULATING POSTERIOR UVEITIS OR CHOROIDITIS</th>
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<tbody>
<tr>
<td>1. Angioid streaks (see p. 526)</td>
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<tr>
<td>2. Central serous retinopathy</td>
</tr>
<tr>
<td>A. Central serous retinopathy and exudative chorioretinopathy associated with systemic vasculitis</td>
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<tr>
<td>B. Central serous retinopathy associated with crater-like holes in the optic disc</td>
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<tr>
<td>C. Choroidal</td>
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<td>D. Chorioretinal</td>
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<td>E. Retinal</td>
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<td>3. Chorioretinopathy with hereditary microcephaly</td>
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<td>4. Circinate retinopathy</td>
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<td>5. Congenital macular dysplasia</td>
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<tr>
<td>6. Doyne homogeneous retinal degeneration</td>
</tr>
<tr>
<td>7. Drug-induced macular disease</td>
</tr>
<tr>
<td>A. Chloroquine (Aralen)</td>
</tr>
<tr>
<td>B. Indomethacin (Indocin)</td>
</tr>
<tr>
<td>C. Thioridazine (Mellaril)</td>
</tr>
<tr>
<td>8. Drusen because of the following:</td>
</tr>
<tr>
<td>A. Disease (vascular, inflammatory, or neoplastic)</td>
</tr>
</tbody>
</table>
B. Heredity (primary degeneration)
C. Senility
9. Fundus flavimaculatus
10. Helicoid peripapillary chorioretinal degeneration
11. Hemangiom of the choroid
12. Idiopathic hyperlipemia
13. Ischemic ocular inflammation
14. Ischemic optic neuropathy (vascular pseudopapillitis)
15. Jensen disease (juxtapapillary retinopathy)
16. Macular degeneration
17. Malignant melanoma
18. Metastatic carcinoma
19. Night-blinding retinochoroidopathies
   A. Predominantly choroidal heredodegenerations
      (1) Choroidal sclerosis
      (2) Choroideremia
      (3) Fuchs spot
      (4) Gyrate atrophy of choroid
      (5) Myopic retinopathy and choroidopathy
   B. Predominantly tapetoretinal heredodegenerations
      (1) Retinitis pigmentosa group
      (2) Retinitis punctata albescens
20. Opacities of the macular retina
   A. Cotton-wool patches (see p. 491-494)
   B. Glial scars
   C. Hemorrhage
   D. Hemosiderin
   E. Inspissated exudates
   F. Lipoid deposits
   G. Pigment epithelium
      (1) Pigment epithelium migration
      (2) Pigment epithelium secretion
      (3) Pigment epithelium seeds
      (4) Proliferation in response to demand for phagocytes
      (5) Proliferation with formation of cuticular masses
      (6) Proliferation with metaplasia
      (7) Simple proliferation
21. Peripheral chorioretinal atrophy
22. Pigmentary perivenous-chorioretinal degeneration
23. Primary familial amyloidosis
24. Relapsing polychondritis
25. Retinal perforation during surgical treatment for strabismus
26. Retinal vasculitis
   A. Involvement of central retinal vein (papillophlebitis)
   B. Retinal periarteritis
   C. Retinal periphlebitis
27. Retinoblastoma
28. Sickle cell retinopathy
29. Solar bums
30. Sorsby pseudoinflammatory (hemorrhagic) macular degeneration
31. Vitreous hemorrhage (see p. 421-422)


**CHOROIDITIS (POSTERIOR UVEITIS)**

1. Anterior and posterior uveitis
   A. Herpes viruses
   B. Peripheral uveitis (cyclitis)
   C. Sarcoidosis syndrome (Schaumann syndrome)
   D. Syphilis (acquired lues)
   E. Toxoplasmosis
   F. Tuberculosis
   G. Unknown
   H. Vogt-Koyanagi-Harada syndrome (uveitis-vitiligo-aloepecia-poliosis syndrome)
2. Acquired immunodeficiency syndrome (AIDS)
3. Acute posterior multifocal placoid pigment epitheliopathy
4. Behçet syndrome
5. Bird-shot choroidopathy
6. Candidiasis
7. Cat-scratch disease
8. Cryptococcosis
9. Cytomegalovirus inclusion disease
10. Coccidioidomycosis
11. Epstein-Barr virus
12. Lyme disease
13. Histoplasmosis
14. Multiple evanescent white dot syndrome
15. Multiple sclerosis
16. *Pneumocystis carinii*
17. Punctate inner choroidopathy
18. Sarcoidosis syndrome (Schaumann syndrome)
19. Serpiginous choroidopathy
20. Syphilis (acquired lues)
21. Systemic lupus erythematosus
22. Toxocariasis
23. Toxoplasmosis
24. Unknown
25. Varicella zoster


**CONDITIONS SIMULATING POSTERIOR UVEITIS IN CHILDREN**

1. Coats syndrome (retinal telangiectasia)
2. Cockayne disease (dwarfism with retinal atrophy and deafness)
3. Cystinosis syndrome (Lignac-Fanconi syndrome)
4. Hypogammaglobulinemia
5. Idiopathic hyperlipemia
6. Leukemia
7. Massive retinal fibrosis
   *8. Retinoblastoma


**CHOROIDITIS (POSTERIOR UVEITIS) IN CHILDREN**

1. Ankylosing spondylitis
2. Anterior and posterior uveitis
   A. Sarcoidosis syndrome (Schaumann syndrome)
   B. Sympathetic ophthalmia
   C. Vogt-Koyanagi-Harada syndrome
3. Arteritis
4. Behçet disease (dermatostomatoophthalmic syndrome)
5. Chorioretinitis of unknown cause
   A. Disseminated chorioretinitis
   B. Juxtapapillary chorioretinitis
6. Cytomegalovirus inclusion disease (cytomegalovirus)
7. Diffuse unilateral subacute neuroretinitis
8. Herpes simplex chorioretinitis
9. Human immunodeficiency virus retinopathy
10. Inability of leukocytes to kill microorganisms
SYNDROMES AND DISEASES ASSOCIATED WITH UVEITIS

1. Arthralgia
   A. Hilding syndrome (destructive iridocyclitis and multiple joint dislocations)
   B. Histoplasmosis
   C. Whipple disease (intestinal lipodystrophy)

2. Arthritis
   A. Ankylosing spondylitis
   B. Behçet syndrome (dermatostomatoophthalmic syndrome)
   C. Bleu syndrome ""
   D. Familial histiocytic dermatoarthritis syndrome
   E. Felty syndrome (rheumatoid arthritis with hypersplenism)
   F. Gonorrhea
   G. Juvenile rheumatoid arthritis (Still disease)
   H. Leprosy (Hansen disease)
   I. Mucocutaneous lymph node syndrome
   J. Progressive systemic sclerosis
   K. Psoriatic arthritis
   L. Reiter syndrome (polyarthritis enteric a)
   M. Rheumatoid arthritis (adult)
   N. Sarcoidosis syndrome (Schaumann syndrome)
   O. Sporotrichosis
   P. Ulcerative colitis
   Q. Van Metre peripheral polyarthritis or mono arthritis
   R. Whipple disease (intestinal lipodystrophy)
   S. Yersiniosis
3. Cataract
   A. Acrodermatitis chronic atrophicans
   B. Andogsky syndrome (dermatogenous cataract)
   C. Anterior segment ischemia syndrome
   D. Arteriovenous fistula
   E. Atopic dermatitis
   F. Carotid artery syndrome (carotid vascular insufficiency syndrome)
   G. Cerebral palsy
   H. Chickenpox
   I. Cockayne syndrome (dwarfism with retinal atrophy and deafness)
   J. Cytomegalic inclusion disease
   K. Electrical injury
   L. Familial histiocytic dermatoarthritis syndrome
   M. Hallermann-Streiff-François syndrome (dyscephalic mandibulooculofacial syndrome)
   N. Herpes simplex
   O. Herpes zoster
   P. Hilding syndrome (destructive iridocyclitis and multiple joint dislocations)
   Q. Histiocytosis X (Hand-Schüller-Christian syndrome)
   R. Hodgkin disease (lymph node disease)
   S. Homocystinuria syndrome
   T. Hypervitaminosis D
   U. Influenza
   V. Juvenile rheumatoid arthritis (Still disease)
   W. Kussmaul disease (periarteritis nodosa)
   X. Leptospirosis (Weil disease)
   Y. Listerellosis
   Z. Malaria
   AA. Measles (rubeola)
   BB. Moniliasis (idiopathic hypoparathyroidism)
   CC. Myotonic dystrophy syndrome
   DD. Oculootoororenoerythropoietic disease
   EE. Passow syndrome (syringomyelia)
   FF. Radiation
   GG. Relapsing polychondritis
   HH. Rubella syndrome
   II. Sarcoidosis syndrome (Schaumann syndrome)
   JJ. Stevens-Johnson syndrome (erythema multiforme exudativum)
   KK. Stickler syndrome (hereditary progressive arthroophthalmyopathy)
   LL. Toxocariasis (visceral larva migrans syndrome)
   MM. Toxoplasmosis
   NN. Werner syndrome (progeria of adults)
   OO. Yersiniosis

4. Conjunctivitis
   A. Acanthamoeba
   B. Actinomycosis
C. African eyeworm disease
D. Amebiasis
E. Andogsky syndrome (dermatogenous cataract)
F. Angular conjunctivitis
G. Ascariasis
H. Atopic dermatitis
I. Bacillary dysentery (shigellosis)
J. Bacterial endocarditis
K. Behçet syndrome (dermatostomato-ophthalmic syndrome)
L. Boutonneuse fever (rickettsia, Marseilles fever)
M. Brucellosis
N. Candidiasis
O. Charlin syndrome (nasal nerve syndrome)
P. Chlamydia
Q. Coccidioidomycosis
R. Coenurusis
S. Cogan syndrome (non syphilitic interstitial keratitis)
T. Crohn disease (granulomatous ileocolitis)
U. Cytomegalic inclusion disease
V. Disseminated lupus erythematosus (Kaposi-Libman-Sacks syndrome)
W. Epidermic keratoconjunctivitis
X. *Escherichia coli*
Y. Felty syndrome (rheumatoid arthritis with hypersplenism)
Z. *Haemophilus aegyptius*
AA. Herpes simplex
BB. Herpes zoster
CC. Hodgkin disease
DD. Infectious mononucleosis
EE. Influenza
FF. Kussmaul disease (periarteritis nodosa)
GG. Leptospirosis (Weil disease)
HH. Listerellosis
II. Lymphogranuloma venereum
JJ. Measles (rubeola)
KK. Meningococcemia
LL. Metastatic bacterial endophthalmitis
MM. Mikulicz-Radecki syndrome (dacryosialoadenopathy)
NN. Moniliasis (idiopathic hypoparathyroidism)
OO. Moraxella lacunata
PP. Mucocutaneous lymph node syndrome
QQ. Mumps
RR. Mycosis fungoides syndrome (Sézary syndrome)
SS. Nocardiosis
TT. Ocular vaccinia
UU. Pneumococcus
VV. Polymysitis dermatomyositis
WW. Progressive systemic sclerosis (scleroderma)
XX. Psoriatic arthritis
YY. Q fever
ZZ. Radiation
AAA. Reiter syndrome (polyarthritis enterica)
BBB. Relapsing fever
CCC. Rocky Mountain spotted fever
DDD. Rubella syndrome
EEE. St. Anthony fire (erysipelas)
FFF. Seborrheic dermatitis
GGG. Sporotrichosis
HHH. Staphylococcus
III. Stevens-Johnson syndrome (erythema multiforme exudativum)
JJJ. Streptococcus
KKK. Syphilis (acquired lues)
LLL. Trichinellosis
MMM. Tuberculosis
NNN. Vaccinia
OOO. Xeroderma pigmentosum
PPP. Yersiniosis

5. Cornea
A. Acanthamoeba
B. Acrodermatitis chronic atrophicans
C. Actinomycosis
D. African eyeworm disease
E. Andogsky syndrome (dermatogenous cataract)
F. Angioedema (Quincke disease)
G. Angular conjunctivitis (Morax-Axenfeld bacillus)
H. Ankylosing spondylitis
I. Anterior segment ischemia syndrome
J. Arteriovenous fistula
K. Atopic dermatitis
L. Bee sting of the cornea
M. Behçet syndrome (dermatostomatoophthalmic syndrome)
N. Brucellosis (Bang disease)
O. Candidiasis
P. Charlin syndrome (nasal nerve syndrome)
Q. Chickenpox
R. Chlamydia
S. Cockayne syndrome (dwarfism with retinal atrophy and deafness)
T. Cogan syndrome (I) (nonsyphilitic interstitial keratitis)
U. Crohn disease (granulomatous ileocolitis)
V. Cystinosis syndrome
W. Cytomegalic inclusion disease
X. Disseminated lupus erythematosus (Kaposi-Libman-Sacks syndrome)
Y. Electrical injury
Z. Epidemic keratoconjunctivitis
AA. E. coli
BB. Felty syndrome (rheumatoid arthritis with hypersplenism)
CC. Gonorrhea
DD. H. aegyptius
EE. Herpes simplex
FF. Herpes zoster
GG. Hilding syndrome (destructive iridocyclitis and multiple joint dislocations)
HH. Histocytosis X (Hand-Schüller-Christian syndrome)
II. Hodgkin disease (lymph node disease)
JJ. Homocystinuria syndrome
KK. Hypervitaminosis D
LL. Hypothermal injury
MM. Influenza
NN. Japanese River fever (mite borne typhus)
OO. Juvenile rheumatoid arthritis (Still disease)
PP. Juvenile xanthogranuloma (nevoxanthoendothelioma)
QQ. Kussmaul disease (periarteritis nodosa)
RR. Leprosy (Hansen disease)
SS. Leptospirosis (Weil disease)
TT. Lewis syndrome (tuberoeruginous syphilid of Lewis)
UU. Listerellosis
VV. Lockjaw
WW. Lymphogranuloma venereum
XX. Malaria
YY. Measles (rubeola)
ZZ. Meningococcemia
AAA. Mikulicz-Radecki syndrome (dacryosialoadenopathy)
BBB. Moniliasis-idiopathic hypoparathyroidism (Addison syndrome)
CCC. Moraxella lacunata
DDD. Mumps
EEE. Mycosis fungoides syndrome (Sézary syndrome)
FFF. Myotonic dystrophy syndrome
GGG. Nocardiosis
HHH. Ocular vaccinia
III. Onchocerciasis syndrome
JJJ. Passow syndrome (status dysraphicus)
KKK. Plague
LLL. Pneumococcus
MMM. Postvaccinial ocular syndrome
NNN. Progressive systemic sclerosis (scleroderma)
OOO. Psoriasis
PPP. Psoriatic arthritis
QQQ. Radiation
RRR. Reiter syndrome (polyarthritis enterica)
SSS. Relapsing fever
TTT. Relapsing polychondritis
UUD. Rheumatoid arthritis (adult)
VVV. Rubella syndrome
WWW. St. Anthony fire (erysipelas)
XXX. Sarcoidosis syndrome (Schaumann syndrome)
YYY. Seborrheic dermatitis
ZZZ. Sporotrichosis
AAAA. Staphylococcus
BBBB. Stevens-Johnson syndrome (erythema multiforme exudativum)
CCCC. Stickler syndrome (hereditary progressive arthroophthalmopathy)
DDDD. Streptococcus
EEEE. Syphilis
FFFF. Thelaziasis
GGGG. Toxoplasmosis
HHHH. Tuberculosis
IIII. Vaccinia
JJJJ. Werner syndrome (progeria of adults)
KKKK. Xeroderma pigmentosum
LLLL. Yersiniosis

Diarrhea
A. Amebiasis
B. Bacillary dysentery
C. Chlamydia
D. Crohn disease (granulomatous ileocolitis)
E. E. coli
F. Listerellosis
G. Psoriatic arthritis
H. Regional enteritis (ulcerative colitis)
I. Rubella syndrome
J. Staphylococcus
K. Ulcerative colitis (regional enteritis)
L. Whipple disease (intestinal lipodystrophy)
M. Yersiniosis

Disc neovascularization
A. Ischemic uveitis of Knox
B. Papillophlebitis

Exophthalmus (proptosis)
A. Actinomycosis
B. Angioedema
C. Arteriovenous fistula
D. Coenurosis
E. Cryptococcosis
F. Disseminated lupus erythematosus (Kaposi-Libman-Sacks syndrome)
G. Hallermann-Streiff-François syndrome (dyscaphalic mandibulooculofacial syndrome)
H. Herpes zoster
I. Histiocytosis X (Hand-Schüller-Christian syndrome)
J. Juvenile xanthogranuloma (nevoxanthoendothelioma)
K. Kussmaul disease (periarteritis nodosa)
L. Mumps
M. Polymyositis dermatomyositis
N. Relapsing polychondritis
O. Streptococcus
P. Trichinellosis
Q. Werner syndrome (progeria of adults)

9. Exudative detachment
A. Acute retinal necrosis syndrome
B. Bacterial endocarditis
C. Boutonneuse fever (rickettsia, Marseilles fever)
D. Cryptococcosis
E. Histiocytosis X (Hand-Schüller-Christian syndrome)
F. Japanese River fever (typhus)
G. Kussmaul disease (periarteritis nodosa)
H. Mycosis fungoides syndrome
I. Oculootoororenoerythropoietic disease
J. Pappataci fever (sandfly fever)
K. Periocular and ocular metastatic tumors
L. Progressive systemic sclerosis (scleroderma)
M. Rheumatic fever
N. Rocky Mountain spotted fever
O. Sarcoidosis syndrome (Schaumann syndrome)
P. Schwartz syndrome (glaucoma associated with retinal detachment)
Q. Stickler syndrome (hereditary progressive arthroophthalmopathy)
R. Toxocariasis (visceral larva migrans syndrome)
S. Toxoplasmosis
T. Weber-Christian syndrome (subcutaneous inflammatory lesions)

10. Glaucoma
A. Acanthamoeba
B. Angioedema
C. Arteriovenous fistula
D. Acarasis
E. Atopic dermatitis
F. Behçet syndrome (dermatostomatoophthalmic syndrome)
G. Brucellosis (late manifestation) (Bang disease)
H. Carotid artery syndrome
I. Coats disease (retinal telangiectasia)
J. Coccidioidomycosis
K. Coenurosis
L. Electrical injury
M. *E. coli*
N. Familial histiocytic dermatoadthritis syndrome
O. Hallermann-Streiff-François syndrome (dyscephalic mandibulooculofacial syndrome)
P. Homocystinuria syndrome
Q. Juvenile rheumatoid arthritis (Still disease)
R. Juvenile xanthogranuloma (nevoxanthoendothelioma)
S. Leptosy (Hansen disease)
T. Listerellosis
U. Measles (rubeola)
V. Oculootoororenoerythropoietic disease
W. Onchocerciasis syndrome
X. Periocular and ocular metastatic tumors
Y. Pneumococcus
Z. Posner-Schlossman syndrome (glaucomatocyclitic crisis)
AA. Pseudouveitis, glaucoma, hyphema syndrome (PUGH syndrome)
BB. Radiation
CC. Relapsing polychondritis
DD. Rubella syndrome
EE. Sarcoidosis syndrome (Schaumann syndrome)
FF. Schwartz syndrome (glaucoma associated with retinal detachment)
GG. Staphylococcus
HH. Stickler syndrome (hereditary progressive arthro-ophthalmopathy)
II. Streptococcus
JJ. Trichinellosis
KK. Uveitis, glaucoma, hyphema syndrome (UGH syndrome)
LL. Weber-Christian syndrome (subcutaneous inflammatory lesions)

11. Hepatomegaly
   A. Cytomegalic inclusions disease
   B. Toxocariasis (visceral larva migrans syndrome)
   C. Toxoplasmosis

12. Influenza-like disease
   A. Acanthamoeba
   B. Amebiasis
   C. Bacillary dysentery
   D. Boutonneuse fever (rickettsia, Marseilles fever)
   E. Brucellosis (Bang disease)
   F. Chickenpox
   G. Chlamydia
   H. E. coli
   I. Gonorrhea
   J. H. aegyptius
   K. Histoplasmosis
   L. Influenza
   M. Japanese River fever (mite-borne typhus)
   N. Leptospirosis (Weil disease)
   O. Malaria
   P. Measles (rubeola)
Q. Meningococcemia
R. Mumps
S. Pappataci fever (sandfly fever)
T. Plague (bubonic plague)
U. Pneumococcus
V. Q fever
W. Relapsing fever
X. Rocky Mountain spotted fever
Y. Staphylococcus
Z. Streptococcus
AA. Toxoplasmosis
BB. Tuberculosis

13. Iris neovascularization, such as Knox ischemic ocular inflammation (rubeosis iridis)
14. Jaundice
   A. Leptospirosis
15. Meningism (meningitis)
   A. Acanthamoeba
   B. Behçet syndrome (dermatostomato-ophthalmic syndrome)
   C. Cryptococcosis
   D. Gonorrhea
   E. Herpes simplex
   F. Histoplasmosis
   G. Leptospirosis (Weil disease)
   H. Listerellosis
   I. Meningococcemia
   J. Sympathetic ophthalmia
   K. Toxoplasmosis
   L. Tuberculosis

16. Microphthalmia
   A. Cytomegalic inclusion disease
   B. Hallermann-Streiff-François syndrome (dyscephalic mandibulooculofacial syndrome)
   C. Mumps
   D. Rubella syndrome
   E. Toxoplasmosis

17. Nodules in the leg
   A. Histoplasmosis
   B. Sarcoidosis syndrome (Schaumann syndrome)
   C. Ulcerative colitis

18. Optic neuritis (papillitis)
   A. Angioedema
   B. Behçet syndrome (dermatostomatoophthalmic syndrome)
   C. Boutonneuse fever (rickettsia, Marseilles fever)
   D. Brucellosis (Bang disease)
   E. Candidiasis
   F. Chickenpox
G. Cytomegalic inclusion disease
H. Disseminated lupus erythematosus (Kaposi-Libman-Sacks syndrome)
I. Electrical injury
J. Felty syndrome (rheumatoid arthritis with hypersplenism)
K. Herpes zoster
L. Hodgkin disease (lymph node disease)
M. Infectious mononucleosis
N. Influenza
O. Juvenile rheumatoid arthritis (Still disease)
P. Leptospirosis
Q. Malaria
R. Measles
S. Meningococcemia
T. Mikulicz-Radecki syndrome (dacryosialoadenopathy)
U. Mumps
V. Ocular vaccinia
w. Onchocerciasis syndrome
X. Pappataci fever (sandfly fever)
Y. Postvaccinial ocular syndrome
Z. Q fever
AA. Reiter syndrome (polyarthritis enteric a)
BB. Regional enteritis (ulcerative colitis)
CC. Rocky Mountain spotted fever
DD. Sarcoidosis syndrome (Schaumann syndrome)
EE. Stevens-Johnson syndrome (erythema multiforme exudativum)
FF. Streptococcus
GG. Sympathetic ophthalmia
HH. Syphilis (acquired lues)
II. Toxocariasis (visceral larva migrans syndrome)
JJ. Toxoplasmosis
KK. Trichinellosis
LL. Tuberculosis
MM. Vaccinia

19. Papilledema
A. Angioedema
B. Arteriovenous fistula
C. Acariasis
D. Bacterial endocarditis
E. Behçet syndrome (dermatostomatoophthalmic syndrome)
F. Brucellosis (Bang disease)
G. Chickenpox
H. Coccidioidomycosis
I. Cryptococcosis
J. Cysticercosis
K. Disseminated lupus erythematosus (Kaposi-Libman-Sacks syndrome)
L. Histiocytosis X (Hand-Schüller-Christian syndrome)
M. Hodgkin disease (lymph node disease)
N. Hypervitaminosis D
O. Malaria
P. Moniliasis-idiopathic hypoparathyroidism (Addison disease)
Q. Mycosis fungoides syndrome (Sézary syndrome)
R. Papillophlebitis
S. Pappataci fever (sandfly fever)
T. Passow syndrome (status dysraphicus)
U. Periocular and ocular metastatic tumors
V. Progressive systemic sclerosis
W. Sarcoidosis syndrome (Schaumann syndrome)
X. Syphilis
Y. Trichinellois
Z. Whipple disease (intestinal lipodystrophy)

20. Paralysis of extraocular muscle
A. African eyeworm disease
B. Arteriovenous fistula
C. Bacterial endocarditis
D. Brucellosis
E. Cerebral palsy
F. Chickenpox
G. Coccidioidomycosis
H. Cysticercosis
I. Disseminated lupus erythematosus
J. Electrical injury
K. Herpes simplex
L. Herpes zoster
M. Hodgkin disease (lymph node disease)
N. Hypothermal injury
O. Infectious mononucleosis
P. Influenza
Q. Kussmaul disease (periarteritis nodosa)
R. Lockjaw (tetanus)
S. Malaria
T. Measles (rubeola)
U. Meningococcemia
V. Multiple sclerosis
W. Mumps
X. Ocular vaccinia
Y. Passow syndrome (status dysraphicus)
Z. Periocular and ocular metastatic tumors
AA. Reiter syndrome (polyarthritis enterica)
BB. Relapsing fever
CC. Relapsing polychondritis
DD. Rocky Mountain spotted fever
EE. Streptococcus
FF. Syphilis (acquired lues)
GG. Trichinelllosis

21. Perivenous sheathing
   A. *Acanthamoeba*
   B. Amebiasis
   C. Boutonneuse fever (rickettsia, Marseilles fever)
   D. Brucellosis (Bang disease)
   E. Candidiasis
   F. Coccidioidomycosis
   G. Metastatic bacterial endophthalmitis
   H. Metastatic fungal endophthalmitis
   I. Multiple sclerosis
   J. Myotonic dystrophy syndrome
   K. Ocular vaccinia
   L. Onchocerciasis syndrome
   M. Plague
   N. Postvaccinial ocular syndrome
   O. Q fever
   P. Sarcoidosis syndrome (Schaumann syndrome)
   Q. Syphilis (acquired lues)
   R. Toxocariasis (visceral larva migrans syndrome)
   S. Toxoplasmosis
   T. Tuberculosis
   U. Vaccinia
   V. Weber-Christian syndrome (subcutaneous inflammatory lesions)

22. Pneumonitis
   A. *Chlamydia*
   B. Cytomegalic inclusion disease
   C. Plague (Bubonic plague)
   D. Pneumococcus
   E. Rubella syndrome
   F. Toxocariasis
   G. Whipple disease (intestinal lipodystrophy)

23. Prostatitis
   A. Gonococcosis
   B. Whipple disease (intestinal lipodystrophy)

24. Salt-and-pepper fundus
   A. Choroideremia in males
   B. Cockayne disease (dwarfism with retinal atrophy and deafness)
   C. Cystinosis
   D. Prenatal influenza
   E. Prenatal syphilis

25. Skin lesions
   A. Acrodermatitis chronica atrophicans
   B. African eyeworm disease
   C. Andogsky syndrome (dermatogenous cataract)
D. Angioedema
E. Atopic dermatitis
F. Behçet syndrome (dermatostomato-ophthalmic syndrome)
G. Chickenpox
H. Disseminated lupus erythematosus (Kaposi-Libman-Sacks syndrome)
I. Familial histiocytic dermatomyositis syndrome
J. Herpes simplex
K. Herpes zoster
L. Histiocytosis X (Hand-Schüller-Christian syndrome)
M. Juvenile xanthogranuloma (nevoxanthoendothelioma)
N. Leprosy (Hansen disease)
O. Lewis syndrome (tuberoserphiginous syphilid of Lewis)
P. Listerellosis
Q. Measles (rubeola)
R. Moraxella lacunata
S. Mucocutaneous lymph node syndrome
T. Mycosis fungoides syndrome (Sézary syndrome)
U. Polymyositis dermatomyositis
V. Postvaccinial ocular syndrome
W. Psoriasis
X. Psoriatic arthritis
Y. St. Anthony fire (erysipelas)
Z. Schistosomiasis (bilharziasis)
AA. Seborrheic dermatitis
BB. Sporotrichosis
CC. Staphylococcus

26. Stomatitis
   A. Behçet syndrome (dermatostomatoophthalmic syndrome)
   B. Disseminated systemic histoplasmosis-not the ocular form
   C. Herpes simplex
   D. Lewis syndrome (tuberoserphiginous syphilid of Lewis)
   E. Reiter syndrome (polyarthritis enteric a)
   F. Regional enteritis
   G. Ulcerative colitis

27. Tonsillitis
   A. Whipple disease (intestinal lipodystrophy)

28. Trauma (nonpenetrating)


CHORIoretinitis JUXtapatIllar is

This large, irregular opaque mass that protrudes three to four diopters and obscures the retinal vessels is seen near the disc and may be confused with acute optic neuritis or a tumor.

1. Acanthamoeba keratitis of fellow eye
2. Bird-shot retinochoroidopathy
3. Coccidioides immitis
4. Histoplasmosis
5. Sarcoïdosis syndrome (Schaumann syndrome)
6. Syphilis (acquired lues)
7. Toxoplasmosis
8. Tuberculosis


CHOROIDAL NEOVASCULARIZATION

Choroidal neovascularization comprises new vessel formation from choriocapillaris through a defect in the Bruch membrane as suggested by fluorescein angiography.

1. Choroidal neovascular ingrowth at the margin of the optic nerve head
   A. Angioid streaks (see p. 526)
   B. Hyaline bodies of optic nerve head
   C. Idiopathic choroidal neovascularization
   D. Macular drusen
   E. Multiple evanescent white-dot and acute idiopathic blind spot enlargement syndrome
   F. Optic-nerve coloboma
   G. Peripapillary choroiditis
   H. Presumed ocular histoplasmosis syndrome
   I. Pseudotumor cerebri (Symond syndrome)
   J. Serpiginous choroiditis
2. Choroidal neovascular ingrowth through breaks in Bruch membrane in the macular area
3. Acute posterior multifocal placoid pigment epitheliopathy
   A. Angioid streaks (see p. 526)
   B. Behçet syndrome (dermatostomatoophthalmic syndrome) or Best disease
   C. Bird-shot retinochoroidopathy
   D. Choroidal rupture
   E. Choroidal tumors
   F. Chronic uveitis
   G. Foveomacular dystrophy
   H. Fundus flavimaculatus
   I. Idiopathic choroidal neovascularization
   *J. Macular drusen
   K. Morning glory syndrome
   L. Myopic degeneration
   M. Parafoveal telangiectasis
   N. Photocoagulation of macular lesions with argon laser
   O. Presumed ocular histoplasmosis syndrome
   P. Osteogenesis imperfecta (van der Hoeve syndrome)
   Q. Retinitis pigmentosa
   R. Rubella syndrome (Gregg syndrome)
   S. Sarcoidosis syndrome (Schaumann syndrome)
   T. Scars from previous deep macular hemorrhage
   U. Senile disciform macular degeneration (Kuhnt-Junius disease)
   V. Serpiginous choroiditis
   W. Sorsby fundus dystrophy
   X. Tilted disc syndrome
   Y. Toxocariasis
   Z. Toxoplasma retinochoroiditis
   AA. Trauma
   BB. Vein occlusion
   CC. Vogt-Koyanagi-Harada syndrome (uveitis-vitiligo-alopecia-poliosis syndrome)


**ISCHEMIC INFARCTS OF CHOROID (ELSCHNIG SPOTS)**

When healed, these may show small, disseminated yellowish scars with central pigment deposits and may be associated with retinal separation when acute.

1. Chronic glomerulonephritis
2. Collagen disease, such as scleroderma
*3. Malignant hypertension
4. Toxemia of pregnancy


**CHORIORETINAL AND CHORIOVITREAL NEOVASCULARIZATION**

This type of new vessel formation from choroid into the retina or vitreous usually occurs after photocoagulation or after any of the following:

1. Atrophic scars in the presumed ocular histoplasmosis syndrome
2. Central serous chorioretinopathy
*3. Diabetes mellitus (Willis disease)
4. Diseases of the retinal pigment epithelium
5. Eales disease (periphlebitis)
6. Leber syndrome (optic atrophy-amaurosis pituitary syndrome)
7. Sarcoidosis syndrome (Schaumann syndrome)
8. Sickle cell disease (Herrick syndrome)


**UVEAL EFFUSION**

Uveal effusion involves leaking of fluid from the choriocapillaris into the choroid or subretinal space or both.
1. Hydrostatic
   A. Dural arteriovenous fistula
   *B. Hypotony, wound leak
   C. Nanophthalmos
2. Idiopathic
3. Inflammatory
   A. After panretinal photocoagulation
   B. HIV
   C. Scleritis, infected scleral buckle
   D. Systemic lupus erythematosus
   E. Trauma, intraocular surgery
   F. Uveitis, sympathetic ophthalmia, Harada disease


**CHOROID COLOBOMA**

1. Aicardi syndrome
2. Basal cell nevus syndrome (Gorlin syndrome)
3. Cat-eye syndrome (partial G-trisomy)
4. CHARGE association among coloboma, heart anomaly, choanal atresia, retardation, genital and ear anomalies
5. Doubtful association
   A. Crouzon syndrome (dysostosis craniofacialis)
   B. Ellis-Van Creveld syndrome (chondroectodermal dysplasia)
   C. Hallerman-Streiff syndrome (dyscephalic mandibulooculofacial)
   D. Incontinentia pigmenti I (Block-Sulzberger syndrome)
   E. Kartagener syndrome (bronchiectasis-dextrocardia-sinusitis)
   F. Laurence-Moon-Bardet-Biedl syndrome (retinitis pigmentosa-polydactyl-adiposogenital syndrome)
   G. Pierre Robin syndrome (micrognathia-glossoptosis syndrome)
   H. Stickler syndrome (hereditary progressive arthro-ophthalmopathy)
   I. Tuberous sclerosis (Bourneville syndrome)
6. Goldenhar syndrome (oculoauriculovertebral dysplasia)
7. Goltz syndrome (focal dermal hypoplasia syndrome)
8. Isolated, sporadic
9. Joubert syndrome with bilateral chorioretinal coloboma (coloboma, chorioretinal with cerebellar vermis aplasia)
10. Klinefelter syndrome (gynecomastia-aspermato genesis syndrome)
11. Lenz microphthalmia syndrome
12. Linear sebaceous nevus syndrome
13. Median facial cleft syndrome
14. Meckel syndrome (dysencephalia splanchnocystic syndrome)
15. Retinal astrocytoma
16. Retinal dysplasia
17. Retinoblastoma
18. Rubinstein-Taybi syndrome (broad-thumbs syndrome)
19. Triploidy
*20. Trisomy (Edward syndrome)
21. Trisomy (Patau syndrome)
22. Turner syndrome
23. Warburg syndrome
24. 13q syndrome


**CHOROIDAL ISCHEMIA**

This condition involves decreased choroidal perfusion demonstrated by fluorescein angiography.

1. Arteritic anterior ischemic optic neuropathy
2. Disseminated intravascular coagulation
*3. Hypertension, severe
4. Renal failure
5. Systemic lupus erythematosus
6. Toxemia of pregnancy
7. Thrombotic thrombocytopenic purpura


**PARS PLANITIS (PERIPHERAL UVEITIS)**

In pars planitis, inferior exudates in the peripheral retina, ora, pars plana, and peripheral vitreous, vitreous ray and cells, posterior cortical cataract, perivasculitis, partial thrombosis of central retinal vein, glaucoma, peripheral retinal hemorrhages, and retinal detachment may be present.
1. Dental infection
2. Hereditary
*3. Idiopathic
4. Multiple sclerosis (disseminated sclerosis)
5. Nematodiasis
6. Rheumatic disease
7. Sarcoidosis syndrome (Schaumann syndrome)
8. Sinus infection
9. Streptococcal hypersensitivity
10. Syphilis (acquired lues)
11. Toxoplasmosis
12. Ulcerative colitis (inflammatory bowel disease)


**Extracted Table Differential Diagnosis of Pars Planitis**
BLURRED OPTIC-NERVE HEADS

CILIOOPTIC VEIN

This vein appears at the disc edge and dips into the optic nerve to anastomose with branches of the central retinal vein

1. Congenital
2. Neurofibromatosis (von Recklinghausen syndrome)
3. Sturge-Weber syndrome (meningocutaneous syndrome)


DRUSEN OF OPTIC NERVE

These white or yellow conglomerate, translucent bodies in the optic nerve may cause field defects.
1. Alagille syndrome
2. Alport syndrome (hereditary nephritis)
3. Angioid streaks (pseudoxanthoma elasticum; Grönblad-Strandberg syndrome)
4. Associated with corneal dystrophy
5. Diabetes mellitus (Willis disease)
6. Friedreich ataxia
7. Glaucoma
8. Hereditary-autosomal dominant
9. High myopia
10. Idiopathic
11. Meningioma (unusual)
12. Pituitary tumor (unusual)
13. Retinal vein occlusion
14. Retinitis pigmentosa
15. Systemic lupus erythematosus (SLE)
16. Status dysrphicus syndrome (Passow syndrome)
17. Syphilis (acquired lues)
18. Tuberous sclerosis (Bourneville syndrome)
19. Wilson disease (hepatolenticular degeneration)


FLUID ENLARGEMENT OF RETROBULBAR OPTIC NERVE OR SHEATH (DEMONSTRATED BY COMPUTED TOMOGRAPHIC SCANNING AND ECHOGRAPHY)

1. Arachnoiditis
2. Basilar artery aneurysm
3. Bilateral temporal lobe cysts
4. Central retinal vein occlusion (see p. 468-472)
5. Facial trauma
6. Iliojejunal bypass
7. Occipital intradural arteriovenous malformation
8. Optic-nerve meningioma
9. Optic-nerve sheath cyst
10. Pseudotumor cerebri
11. Subdural hematoma  
12. Trauma (intraneural hemorrhage of optic nerve)  
13. Uveal meningeal syndrome


**HYPEREMIA OF OPTIC DISC**

1. Central retinal vein thrombosis (see p. 468-472)  
2. Hemangioma  
3. Hypermetropia  
4. Hypertensive retinopathy  
5. Ischemic optic neuropathy  
6. Neovascularization  
7. Optic neuritis (see p. 578)  
8. Papilledema (see p. 593)  
9. Polycythemia vera (Vaquez disease)  
10. von Hippel-Lindau disease (retinocerebral angiomatosis)


**ISCHEMIC OPTIC NEUROPATHY**

The anterior form is occlusive disease of the optic-nerve head and retrolaminar region of the optic nerve; the posterior form is occlusion of one or more nutrient arteries to the rest of the optic nerve. Onset is usually sudden, with painless unilateral visual loss and visual-field defect.

1. Compression  
   A. Orbital hemorrhage (trauma)  
   B. Thyroid disease (Graves disease)  
2. Drugs  
   A. Sumatriptan  
   B. Vigabatrin  
3. Systemic diseases (often in combination)  
   A. Arteriosclerosis  
   B. Arrhythmia  
   C. Atherosclerosis  
   D. Cerebrovascular disease  
   E. Diabetes mellitus  
   F. Gastrointestinal ulcer  
   G. Hypercholesterolemia  
   H. Hyperhomocystinemia  
   I. Hyperparathyroidism  
   J. Hypertension, nocturnal hypotension  
   K. Ischemic heart disease
L. Sickle cell disease
M. Takayasu disease

4. Vasculitis
   A. Allergic vasculitis
   B. Buerger disease
   C. Churg-Strauss syndrome (allergic granulomatosis and angiitis)
   D. Collagen diseases, including polyarteritis nodosum and systemic lupus
   E. Giant cell (temporal) arteritis
   F. Postimmunization
   G. Postviral vasculitis
   H. Syphilis

5. Miscellaneous
   A. Acute anemia
   B. Anemia combined with hypotension
   C. Carotid artery disease
   D. Fabry disease (angiokeratoma corporis diffusum)
   E. Hypertensive with peritoneal dialysis
   F. Low tension glaucoma
   G. Glucose-6-phosphate dehydrogenase deficiency
   H. Migraine
   I. Polycythemia vera
   J. Radiation
   K. Retinal artery occlusion
   L. Tobacco
   M. Various vascular disorders (e.g., cavernous sinus thrombosis)


**LINEAR HEMORRHAGE ON OPTIC DISC**

1. Diabetes mellitus
2. Drusen of optic nerve
3. Glaucoma
4. Ischemic optic neuropathy
   A. Altitudinal field loss
   B. Dense arcuate field loss
   C. Sector-shaped field loss
5. Isolated finding
6. Leukemia
7. Systemic hypertension


**NEOVASCULARIZATION OF OPTIC DISC (GROWTH OF BLOOD VESSELS ONTO OPTIC DISC)**

1. Anemia
2. Age-related macular degeneration (AMD)
3. Arterial insufficiency
4. Behçet disease (oculobuccogenital syndrome)
5. Buerger disease (thromboangiitis obliterans)
6. Coats disease (retinal telangiectasia)
7. Choroidal rupture
8. Diabetes mellitus
9. Drusen of optic nerve head
10. Eales disease (periphlebitis)
11. Geographic helicoid peripapillary choroidopathy
12. Glaucoma, chronic uncontrolled
13. Hereditary drusen of Bruch membrane
14. von Hippel-Lindau disease (retinocerebral angiomatosis)
15. Hypertensive retinopathy, advanced
16. Incontinentia pigmenti achromians (hypomelanosis of Ito syndrome)
17. Infection
   A. Endophthalmitis
   B. Congenital rubella syndrome (Gregg syndrome)
   C. Histoplasmosis
   D. Toxoplasmosis
18. Intraocular inflammation
   A. Rheumatoid arthritis
   B. Sarcoidosis syndrome (Schaumann syndrome)
   C. Uveitis (unspecified)
19. Myopia, severe
20. Norrie disease (fetal iritis syndrome)
21. Pseudotumor cerebri (Symond syndrome)
22. Retinal vein occlusion (see p. 468-472)
23. Retinitis pigmentosa
24. Retinopathy of prematurity (ROP)
25. Sickle cell disease (Herrick syndrome)
26. Takayasu disease (aortic arch syndrome)
27. Trauma
28. Tumors
   A. Benign
      (1) von Hippel-Lindau disease
      (2) Juxtapapillary capillary hemangioma
      (3) Nevus
   B. Malignant
      (1) Choroidal melanoma
      (2) Leukemia
      (3) Lymphoma
      (4) Metastatic tumors


**NEURORETINITIS (INFLAMMATION OF OPTIC NERVE AND ADJACENT RETINA)**

1. *Bartonella henselae* infection
2. Cat-scratch disease
3. Herpes simplex
4. Idiopathic
5. Mumps (epidemic parotitis)
6. *Nematode*
7. *Salmonella*
8. Syphilis
9. *Toxocara canis*
10. Toxoplasmosis


OPTIC-NERVE ATROPHY

1. Chromosome disorders
   A. Angelman syndrome (happy puppet syndrome; microdeletion of chromosome 15q11-13)
   B. Chromosome deletion (long arm) syndrome (de Grouchy syndrome)
   C. Cri-du-chat syndrome (cat-cry syndrome; deletion of short arm of chromosome 5)
   D. Patau syndrome (trisomy syndrome)
   E. Subacute sclerosing panencephalitis (Dawson disease)

2. Demyelinating and degenerative diseases
   A. Arylsulfatase A deficiency syndrome (ADL, metachromatic leukodystrophy)
   B. Devic syndrome (optical myelitis)
   C. Hereditary motor sensory neuropathy (HMSN I; Charcot-Marie-Tooth syndrome)
   D. Hereditary optic atrophy (Behr syndrome)
   E. Multiple sclerosis

3. Dermatologic disorders
   A. Keratodermia palmaris et plantaris
   B. McCune-Albright syndrome (fibrous dysplasia)
   C. Naegeli syndrome (reticular pigmented dermatosis)
   D. Porphyria cutanea tarda
   E. Pseudoxanthoma elasticum (Grönblad-Strandberg syndrome)
   F. Wrinkly-skin syndrome

4. Drugs, poisons, and vaccines, including the following:
   acetylphenazine  barbital  chloroquine
   acetylsalicylic acid beclomethasone  chlorpromazine
   adrenal cortex injection benzathine penicillin G  cholecalciferol
   aldosterone  betamethasone  cisplatin
   allobarbitral  bromide (?)  clindamycin
   amobarbital  bupivacaine (?)  cocaine
   antimony lithium  butobarbital  colchicine compounds
   thiomalate  butalbital  cortisone
   antimony potassium tartrate  butallylonal  cyclobarbital
   tartrate  butaperazine  cyclopentobarbital
   antimony sodium  butethal  danazol
   tartrate  calcifediol  dapsone
   antimony sodium  calcitriol  desoxycorticosterone
   thioglycollate  carbamazepine  dexamethasone
   antipyrine  carbon dioxide  diatrizoate meglumine
   aprobarbital  carbromal  or sodium
   aspirin  carphenazine  diethazine
   auranofin (?)  cephaloridine (?)  ergocalciferol
   aurothioglucose (?)  chloramphenicol (?)  ethambutol
   aurothioglycanide (?)  chlorprocaine (?)  ethopropazine
etidocaine (?)
fludrocortisone
fluorometholone
fluphenazine
fluprednisolone
gentamicin
glutethimide
gold Au 198
gold sodium thiomalate (?)
gold sodium thiosulfate (?)
heptabarbital
hexachlorophene
hexamethonium
hexethal
hexobarbital
hydrabamine
benoxymethyl penicillin
hydrocortisone
hydroxychloroquine
ibuprofen
interferon
iodine and iodine solutions and iodochlor-hydroxyquin
iodoquinol
isocarboxazid (?)
isoniazid
lidocaine (?)
medrysone
mephabarbital
mepivacaine (?)
meprednisone
mesoridazine
methaqualone (?)
metharbital
methdilazine
methiturial
methohexital
methotrimeprazine
methyl alcohol
methylene blue
methyprednisolone
metylpromyrol
mitotane
nadolol (?)
nalidixic acid
naprofen (?)
oxyphenbutazone
penicillamine
pentobarbital
perazine
pericyazine
perphenazine
phenelzine (?)
phenobarbital
phenoxymethyl penicillin
phenylbutazone
pipерacetazine
potassium penicillin G
potassium penicillin V
potassium phenethicillin
prednisolone
prilocaine (?)
primidone
probarbital
procaïne (?)
procaine penicillin G
procarbazine
prochlorperazine
promazine
promethazine
propiomazine
propoxycaïne (?)
quiline
radioactive iodides
secobarbital
silicone
sodium antimonylgluconate
sodium salicylate
stibocaptate
stibogluconate
stibophen
streptomycin
sulfacetamide
sulfachlorpyridazine
sulfacycline
sulfadimethoxine
sulfamerazine
sulfameter
sulfamethazine
sulfamethizole
sulfamethoxazole
sulfamethoxypyridazine
sulfanilamide
sulfaphenazole
sulfapyridine
sulfasalazine
sulfathiazole
sulfisoxazole
sulthiame
talbutal
tamoxifen
thiamylal
thiethylperazine
thiopental
thiopropazate
thiopropazine
thioridazine
tobramycin
tranylcypromine (?)
triamcinolone
trichloroethylene
trifluoperazine
triflupromazine
trimeprazine
tryparsamide
vaccines- influenza
vinbarbital
vinblastine
vincristine
vitamin A
vitamin D (retinol)

vitamin D2 (ergocalciferol)

vitamin D3 (cholecalciferol)
warfarin
5. Endocrine
   A. Cretinism (hypothyroidism)
   B. Cushing syndrome (adrenocortical syndrome)
   C. Diabetes mellitus
   D. DIDMOAD (diabetes mellitus and insipidus with optic atrophy and deafness) syndrome; Wolfram syndrome, Marquardt-Loriaux syndrome)
   E. Fröhlich syndrome (dystrophia adiposogenitalis)
   F. Hyperparathyroidism
   G. Hypophosphatasia (phosphoethanolaminuria)
   H. Juvenile diabetes mellitus (Mauriac syndrome)
   I. Pituitary gigantism syndrome (Launois syndrome)
   J. Simmonds syndrome (hypopituitarism syndrome)
   K. Retinohypophysis syndrome (LiJo Pavia-Lis syndrome)
   L. Zollinger-Ellison syndrome (polyglandular adenomatosis syndrome)

6. Granulomatoses
   A. Sarcoidosis
   B. Tuberculosis
   C. Wegener syndrome (Wegener granulomatosis)

7. Infectious
   A. African eye-worm disease (loiasis)
   B. Anthrax
   C. Congenital cytomegalic inclusion disease
   D. Congenital rubella syndrome (Gregg syndrome)
   E. Cysticercosis
   F. Deerfly fever (tularemia)
   G. Encephalitis
   H. Encephalomeningitis
   I. Echinococciosis (hydatid cyst)
   J. Lyme disease (borreliosis, relapsing fever)
   K. Malaria
   L. Meningitis
   M. Measles (morbilli)
   N. von Mikulicz-Radecki syndrome (dacryosialoadenopathy)
   O. Mumps (epidemic parotitis)
   P. Mycoplasma pneumoniae
   Q. Onchocerciasis syndrome (river blindness)
   R. Rocky Mountain spotted fever
   S. Syphilis (congenital or acquired)
   T. Toxoplasmosis
   U. Yellow fever

8. Inherited
   A. Congenital optic atrophy (autosomal dominant or recessive)
   B. Jensen syndrome (opticoacoustic nerve atrophy with dementia; X-linked)
   C. Juvenile optic atrophy (autosomal dominant)
   D. Metaphyseal dysplasia, anetoderma and optic atrophy (autosomal recessive)
   E. Myotonic dystrophy
F. Optic atrophy, cataract and neurologic disorder (dominant)
G. Optic atrophy, non-Leber type, early onset (X-linked)
H. Optic atrophy, polyneuropathy, and deafness (X-linked)
I. Optic atrophy, spastic paraplegia syndrome (X-linked)
J. Optic atrophy, spastic paraplegia, dementia (autosomal dominant)
K. Optic atrophy, nerve deafness, and distal neurogenic amyotrophy (recessive)
L. Optic atrophy with demyelinating of central nervous system (autosomal dominant)
M. Optic atrophy hypoplasia, familial, bilateral (autosomal dominant)

9. Inherited metabolic disorders
   A. Leukodystrophies
      (1) Adrenoleukodystrophy (ALD)
      (2) Canavan disease (spongy degeneration of the nervous system)
      (3) Cockayne syndrome
      (4) Homocystinuria syndrome
      (5) Krabbe disease
      (6) Maple syrup urine disease
      (7) Menkes disease (kinky-hair syndrome)
   B. Peroxisome abnormalities
      (1) Defective biogenesis:
         a. Infantile Refsum syndrome (heredopathia atactica polyneuritiformis)
         b. Neonatal ALD (adrenal cortical atrophy, patchy brain demyelination)
         c. Zellweger syndrome (cerebrohepatorenal syndrome)
      (2) Refsum syndrome (heredopathia atactica polyneuritiformis)
      (3) Rhizomeric chondrodysplasia punctata
      (4) Single enzyme deficiency
         a. Primary hyperoxaluria type I
         b. X-linked ALD
   C. Storage disorders
      (1) Lipoidoses
         a. Generalized gangliosidosis
            (i) Gangliosidosis GM2, type
            (ii) Generalized gangliosidosis GM type
            (iii) Juvenile gangliosidosis GM type
         b. Sphingolipidoses (arylsulfatase A deficiency syndrome)
            (i) Arylsulfatase A deficiency syndrome (metachromatic leukodystrophy)
               (a) Late infant form: Greenfield disease
               (b) Adult form: Bogaert-Nijssen-Peiffer syndrome
               (c) Austin disease (multiple sulfatase deficiency
               (d) Fabry disease (angiokeratoma corporis diffusum)
               (e) Krabbe disease (globoid cell leukodystrophy)
(f) Niemann-Pick syndrome (essential lipoid histiocytosis)
(g) Tay-Sachs syndrome (hexosaminidase deficiency)

(2) Glucose-phosphate dehydrogenase deficiency (von Gierke disease)
(3) Mucolipidoses IV (ML IV)
(4) Mucopolysaccharidoses (MPS) or lysosomal storage diseases
   a. MPS I-H (Hurler syndrome; chondroosteodystrophy or lipochondrodystrophy)
   b. MPS I-S (Scheie syndrome)
   c. MPS II (Hunter syndrome)
   d. MPS III (Sanfilippo syndrome)
   e. MPS IV (Morquio syndrome)
   f. MPS VI (Maroteaux-Lamy syndrome)
(5) Neural ceroid lipofuscinosis
   a. Infantile type: Haltia-Santavuori disease
   b. Late infantile type: Jansky-Bielschowsky disease (internuclear ophthalmoplegia)
   c. Juvenile type: Batten disease (Spielmeyer-Vogt-Sjögren syndrome; cerebroretinal degeneration)
(6) Other disorders involving lipids
   a. Bassen-Kornzweig syndrome (familial hypolipoproteinemia)
   b. Refsum syndrome (heredopathia atactica polyneuritiformis)

10. Local
    A. Aphakic cystoid macula edema (ACME; Irvine-Gass syndrome)
    B. Bird-shot chorioretinopathy
    C. Coats disease
    D. Drusen of optic nerve
    E. Glaucoma
    F. Vascular occlusion

11. Mental and psychomotor deficiency, retardation
    A. Drummond syndrome (idiopathic hypercalcemia)
    B. Familial dysautonomia (Riley-Day syndrome)
    C. Hallervorden-Spatz syndrome
    D. Hallgren syndrome (retinitis pigmentosa-deafness-ataxia syndrome)
    E. Kloepfer disease
    F. Rubinstein-Taybi syndrome

12. Miscellaneous
    A. Albinism
    B. Anemia
    C. Arachnoidal adhesion (e.g., caused by tabes)
    D. Bloch-Sulzberger disease (incontinentia pigmenti)
    E. Bobble-head doll syndrome (massive dilatation of third ventricle)
    F. Bonnet-Dechaume-Blanc syndrome (neuroretinoangiomatosis)
    G. Brown-Sequard syndrome
    H. Cerebellar ataxia (Louis-Bar syndrome)
I. Cerebral palsy
J. Cystic fibrosis syndrome
K. Foster-Kennedy syndrome (basal-frontal syndrome)
L. Histiocytosis X eosinophil granuloma (Hand-Schuller-Christian syndrome)
M. Incipient prechiasmal optic nerve compression syndrome
N. Laurence-Moon-Bardet-Biedl syndrome (retinitis pigmentosa-polydactyly-adiposogenital syndrome)
O. Leber syndrome
P. Oculodental syndrome (Peter syndrome)
Q. Optic cochleodental degeneration syndrome
R. Pelizaeus-Merzbacher disease (aplasia axialis extracorticalis congenita)
S. Posthypoxic syndrome
T. Pseudotumor cerebri
U. Rieger syndrome
V. Russell syndrome
W. Status dysraphicus syndrome (Passow syndrome)
X. Sphenomaxillary fossa syndrome (pterygopalatine fossa syndrome)
Y. Wagner disease (hereditary vitreoretinal degeneration)

13. Nutritional deficiency
   A. Avitaminosis B (Wernicke syndrome, beriberi)
   B. Avitaminosis B (pellagra)
   C. Garland syndrome (central nervous system deficiency)
   D. (?) Infantile neuroaxonal dystrophy (possible vitamin E deficiency, Seitelberger disease II)
   E. Kwashiorkor syndrome (hypoproteinemia syndrome)
   F. (?) Oculoorogenital syndrome (avitaminosis B with possible avitaminosis A)

14. Orbital
   A. Hutchinson- Pepper syndrome (metastatic infraorbital neuroblastoma)
   B. Rollet syndrome (orbital apex syndrome)
   C. Superior orbital fissure syndrome (Rochon-Duvigneaud syndrome)

15. Rheumatoid
   A. von Bechterew - Stumpelld syndrome (ankylosing spondylitis)
   B. Behçet disease (oculobuccogenital syndrome)
   C. Polymyalgia rheumatica
   D. SLE

16. Skeletal disorders
   A. Achondroplasia
   B. Albers-Schonberg syndrome (osteopetrosis)
   C. Anencephaly
   D. Apert syndrome (acrocephalosyndactylysm syndrome)
   E. Brachmann-de Lange syndrome
   F. Camurati-Engelmann syndrome (progressive diaphyseal dysplasia)
   G. Chondrodystrophia calcificans congenita (Conradi syndrome)
   H. Cloverleaf skull syndrome (Kleeblattschädel deformity)
   I. Craniometaphyseal dysplasia (Pyle syndrome)
   J. Craniostenosis
1. Oxycephaly
2. Plagiocephaly
3. Scaphocephaly
4. Trigonocephaly
K. Crouzon syndrome (craniofacial dysotosis)
L. Enchondromatosis syndrome (Ollier syndrome)
M. Generalized gangliosidosis GM type
N. Greig syndrome (hypertelorism ocularis)
O. Hallermann-Streiff-François syndrome (oculomandibulofacial dyscephaly)
P. Hutchinson-Gilford progeria syndrome (progeria)
Q. Marchesani syndrome
R. McCune-Albright syndrome (fibrous dysplasia)
S. Metaphyseal dysplasia, anetoderma, and optic atrophy
T. Microcephaly
U. Osteogenesis imperfecta (van der Hoeve syndrome)
V. Paget syndrome (osteitis deformans)
W. Primary hyperoxaluria type (osteodystrophy hydrocephalus)
X. Zellweger syndrome (cerebrohepatorenal syndrome)

17. Trauma
A. Direct and indirect optic nerve trauma
B. Electrical injury
C. Mechanical injury/surgical trauma (orbital floor fracture, malar fractures, Krönlein lateral orbitotomy)
D. Ocular contusion
E. Optic-nerve evulsion
F. Radiation

18. Tumors
A. Craniopharyngiomas
B. Ectopic pinealomas
C. Gliomas
D. Hemangiomas
E. Meningiomas
F. Nasopharyngeal carcinomas
G. Neuroblastomas
H. Pituitary adenomas
I. Pseudo-Foster Kennedy syndrome
J. Pseudo-pseudo-Foster Kennedy syndrome
K. von Recklinghausen syndrome (neurofibromatosis)
L. Tumors extending into fourth ventricle and cerebellum causing papilledema

19. Vascular
A. Aneurysm of internal carotid artery (foramen lacerum syndrome)
B. Arteriosclerosis
C. Cavernous sinus thrombosis (Foix syndrome)
D. Giant cell (temporal arteritis)
E. Hollenhorst syndrome (chorioretinal infarction syndrome)
F. Kussmaul disease (necrotizing angiitis)
G. Occlusion of the carotid artery
H. Sickle cell disease (Herrick syndrome)
I. Takayasu syndrome (aortic arch syndrome)


### Optic-Nerve Atrophy and Deafness

1. Adult form of arylsulfatase A deficiency (Bogaert-Nijssen-Peiffer syndrome; opticocochleodental degeneration)
2. Camuati-Engelmann syndrome (progressive diaphyseal dysplasia)
3. HMSN I (Charcot-Marie-Tooth syndrome)
4. Congenital rubella syndrome (German measles, Gregg syndrome)
5. Cockayne syndrome (dwarfism with retinal atrophy and deafness)
6. Craniometaphyseal dysplasia (Pyle syndrome)
7. DIDMOAD syndrome (optic atrophy, sensorineural deafness, diabetes mellitus and diabetes insipidus)
8. Dominant inheritance-congenital deafness and progressive optic nerve atrophy
9. Friedreich ataxia (optic atrophy, ataxia, and progressive hearing loss)
10. Generalized gangliosidosis GM type
11. Hallgren syndrome (retinitis pigmentosa-deafness-ataxia syndrome)
12. Juvenile diabetes mellitus
13. Krabbe syndrome (infantile globoid [II] cell leukodystrophy)
14. MLIV
15. MPS I-H (Hurler syndrome)
16. MPS II (Hunter syndrome)
17. MPS IV (Morquio syndrome)
18. (?) MPS (Maroteaux-Lamy syndrome)
19. (?) Niemann-Pick syndrome (essential lipoid histiocytosis)
20. Osteogenesis imperfecta
21. Recessive: nerve deafness, optic atrophy, and distal neurogenic amyotrophy
22. Refsum syndrome (phytanic acid oxidase deficiency)
23. Rosenberg-Chutorian syndrome
24. Sylvester disease
25. Treft syndrome


**Syndromes and Diseases Associated with Optic Atrophy**

1. Achondroplasia
2. Acquired lues (syphilis)
3. African eye worm disease (loiasis)
4. Albers-Schonberg syndrome (osteopetrosis)
5. Albinism
6. Albright syndrome (fibrous dysplasia)
7. Anemia
8. Anencephaly
9. Aneurysm of internal carotid artery syndrome (foramen lacerum syndrome)
10. Anthrax
11. Apert syndrome (acrocephalosyndactylism syndrome)
12. Arachnoidal adhesion, including tabes
13. Arteriosclerosis
14. Arylsulfatase A deficiency syndrome (metachromatic leukodystrophy)
15. Avitaminosis B2 (pellagra)
16. Bassen-Kornzweig syndrome (familial hypolipoproteinemia)
17. Batten-Mayou syndrome (cerebroretinal degeneration)
18. Behçet syndrome (oculobuccocentral syndrome)
19. Behr syndrome (optic atrophy-ataxia)
20. Bielschowsky-Jansky disease (internuclear ophthalmoplegia)
21. Bloch-Sulzberger disease (incontinentia pigmenti)
22. Bobble-head doll syndrome
23. Bonnet-Dechaume-Blanc syndrome (neuroretinointerstitialitis)
24. Brown-Marie syndrome (hereditary ataxia syndrome)
25. Brown-Sequard syndrome (lesion of spinal cord)
26. Carbon monoxide
27. Central nervous system deficiency-bitemporal pallor because of deficient diet (Garland syndrome)
28. Cerebral palsy
29. Cerebellar ataxia
30. Charcot-Marie-Tooth syndrome (progressive neuritic muscular syndrome)
31. Chondrodystrophy calcificans congenita (Conradi syndrome)
32. Chromosome deletion (long-arm) syndrome (de Grouchy syndrome)
33. Coats disease (retinal telangiectasia)
34. Cockayne syndrome (dwarfism with retinal atrophy and deafness)
35. Congenital cytomegalic inclusion disease
36. Congenital optic atrophy-autosomal dominant or recessive
37. Congenital syphilis
38. Craniometaphyseal dysplasia (Pyle syndrome)
39. Craniostenosis (including oxycephaly, scaphocephaly, trigonocephaly, and plagiocephaly)
40. Cretinism (hypothyroidism)
41. Cri-du-chat syndrome (cat-cry syndrome)
42. Crouzon syndrome (craniofacial dysostosis)
43. Cushing syndrome (adrenocortical syndrome)
44. Cystic fibrosis syndrome (fibrocystic disease of pancreas)
45. Cysticercosis
46. Dawson disease (subacute sclerosing panencephalitis)
47. Deerfly fever (tularemia)
48. de Lange syndrome (congenital muscular hypertrophy-cerebral syndrome)
49. Devic syndrome (optical myelitis)
50. Diabetes mellitus
51. Didmoad-Wolfram syndrome (diabetes mellitus and insipidus with optic atrophy and deafness-autosomal recessive)
52. Diencephalic syndrome (Penfield syndrome)
53. Disseminated lupus erythematosus (Kaposi-Libman-Sacks syndrome)
54. Dollinger-Bielschowsky syndrome (lipidosis)
55. Drugs, including the following:
    acetophenazine bromisovalum cyclopentobarbital
    allobarbital broxyquinoline cycloserine (?)
    alseroxylon (?) bupivacaine (?) dapsone
    aminosalicylate (?) butabarbital deferoxamine
    aminosalicylic acid (?) butalbital deserpidine(?)
    amobarbital butallylonal dexamethasone
    amodiaquine butaperazine dextrothyroxine (?)
    antimony lithium, butethal diethazine
    thiomalate calcitriol digitalis (?)
    antimony potassium tartrate carbromal diiodohydroxyquin
    tartrate carphenazine ergocalciferol
    antimony sodium chloramphenicol ergonovine (?)
    tartrate chloroprocaine (?) ergot (?)
    antimony sodium chloroquine ergotamine (?)
    thioglycollate chlorpromazine ethambutol
    antipyrine cholecalciferol ethopropazine
    aprobarbital clindamycin etidocaine (?)
    aspirin cobalt (?) ferrocholinate (?)
    barbital cocaine ferrous fumarate (?)
    betamethasone cortisone ferrous gluconate (?)
    bromide (?) cyclobarbital ferrous succinate (?)
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<td>oxyphenbutazone</td>
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<td>sulfacetamide (?)</td>
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<td>sulfachlorpyridazine (?)</td>
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<td>sulfadiazine (?)</td>
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56. Drummond syndrome (idiopathic hypercalcemia)
57. Drusen of optic nerve
58. Dyschondroplasia syndrome (Ollier syndrome)
59. Electrical injury
60. Encephalitis, acute
61. Engelmann syndrome (diaphyseal dysplasia)
62. Exfoliation syndrome
63. Foix syndrome (cavernous sinus thrombosis)
64. Foster-Kennedy syndrome (basal-frontal syndrome)
65. Friedreich ataxia (optic atrophy and sensorineural deafness)-recessive
66. Fröhlich syndrome (dystrophia adiposogenitalis)
67. Galactosyl ceramide lipidosis (globoid cell leukodystrophy)
68. Gangliosidosis GM 1, type
69. Generalized gangliosidosis (infantile)
70. Greig syndrome (hypertelorism ocularis)
71. Glaucoma
72. Glucose-phosphate dehydrogenase deficiency (von Gierke disease)
73. Grönblad-Strandberg syndrome (systemic elastodystrophy)
74. Hallermann-Streiff-François syndrome (oculomandibulofacial dyscephaly)
75. Hallervorden-Spatz syndrome (pigmentary degeneration of globus pallidus)
76. Hallgren syndrome (retinitis-pigmentosa deafness-ataxia syndrome)
77. Happy-puppet syndrome (puppet children)
78. Herrick syndrome (sickle cell disease)
79. Histiocytosis X (Hand-Schüller-Christian syndrome)
80. Hollenhorst syndrome (chorioretinal infarction syndrome)
81. Homocystinuria syndrome
82. Hutchinson syndrome (progeria)
83. Hydatid cyst
84. Hydrocephalus chondrodystrophicus congenita (Kleeblattschädel syndrome)
85. Hyperparathyroidism
86. Hypophosphatasia (phosphoethanolaminuria)
87. Incipient prechiasmal optic nerve compression syndrome
88. Infantile neuroaxonal dystrophy (Seitelberger disease II)
89. Infantile type of neuronal ceroid lipofuscinosis
90. Infections such as basal meningitis, infectious encephalomeningitis (especially measles epidemic parotitis), congenital neurosyphilis (rare before 2 years of age), and toxoplasmosis
91. Irvine syndrome (spontaneous rupture of vitreous face with vitreous adhesions to the wound followed by macular edema)
92. Jensen syndrome (opticoacoustic nerve atrophy with dementia)-X-linked
93. Juvenile diabetes-rare
94. Juvenile optic atrophy-autosomal dominant
95. Keratodermia palmaris et plantaris
96. Kloepfer disease
97. Krabbe disease (meningocutaneous syndrome)
98. Kussmaul disease (necrotizing angiitis)
99. Kwashiorkor (hypoproteinemia syndrome)
100. Laurence-Moon-Bardet-Biedl syndrome (retinitis pigmentosa-polydactyly-adiposogenital syndrome)
101. Leber syndrome (optic atrophy-amaurosis-pituitary syndrome)
102. Leigh disease (subacute necrotizing encephalomyelopathy)
103. Leukemia
104. Malaria
105. Maple syrup urine disease (branched-chain ketoaciduria)
106. Marchesani syndrome (brachymorphy with spherophakia)
107. Maroteaux-Lamy disease (MPS VI)
108. Marquardt-Loriaux syndrome
109. Measles (morbilli)
110. Menkes disease (kinky-hair disease)
111. MERRF syndrome
112. Metachromatic leukodystrophy (Greenfield disease)
113. Metaphyseal dysplasia, anetoderma, and optic atrophy - autosomal recessive
114. Microcephaly
115. Micro syndrome
116. Mikulicz-Radecki syndrome (dacryosialoadenopathy)
117. ML IV
118. MPS IH (Hurler syndrome)
119. MPS IS (Scheie syndrome)
120. MPS II (Hunter syndrome)
121. MPS IV (Morquio syndrome)
122. Multiple sclerosis (disseminated sclerosis)
123. Mumps
124. Naegeli syndrome (reticular pigmented dermatosis)
125. Niemann-Pick syndrome (essential lipoid histiocytosis)
126. Occlusion of carotid artery
127. Oculodental syndrome (Peter syndrome)
128. Oculoorogenital syndrome (riboflavin deficiency syndrome)
129. Onchocerciasis syndrome (river blindness)
130. Optic atrophy, cataract and neurologic disorder-dominant
131. Optic atrophy, nerve deafness, and distal neurogenic amyotrophy-recessive
132. Optic atrophy, non-Leber type, with early onset-x-linked
133. Optic atrophy, polynueropathy and deafness-x-linked
134. Optic atrophy, spastic paraplegia syndrome-x-linked
135. Optic atrophy with demyelinating of central nervous system-autosomal dominant
136. Optic nerve hypoplasia, familial, bilateral-autosomal dominant
137. Opticocochleodental degeneration syndrome
138. Orbital operation, such as following orbital floor fracture, reduction of malar fractures and Krönlein lateral orbitotomy
139. Osteogenesis imperfecta (van der Hoeve syndrome)
140. Paget syndrome (osteitis deformans)
141. Passow syndrome (syringomyelia)
142. Pelizaeus-Merzbacher disease (aplasia axialis extracorticalis congenita)
143. Pituitary gigantism syndrome (Launois syndrome)
144. Polymyalgia rheumatica
145. Porphyria cutanea tarda
146. Posthypoxic encephalopathy syndrome
147. Refsum syndrome (phytanic acid oxidase deficiency)
148. Relapsing fever
149. Retinohypophysary syndrome (Lijo Pavia-Lis syndrome)
150. Rieger syndrome (hypodontia and iris dysgenesis)
151. Riley-Day syndrome (congenital familial dysautonomia)
152. Rochon-Duvigneaud syndrome (superior orbital fissure syndrome)
153. Rocky Mountain spotted fever
154. Rollet syndrome (orbital apex-sphenoidal syndrome)
155. Rosenberg-Chutorian syndrome
156. Rubella syndrome (German measles)
157. Rubinstein-Taybi syndrome (broad thumb syndrome)
158. Russell syndrome
159. Sabin-Feldman syndrome
160. Sanfilippo disease (MPS III)
161. Scaphocephaly syndrome
162. Schaumann syndrome (sarcoidosis syndrome)
163. Schilder syndrome (encephalitis periaxialis diffusa)
164. Simmonds syndrome (hypopituitarism syndrome)
165. Spastic paraplegia, optic atrophy, dementia-autosomal dominant
166. Sphenomaxillary fossa syndrome (pterygopalatine fossa syndrome)
167. Spielmeyer-Vogt syndrome (cerebroretinal degeneration)
168. Spongy degeneration of the white matter
169. Suprarenal-sympathetic syndrome (adrenal medulla tumor syndrome)
170. Sylvester disease
171. Simmond syndrome (benign intracranial hypertension)
172. Takayasu syndrome (aortic arch syndrome)
173. Tay-Sachs syndrome (hexosaminidase deficiency)
174. Temporal arteritis syndrome (Hutchinson-Horton-Magath-Brown syndrome)
175. Toxins, including lead, chronic cyanide intoxication such as from eating cassava, thallium (used for treatment of scalp fungi)
176. Trauma, evulsion of optic nerve, and ocular contusion
177. Treft syndrome
178. Trisomy D syndrome (Patau syndrome)
179. Tuberculosis
180. Tumors, including craniopharyngiomas, ectopic pinealomas, gliomas, hemangiomas, meningiomas, nasopharyngeal carcinomas, neuroblastomas, pituitary adenomas, and tumors extending into fourth ventricle and cerebellum causing papilledema
181. Tunbridge-Paley disease
182. Vaccinia
183. Vascular accident
184. von Bekterev-Strumpell syndrome (ankylosing spondylitis)
185. von Recklinghausen syndrome (neurofibromatosis)
186. Wagner syndrome (hyaloideoretinal degeneration)
187. Wegener syndrome (Wegener granulomatosis)
188. Wernicke syndrome (thiamine deficiency)
189. Wrinkly-skin syndrome
190. Yellow fever
191. Zellweger syndrome (cerebrohepatorenal syndrome)
192. Zollinger-Ellison syndrome (polyglandular adenomatosis syndrome)


**OPTIC-NERVE HYPOPLASIA**

1. Chromosome disorders
   A. Down syndrome (trisomy 21)
   B. Deletion of long chromosome (13)
   C. Edward syndrome (trisomy 18)
   D. Patau syndrome (trisomy 13)
   E. Ring chromosome mosaicism
   F. 13q deletion syndrome
2. Idiopathic
3. Neurologic conditions
   A. Agenesis of the corpus callosum
   B. Anencephaly
   C. Basal encephalocele
   D. Behavioral problems
   E. Cerebellar atrophy
   F. Cerebral atrophy
   G. Cerebral infarcts
   H. Cerebral palsy
   I. Colpocephaly
   J. Congenital suprasellar tumors
   K. Congenital third, fourth, and sixth nerve palsy and up-gaze palsies
   L. Encephaloceles
   M. Hydranencephaly
   N. Hydrocephaly
   O. Mental retardation
   P. Migration disturbances
   Q. Perinatal encephalopathy
   R. Porencephaly
   S. Posterior pituitary ectopia
4. Ocular conditions
   A. Albinism
   B. Aniridia
   C. Astigmatism
   D. Blepharophimosis
   E. Colobomas (optic disc and chorioretinal)
   F. High myopia
   G. Microphthalmos
   H. Retinal vascular tortuosity

5. Systemic conditions
   A. Aicardi syndrome
   B. Albinism
   C. Chondrodysplasia punctata
   D. Cleft lip and palate
   E. Diabetes mellitus (maternal)-segmental optic nerve hypoplasia in infant
   F. Duane retraction syndrome
   G. Fetal alcohol syndrome, especially pituitary abnormalities with isolated
tortuosity of retinal veins
   H. Goldenhar-Gorlin syndrome
   I. Hemifacial atrophy
   J. Hypertelorism
   K. Intrauterine infections-including cytomegalovirus and hepatitis
      (1) Cytomegalovirus
      (2) Hepatitis
   L. Inherited (autosomal dominant or recessive)
   M. Klippel-Trenaunay-Weber syndrome
   N. Meckel syndrome
   O. Median cleft face syndrome
   P. de Morsier syndrome (septooptic dysplasia and mutations in the homeobox
gene HESX1/hesxl)
   Q. Osteogenesis imperfecta
   R. Potter syndrome
   S. Syndrome of nevus sebaceus of Jadassohn
   T. Topless optic disk syndrome

6. Toxins (maternal use of)
   A. Alcohol
   B. Lysergic acid diethylamide (LSD)
   C. Phencyclidine (PCP)
   D. Phenytoin
   E. Quinine
   F. Tobacco

32:89-112.


**OPTIC NEURITIS (PAPILLITIS AND RETROBULBAR NEURITIS)**

This condition is characterized by progressive loss of vision and possibly complete amaurosis; pain in or behind the eye, especially on lateral movement; Marcus Gunn pupillary phenomenon; and central or paracentral scotoma.

1. Demyelinating and degenerative diseases
   A. Adrenoleukodystrophy
   B. Hereditary ataxia (Brown-Marie syndrome)
   C. Multiple sclerosis
   D. Opticomyelitis (Devic disease)

2. Drugs, poisons, vaccines
   A. Drugs, including the following:
   - acetohexamide
   - acetyldigitoxin
   - alcohol
   - allobarbital
   - aminosalicylate
   - aminosalicylic acid (?)
   - amiodarone (?)
   - amitriptyline
   - amobarbital
   - aprobarbital
   - barbital
   - bromisovalum
   - broxyquinoline
   - bupivacaine (?)
   - butabarbital
   - butalbital
   - butallylona
   - butethal
   - calcitriol (?)
   - caramiphen
   - carbfomal
   - carmustine
   - chloral hydrate (?)
   - chloramphenicol
   - chloroprocaine (?)
   - chlorpropamide (?)
   - cholecalciferol
   - cisplatin
   - clindamycin
   - clophene (?)
   - cyclobarbital
   - cyclopentobarbital
   - cycloserine (?)
   - dexamethasone
   - desipramine
   - deslanoside
   - dextrothyroxine (?)
   - didanosine
   - didanosine
   - diethylpropion (?)
   - digitalis
   - digitoxin
   - digoxin
   - diiodohydroxyquin
   - diphtheria and tetanus toxoids (adsorbed)
   - diphtheria and tetanus toxoids and pertussis
diphtheria toxoid (adsorbed)
disulfiram
ergocalciferol (?)
ergonovine
ergotamine
ethambutol
etchlorvynol
ethionamide
etidocaine (?)
etoposide
ettetinate
fenoprofen
ferrocholinate (7)
ferrous fumarate (7)
ferrous gluconate (7)
ferrous succinate
ferrous sulfate (7)
fluorouracil (7)  methysergide  smallpox vaccine
gitalin  metronidazole(?)  streptomycin
glyburide  minoxidil(?)  sulfacetamide
heptabarbital  mumps virus vaccine  sulfachloropyridazine
hexethal  (live)  sulfacytine
hexobarbital  naproxen  sulfadiazine
ibuprofen  nialamide (?)  sulfadimethoxine
imipramine  nitrofurantoin (?)  sulfamerazine
indomethacin (?)  nortriptyline  sulfameter
influenza virus vaccine  nystatin  sulfamethizole
interferon alpha, beta, or gamma  oral contraceptives  sulfamethoxazole
iodide and iodine  oxyphenbutazone  sulfamethoxypyridazine
solutions and compounds  penicillamine  sulfanilamide
iodochlorhydroxyquin  pentobarbital  sulfaphenazole
iodoquinol  phenobarbital  sulfapyridine
iron dextran (?)  phenylbutazone  sulfasalazine
iron sorbitex (?)  piroxicam (?)  sulfathiazole
isoniazid  poliovirus vaccine (?)  sulindac (?)
isocarboxazid (?)  polysaccharide-iron complex (?)  talbutal
tlamoxifen (?)  methysergide  tetanus immune globulin
isotretinoin  prilocaine (?)  tetanus toxoid (?)
kamamycin (?)  primidone  thiamylal
kanamycin (?  probarbital  thiopental
lanatoside C  procaine (?)  thyroglobulin (?)
levothyroxine (?)  procarbazine  thyroid (?)
lidocaine  protriptyline  tolamide
tliothyronine (?)  quinacrine  tolbutamide
liotrix (?)  quinidine  tolypetaamine
measles and rubella virus vaccine  rabies immune globulin  trichlorehylene
measles, mumps, and rubella virus vaccine  rabeys vaccine  tryparsamide
mephobarbital  radioactive iodides  vaccine (adsorbed)
mepivacaine (?)  rifampin (?)  vinbarbital
metharbital  rubella and mumps virus vaccine (live)  vinblastine
methitural  methohexital  vincristine
methysergide  rubella virus vaccine  vitamin D
methysergide (7)  (live)  vitamin D2 (?)
methyl alcohol  secobarbital  vitamin D3 (?)
methylergonovine

B. Poisons (inhalation, skin absorption, or ingestion): alcohol, arsenicals (inorganic, gaseous, or organic), carbon disulfide, carbon tetrachloride, chlorodinitrobenzene and dinitrobenzene, copper, dinitrotoluene, Lysol solution, mercury, methyl bromide, methyl alcohol, siderosis (exogenous: intraocular foreign body or endogenous: iron metabolism disorders), tobacco, toluene (methyl benzene), trichlorethylene, tricresil phosphate, venoms (e.g., bee sting), vinyl benzene (styrene)
C. Vaccines and toxoids: Bacille Calmette-Guérin (bCG) vaccination, diphtheria toxoid (absorbed), diphtheria and tetanus toxoids (absorbed), influenza virus vaccine, measles or mumps or rubella live vaccine, poliovirus vaccine, rabies immune globulin, rabies vaccine, smallpox vaccine, tetanus immune globulin (?), tetanus toxoid (?), bee and wasp sting.

3. Infection and inflammation

A. Bacterial
   (1) Anthrax
   (2) Botulism (toxin from clostridium botulinum)
   (3) Brucellosis (undulant fever)
   (4) Diphtheria
   (5) Endocarditis
   (6) Leptospirosis (Weil syndrome)
   (7) Lyme disease (borreliosis, relapsing fever)
   (8) Mycoplasma pneumoniae
   (9) Pertussis (whooping cough)
   (10) Streptococcus (scarlet fever)
   (11) Syphilis (acquired lues)
   (12) Tuberculosis
   (13) Typhoid fever (abdominal typhus)

B. Fungal
   (1) Candidiasis
   (2) Coccidioidomycosis
   (3) Mucormycosis
   (4) Torulosis (cryptococcus)

C. Viral
   (1) Acquired immune deficiency syndrome (AIDS)
   (2) Bornholm disease (epidemic pleurodynia)
   (3) Chickenpox (varicella)
   (4) Epidemic keratoconjunctivitis
   (5) Equine encephalitis
   (6) Hepatitis A, B, C
   (7) Infectious mononucleosis
   (8) Influenza
   (9) Measles (rubeola)
   (10) Mumps
   (11) Pappataci fever (sandfly fever)
   (12) Poliomyelitis
   (13) Smallpox
   (14) Yellow fever

D. Protozoan
   (1) Malaria
   (2) Toxoplasmosis
   (3) Trypanosomiasis

E. Rickettsia
   (1) Boutonneuse fever rickettsia (Marseilles fever)
2. Japanese river fever (typhus)
3. Q fever
4. Rocky Mountain spotted fever

F. Orbit
1. Herpes zoster
2. Infections of the gasserian ganglion
3. von Mikulicz-Radecki syndrome (dacryosialoadenopathy)
4. Rollet syndrome (orbital apex syndrome)
5. Tolosa-Hunt syndrome (painful ophthalmoplegia)

G. Helminth infestations
1. Acanthamoeba
2. Echinococcosis (hydatid cyst)
3. Onchocerciasis (river blindness)
4. Toxocariasis (nematode ophthalmia syndrome)
5. Trichinellosis

H. Spread from sphenoid and posterior ethmoidal sinuses

I. Postinfectious
1. Guillain-Barré syndrome (acute infectious neuritis)
2. Reye syndrome (acute encephalopathy syndrome)
3. Subacute sclerosing panencephalitis (Dawson disease)

4. Noninfectious arteritis, hypersensitivity vasculitis
A. Involving small vessels
1. Drugs
2. Henoch-Schönlein

B. Involving small and medium-sized vessels
1. Polyarteritis nodosa (Kussmaul disease)
2. Necrotizing granulomatous arthritis
   a. Sarcoidosis
   b. Wegener granulomatosis (Wegener syndrome)
3. Buerger disease (thromboangiitis obliterans)
4. Localized arteritis
   a. Idiopathic
   b. Polyarteritis nodosa

C. Involving large, medium and small vessels
1. Arteritis in collagen vascular disease
   a. Behçet disease (oculobuccogenital syndrome)
   b. Progressive systemic sclerosis (PSS; scleroderma)
   c. Rheumatoid arthritis
   d. SLE
2. Giant cell (temporal) arteritis
3. Takayasu syndrome (aortic arch syndrome)

D. Idiopathic paroxysmal digital cyanosis (Raynaud disease)

E. Multiple myeloma (Kahler disease)

5. Others
A. Chorioretinitis
B. Cystic fibrosis syndrome
C. Hutchinson-Gilfor (progeria) syndrome
D. Hysteria
E. McCune-Albright syndrome (fibrous dysplasia)
F. Naegeli syndrome (melanophoric nevus)
G. Paget disease (osteitis deformans)
H. Parkinson syndrome (paralysis agitans)
I. Relapsing polychondritis
J. Stevens-Johnson syndrome (erythema multiforme exudativum)
K. Uveitis, including sympathetic ophthalmia

6. Systemic diseases
   A. Endocrine
      (1) Diabetes mellitus
      (2) Hypoparathyroidism
      (3) Hyperthyroidism (Basedow syndrome)
      (4) Hyperthyroidism
      (5) Juvenile diabetes-dwarfism-obesity syndrome (Mauriac syndrome)
      (6) Lactation
      (7) Pregnancy
      (8) Puberty
      (9) Retinohypophysary syndrome (Lijo Pavia-Lis syndrome)
   B. Nutritional diseases
      (1) Beriberi (vitamin B deficiency)
      (2) Carcinomatosis
      (3) Hyperemesis gravidarum
      (4) Pellagra (vitamin B deficiency)
   C. Rheumatic disease, arthritis
      (1) Felty syndrome
      (2) Juvenile rheumatoid arthritis (Still disease)
      (3) Polymyalgia rheumatica
      (4) Reiter syndrome (polyarthritis enterica)
      (5) Rheumatoid arthritis
   D. Miscellaneous
      (1) Amyloidosis (Lubarsch-Pick syndrome)
      (2) Chronic glomerulonephritis with secondary renal hypertension or pyelonephritis
      (3) Emphysema
      (4) Hepatic failure
      (5) Hypertension
      (6) Porphyria
      (7) Trauma
   A. Mechanical
   B. Radiation
      (1) Electromagnetic
         a. High voltage/lighting
b. Microwave
c. Laser bum
d. X-ray
(2) Radioactive source
a. a-ruthenium
b. b-betatron
c. g-cobalt
d. pisotope

8. Tumors
   A. Craniopharyngioma
   B. Hemangiopericytoma of optic nerve
   C. Myeloproliferative diseases
      (1) Hodgkin disease
      (2) Leukemia
      (3) Lymphoma
   D. Neuroblastoma


Extracted Table (Several unreadable titles)

PSEUDO OPTIC NEURITIS (LESSONS THAT MIMIC OPTIC NEURITIS)

1. Congenital retinoschisis
2. Hematoma
3. Ischemic optic neuropathy
4. Papilledema (see p. 593)
5. Retinal lesions that also exhibit metamorphopsia, e.g., serous or angiospastic retinopathy
6. Tumors
   A. Disc
      (1) Gliomas
      (2) Meningiomas
      (3) Metastatic carcinoma
      (4) Neurofibromas
B. Expanding lesions of anterior and middle cranial fossa producing central scotoma
   (1) Craniopharyngiomas
   (2) Ectopic pinealomas
   (3) Meningiomas
   (4) Metastatic carcinomas
   (5) Myeloproliferative diseases
       a. Hodgkin disease
       b. Lymphomas
       c. Plasmocytoma
   (6) Nasopharyngeal carcinomas
   (7) Pituitary adenomas


**OPTICOCILIARY SHUNTS (TORTUOUS, ECTATIC CHANNELS FROM OPTIC NERVE TO CHOROID)**

1. Arachnoid cyst of the optic nerve
2. Central retinal vein occlusion (see p. 468-472)
3. Chronic atrophic papilledema
4. Drusen of the optic nerve
5. Optic-nerve glioma
6. Primary nerve sheath meningioma
7. Sickle cell trait
8. Sphenoorbital meningioma


**PAPILLEDEMA (SWELLING OF OPTIC DISC)**

1. Drugs, poisons, and vaccines
   A. Drugs (those in all capitalized letters are drugs that also cause pseudotumor cerebri):
   - acetophenazine
   - amiodarone
   - ADRENAL CORTEX INJECTION
     - amobarbital
     - AMPHOTERICIN B
     - antimony potassium tartrate
     - antimony sodium
     - antimony lithium
   - ALDOSTERONE
     - thiomalate
   - alobarbital

antimony sodium
thioglycollate
aprobarbital
aspirin
auranofin (?)
aurothioglucone (?)
aurothioglycanide (?)
Azathioprine
barbital
benzathine penicillin G
bromide (?)
bupivacaine (?)
butabarbital
butalbital
butallylonal
butaperazine
butethal
calcitriol
carbamazepine
carbon dioxide
carphenazine
cephaloridine (?)
chlorambucil
chloramphenicol (?)
chloroprocaïne (?)
chlorpromazine
CHLORTETRACYCLINE
cholecalciferol
cisplatin
colchicine
CORTISONE
cyclobarbital
cyclopetobarbital
DANAZOL
demeclocycline
desoxydortico-sterone
dexamethasone
dextrothyroxine
diadosine
diethazine
doxycycline
e elt roxin
ethambutol
ethopropazine
etidocaine (?)
etoposide
FLUOROCORTISONE
fluorometholone
fluphenazine
FLUPREDNISOLONE
GENTAMICIN
glutethimide
gold Au 198
gold sodium
thiomalate (?)
gold sodium
thiosulfate (?)
heptabarbital
HEXACHLOROPHENE
hexathal
hexobarbital
hydrabamine
phenoxyethyl penicillin
HYDROCORTISONE
IBUPROFEN
INDOMETHACIN
INSULIN-LIKE GROWTH FACTOR I
interferon
interferon alpha, beta, or gamma
isocarboxazid (?)
isoniazid
ISOTRETINOIN
KETOPROFEN
LEVODOPA
LEVOTHYROXINE
lidocaine
LIOPTHYRONINE
LITHIUM CARBONATE
LITHIUM CITRATE
MANGANESE
mephobarbital
mepivaçaine (?)
MEPREDNISONE
mesoridazine
METHACYCLINE
methaqualone (?)
metharbital
methdilazine
methitural
methohexital
methotrimemprazine
methyl alcohol
methylene blue
METHYLPREDNISOLONE
methypyrrolon
mitotane
nadolol (?)
NALIDIXIC ACID
Naproxen (?)
NITROFURANTOIN
NITROGLYCERIN
NORPLANT
ofloxacin
ORAL CONTRACEPTIVES
OXYTETRACYCLINE
PARAMETHASONE
penicillamine
pentobarbital
perazine
PERHEXILINE
pericyazine
perphenazine
phenelzine (?)
phenobarbital
phenoxyethyl penicillin
PHENYLPROPANOL-AMINE
PHENYTOIN
piperacetaçine
POTASSIUM PENICILLIN G
POTASSIUM PENICILLIN V
POTASSIUM PHENETHICILLIN
PREDNISOLONE
PREDNISONE
prilocaine (?)
primidone
probarbital
**B. Poisons (inhalation, skin absorption, or ingestion)**

1. Carbon dioxide
2. Lead
3. Methyl alcohol

**C. Vaccines**

1. Diphtheria-tetanus toxoids-pertussis vaccine (absorbed)
2. Influenza virus vaccine
3. Measles or mumps or rubella live vaccine

**2. Intracranial causes-usually bilateral**

**A. Tumors**

*(1)* Frontal lobe lesion-mental changes (apathy, euphoria, and social behavioral changes); normal visual field if confined to frontal lobe; most likely, tumors are medulloblastoma, meningioma, astrocytoma, glioblastoma, or metastasis from lung or breast

*(2)* Temporal lobe lesions-formed hallucinations, superior homonymous quadrantanopia, or homonymous hemianopia, ipsilateral mydriatic fixed pupil and oculomotor paresis, and contralateral facial palsy; most likely, tumors are medulloblastoma, meningioma, astrocytoma, glioblastoma, or metastasis from lung or breast

*(3)* Parietal lobe lesions-visual agnosia such as alexia or dyslexia, complete homonymous hemianopia, or inferior homonymous quadrantanopia, disturbances of trigeminal nerve, including decreased corneal sensation and positive (asymmetric response) optokinetic nystagmus; most likely caused by:

a. Astrocytoma
b. Glioblastoma
c. Medulloblastoma
d. Meningioma
e. Metastasis from lung or breast

(4) Occipital lobe lesions - unformed visual hallucinations and homonymous congruous visual field defect; most likely caused by the following:
   a. Astrocytoma
   b. Glioblastoma
c. Hemangioma
d. Meningioma
e. Metastasis from lung or breast

(5) Third-ventricle and sellar lesions - visual field of bitemporal hemianopia or unilateral blindness and contralateral temporal hemianopia; most likely, tumors are craniopharyngioma

(6) Fourth-ventricle and cerebellum lesions - ataxia, asynergy, dysmetria, hypotonia, and acquired jerk nystagmus, usually horizontal and more pronounced in lateral gaze; most likely caused by:
   a. Astrocytoma
   b. Hemangioblastoma
c. Medulloblastoma
d. Metastasis from lung or breast

(7) Cerebellar-pontine angle tumor such as Cushing syndrome II (acoustic neuroma syndrome)

(8) Base skull tumor, such as Garcin syndrome (half-base syndrome)

(9) Chiasmal tumor such as Fröhlich syndrome (dystrophia adiposogenitalis)

(10) Neuroblastoma

(11) Russell syndrome (diencephalic syndrome)

(12) Zollinger-Ellison syndrome (polyglandular adenomatosis syndrome)

B. Decreased intracranial capacity, such as in acrocephalosyndactyly (Apert disease), Arnold-Chiari syndrome (cerebellomedullary malformation syndrome), craniofacial dysostosis (Crouzon disease), craniostenosis, hypertelorism, and tower skull (oxycephaly)

C. Pseudotumor cerebri (Symonds syndrome) - bilateral papilledema and increased intracranial pressure but negative neurologic and general physical findings
   (1) Addison disease (adrenal cortical insufficiency)
   (2) Autosomal dominant endosteal hyperostosis
   (3) Chronic respiratory insufficiency
   (4) Familial Mediterranean fever
   (5) Hypertension
   (6) Multiple sclerosis
   (7) Polyangiitis overlap syndrome
   (8) Psittacosis
   (9) Renal disease
   (10) Reye syndrome
(11) Sarcoidosis
(12) SLE
(13) Thrombocytopenia purpura
(14) Vitamin A (excessive) after overeating carrots in a weight loss program
(15) Drugs including the following:

<table>
<thead>
<tr>
<th>Drug Name</th>
<th>Table 1</th>
</tr>
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<tbody>
<tr>
<td>absorbed levodopa (?)</td>
<td>hexachlorophene</td>
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<tr>
<td>adrenal cortex injection</td>
<td>hydralazine</td>
</tr>
<tr>
<td>aldosterone</td>
<td>penicillin V</td>
</tr>
<tr>
<td>amiodarone</td>
<td>hydrocortisone</td>
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<tr>
<td>benzathine penicillin G</td>
<td>ibuprofen (?)</td>
</tr>
<tr>
<td>betamethasone</td>
<td>indomethacin</td>
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<tr>
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<td>chlorotetracycline</td>
<td>ketoprofen</td>
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<td>leuprolide acetate</td>
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<td>levothyroxine</td>
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<tr>
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<td>liothyronine</td>
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<tr>
<td>dexamethasone</td>
<td>liotrix</td>
</tr>
<tr>
<td>diphtheria and tetanus toxoids and pertussis vaccine</td>
<td>lithium carbonate</td>
</tr>
<tr>
<td>doxycycline</td>
<td>medroxyprogesterone</td>
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<td>etretinate</td>
<td>medrysone</td>
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<tr>
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<td>meprednisone</td>
</tr>
<tr>
<td>fluprednisolone</td>
<td>methylprednisolone</td>
</tr>
<tr>
<td>gentamicin</td>
<td>minocycline</td>
</tr>
<tr>
<td>griseofulvin</td>
<td>nalidixic acid</td>
</tr>
</tbody>
</table>

(16) Frankl-Hochwart syndrome (pineal-neurologic-ophthalmic syndrome)
(17) Glomus jugulare tumor
(18) Iron-deficiency anemia
(19) Menarche
(20) Pregnancy
(21) Thrombosis of the sagittal or lateral sinus, such as that following otitis media in children
(22) Yersinia pseudotuberculosis

3. Neurologic disorders
   A. Cerebral palsy
   B. Foster-Kennedy syndrome
      (1) Aneurysm of internal carotid, anterior cerebral, or anterior communicating artery
      (2) Arteriosclerotic plaques of internal carotid or anterior cerebral arteries
      (3) Chiasmal arachnoiditis secondary to trauma, spinal anesthesia, or syphilis
      (4) Craniopharyngioma with forward extension
(5) Frontal lobe tumors or abscess
(6) Glioma of the intracranial portion of optic nerve
(7) Internal hydrocephalus because of tumor of posterior fossa
(8) Old unilateral optic nerve atrophy (e.g., consecutive ischemic optic neuropathies)
(9) Olfactory groove, sphenoid ridge and suprasellar meningioma
  C. High cerebrospinal fluid protein content and defective absorption (e.g., Guillain-Barré syndrome (Acute Infectious Neuritis))
  D. Muscular dystrophy
  E. Parkinson syndrome (shaking palsy)
  F. Status dysraphicus syndrome (Passow syndrome, syringomyelia)
  G. Subdural or subarachnoid hemorrhage

4. Miscellaneous
   A. Abscess
   B. Angioedema
   C. Brown-Séquard syndrome
   D. Camurati-Engelmann syndrome (progressive diaphyseal dysplasia)
   E. Chediak-Higashi syndrome (anomalous leukocytic inclusions with constitutional stigma)
   F. Churg-Strauss syndrome (allergic granulomatosis and angiitis)
   G. Citrullinemia (late onset)
   H. Degos syndrome (malignant atrophic papulosus)
   I. Fabry disease (angiokeratoma corporis diffusum)
   J. Hydrocephalus
   K. Kenny syndrome
   L. McCune-Albright syndrome (fibrous dysplasia)
   M. Nocturnal hypoventilation
   N. Pelizaeus-Merzbacher syndrome (aplasia axialis extracorticalis congenita)
   O. Polymyalgia rheumatica
   P. Primary hyperoxaluria type
   Q. Renal insufficiency

5. Ocular cause-usually unilateral
   A. Acute glaucoma
   B. ACME (Irvine-Gass syndrome)
   C. Central retinal vein or artery occlusion
   D. Hypotony, including that following intraocular surgery
   E. Inflammatory
      (1) Bird-shot retinochoroidenopathy
      (2) Gumma of nerve head
      (3) Juxtapapillary choroiditis
      (4) Neuroretinitis (see p. 563)
      (5) Retinal vasculitis
      (6) Rocky Mountain spotted fever
      (7) Sarcoidosis
      (8) Tuberculoma of nerve head
      (9) Uveitis
10. Vasculitis

F. Trauma

G. Tumors

1. Glioma
2. Hemangioma
3. Melanocytoma
4. Melanotic sarcoma
5. Neurofibromatosis (von Recklinghausen disease)
6. Periocular and ocular metastatic tumors
7. Secondary carcinoma
8. Tuberous sclerosis

6. Orbital cause—usually unilateral, may have exophthalmos

A. Aneurysm of the ophthalmic artery
B. Orbital abscess
C. Rollet syndrome (orbital apex syndrome)
D. Scaphocephaly syndrome (craniofacial dysostosis)
E. Sinusitis
F. Superior orbital fissure syndrome (Rochon-Duvigneaud syndrome)
G. Trauma
H. Tumors

1. Benign
   a. Cystic adenoma
   b. Dermoid cyst
   c. Osteopetrosis (Albers-Schonberg disease)
   d. Paget disease

2. Malignant
   a. Fibrosarcoma
   b. Glioma
   c. Hutchinson- Pepper syndrome
   d. Lacrimal gland
   e. Lymphosarcoma
   f. Myosarcoma
   g. Osteosarcoma
   h. Secondary metastasis and extension from nasopharynx or sinuses

3. Orbital invasion by intracranial tumor (e.g., chordoma)

7. Systemic diseases—usually bilateral

A. Blood dyscrasias
   1. Iron-deficiency anemias
   2. Pernicious anemia
   3. Thrombocytopenic purpura

B. Carbohydrate metabolisms disorders
   1. Diabetes mellitus
   2. ML III
   3. MPS II (Hunter syndrome)
   4. MPS VI (Maroteaux-Lamy syndrome)
C. Cardiopulmonary insufficiency
   (1) Chronic bronchitis
   (2) Congenital heart disease
   (3) Cystic fibrosis of lungs
   (4) Pickwickian syndrome
   (5) Pulmonary emphysema

D. Collagen diseases
   (1) Polyarteritis nodosa
   (2) PSS (scleroderma)
   (3) Relapsing polychondritis
   (4) SLE

E. Endocrine
   (1) Addison disease (adrenal cortical insufficiency)
   (2) Diabetes mellitus (Willis disease)
   (3) Hyperparathyroidism
   (4) Hyperthyroidism (Basedow syndrome)
   (5) Hypothyroidism
   (6) Hypocalcemia
   (7) Hypoparathyroidism
   (8) Hypophosphatiasia
   (9) Idiopathic hypercalcemia (Drummond syndrome)
   (10) Menses
   (11) Pituitary deficiency
   (12) Pregnancy
   (13) Pseudohypoparathyroidism syndrome
   (14) Suppression of adrenal function from prolonged use of steroids
   (15) Suprarenal-sympathetic syndrome

F. Giant cell (temporal arteritis)

G. Hypertension/arteriosclerosis

H. Infectious (rare usually optic neuritis)
   (1) AIDS
   (2) Anterior poliomyelitis
   (3) Bang disease (brucellosis)
   (4) Chickenpox
   (5) Coccidioidomycosis
   (6) Echinococcosis (hydatid cyst)
   (7) Encephalitis
   (8) Infectious mononucleosis
   (9) Lyme disease (borreliosis, relapsing fever)
   (10) Malaria
   (11) Meningitis
   (12) Mycoplasma pneumoniae
   (13) Parasitic infections (e.g., cysticercosis, cryptococcus)
   (14) Parinaud syndrome (divergence paralysis)
   (15) Pertussis (whooping cough)
   (16) Presumed ocular histoplasmosis
(17) Psittacosis
(18) Sandfly fever (Pappataci fever)
(19) Trichinellosis
(20) Whipple disease (intestinal lipodystrophy)

I. Postinfectious
(1) Guillain-Barré syndrome (acute infectious neuritis)
(2) Reye syndrome (acute encephalopathy syndrome)
(3) Subacute sclerosing panencephalitis (Dawson disease)
(4) Vogt-Koyanagi-Harada syndrome (uveitis-vitiligo-alopecia-poliosis)

J. Myeloproliferative diseases
(1) Histiocytosis X (lipoid granuloma)
(2) Hodgkin disease
(3) Leukemia
(4) Multiple myeloma
(5) Mycosis fungoides (Sézary syndrome)
(6) Polycythemia vera

K. Paraproteinemias
(1) Cryoglobulinemia
(2) Macroglobulinemia
(3) Mediterranean fever
(4) POEMS syndrome (polyneuropathy, organomegaly, endocrinopathy, M protein, and skin changes)

L. Nutritional diseases
(1) Beriberi (thiamine deficiency)
(2) Pellagra (niacin deficiency)
(3) Plummer-Vinson syndrome (deficiency of vitamin B complex and iron)
(4) Vitamin B12 deficiency

M. Sarcoidosis (Heerford syndrome, Schaumann syndrome)

8. Trauma
A. Battered/shaken baby syndrome
B. Cerebral hemorrhage
C. Purtscher syndrome

9. Vascular malformations
A. Arteriovenous fistula
B. Aneurysms
(1) Bonnet-Dechaume-Blanc syndrome (neuroretinocardioglioma)
(2) Foramen lacerum syndrome (aneurysm of internal carotid artery syndrome)
(3) Superior vena cava syndrome
C. Cavernous sinus thrombosis (Foix syndrome)


**PSEUDOPAPILLEDEMA (MAY BE MISTAKEN FOR SWELLING OF OPTIC NERVE)**

1. Arteriovenous aneurysms (racemose aneurysms) of the retina (Wyburn-Mason syndrome)
2. Bergmeister papilla
3. Cervicooculoacousticus syndrome
4. Down syndrome
5. Drusen of optic nerve (see p. 559)
6. Epipapillary membrane and Bergmeister papilla
7. Fuchs coloboma (partial)
8. Hematoma
9. High hyperopia or astigmatism
10. Juvenile diabetes mellitus (Mauriac syndrome)
11. Medullated nerve fibers (opaque nerve fibers)
12. Normal variant
13. Opacities or haziness of the media, especially nuclear sclerosis of the lens
14. Optic neuritis or papillitis (see p. 578)
15. Peripapillary retinal hemangioma
16. Sarcoidosis (Schaumann syndrome)
17. Tilted disc (partial)
18. Tortuosity and anomalous early branching of the retinal vessels
19. Tumors of disc
   A. Gliomas
   B. Meningiomas
   C. Metastatic
   D. Neurinoma
   E. Neurofibroma


PERIPAPILLARY SUBRETINAL NEOVASCULARIZATION

1. Excessive laser treatment
2. Optic disc drusen
3. Optic nerve coloboma
4. Presumed histoplasmosis syndrome
5. Presumed sarcoidosis
6. Serpiginous peripapillary choroiditis


PIGMENTED TUMORS OF OPTIC DISC

1. Drusen
2. Bourneville syndrome (tuberous sclerosis)
3. Hemangioma of the disc with hemorrhages and secondary pigmentation
4. Malignant melanoma
5. Melanocytomas
6. Metastases


PSEUDOGLAUCOMATOUS ATROPHY OF OPTIC DISC

This condition involves cupping of the nerve head with optic atrophy and field defects simulating true glaucoma but without ocular hypertension.

1. Arteriosclerosis
2. Congenital anomalies of the optic disc
   A. Branching of vessels behind the lamina so that the individual branches appear at the disc margins
   B. Coloboma within the nerve sheath
   C. Congenital coloboma of the optic disc
   D. Morning-glory anomaly
   E. Oblique insertion of the optic nerve
   F. Traction of the disc with bowing of the scleral crescent
3. Giant cell (temporal) arteritis
4. Optic pit
5. Patients using digitalis
6. Reduced blood supply to optic nerve (e.g., acute hypotension, blood loss [severe] carotid insufficiency, gastrointestinal bleeding, ischemic optic neuropathy, myocardial infarction, pernicious anemia)
7. Schnobel cavernous atrophy
8. Syphilitic optic-nerve atrophy
9. Tumors arising near the chiasm


**TEMPORALLY DISPLACED DISC (DRAGGED DISC)**

1. Abnormal tortuous retinal vessels temporally
2. Ectopic macula
3. ROP
4. Temporally displaced vessels

18
Visual-Field Defects

PSEUDO- VISUAL-FIELD DEFECTS

1. Facial contour
   A. Prominent nose
   B. Bushy, projecting eyebrows
   C. High cheekbones
   D. Ptosis or blepharochalasis
   E. Sunken globes
   F. Fracture of orbit
2. Corneal opacities
3. Lenticular opacities, especially if miotics are used, will depress fields and exaggerate existing scotomas
4. Aphakia without lens or with convex lens; little distortion with contact lens or intraocular lens
5. Dull patient; patient may be mentally defective, have toxemia, arteriosclerosis, cerebral tumor, brain abscess, or increased intracranial pressure
6. Pupillary size
   A. Decrease in miotic field, especially with opacities of ocular media
7. Uncorrected refractive errors-correct for distance testing
8. Head tilting when the head is tilted toward the left shoulder; the right blind spot is elevated; when the head is tilted toward the right shoulder, the right blind spot is lowered
9. Environmental artifacts
   A. Reduction in illumination of screen and test objects magnifies field defect
   B. Variation in size of test object changes field defect
   C. Standard distance of patient from screen
   D. Attention of patient
   E. Technique of examiner
10. Psychologic artifacts
   A. Patient's misunderstanding of test
   B. Tiring of patient by prolonged testing
   C. Malingering-isopters at different distances are inconsistent
   D. Hysteria-spiral field defects may be found
11. Frames of glasses and segments of multifocal lenses
12. Colored contact lenses


**BILATERAL CENTRAL SCOTOMAS**

These are bilateral macular defects with decreased visual acuity; scotomas may be central or centrocecal.

1. Bilateral macular lesions, such as cysts or those due to hemorrhage, edema, degeneration, detachment, hole, or infection (see p. 439)
2. Bilateral optic-nerve lesions
   A. Papilledema with macular edema (see p. 593)
   B. Bipituitary adenoma compressing the prechiasmatic segment of the distal optic nerve
   C. Papillitis (see p. 578)
   D. Retrobulbar neuritis (see p. 578)
3. Diabetes mellitus
4. Familial optic atrophies (see p. 564)
5. Hyperbaric oxygen
6. Migraine-forerunner of visual aurae
*7. Nutritional deficiency, such as thiamine or vitamin B₁₂ deficiency
8. Pernicious anemia
9. Occipital cortex lesions
10. Toxic agents
    A. Aromatic aminocompounds and nitrocompounds-aniline, nitrobenzene, trinitrotoluene
    B. Carbon disulfide
    C. Drugs, including:
        acetophenazine         allobarbital          antazoline
        acetyldigitoxin        aluminum nicotinate (?)     aprobarbital
        adrenal cortex injection aminosalicylic acid (?)  aspirin
        alcohol                amiodarone             barbital
        aldosterone            amobarbital           beclomethasone
        alkavervir             amodiaquine            betamethasone
bromide
bromisovalum
brompheniramine
butabarbital
butalbital
butallylonal
butaperazine
butethal
caramiphen (?)
carboxamine
carbon dioxide
carbromal
carisoprodol
carphazinone
chloramphenicol
chloroquine
chlorpheniramine
chlorpromazine
chlorpropamide (?)
chlorotetracycline
ciprofloxacin
cisplatin
clemastine
clophene
cobalt (?)
contraceptives
cortisone
cyclobarbital
danazol
dapiprazole
  hydrochloride
deferoxamine
demeclocycline
deslanoside
desoxycorticosterone
dexamethasone
dexbrompheniramine
dexchlorpheniramine
diatrizoate meglumine
  and sodium
diazoxide
diethazine
diethylcarbamazine
digitalis
digitoxin
digoxin
dimethindene
diphenhydramine
diphenylpyraline
diphtheria and tetanus
  toxoids absorbed
disulfiram
doxylamine
emetine
epinephrine
ergonovine
ergot
ergotamine
ethambutol
ethchloryvnl
ethopropazine
fludrocortisone
fluorometholone
fluphenazine
fluprednisolone
gitalin
heptabarbital
hexamethonium
hexethal
hexobarbital
hydrocortisone
hydroxychloroquine
ibuprofen
indomethacin (?)
influenza virus vaccine
interleukin 2, 3, and 6
iodide and iodine
  solutions and
  compounds
iothalamic acid
isoniazid
lanatoside C
lidocaine
lithium carbonate
medrysone
metharbital
methdilazine
methitural
methohexital
methotrimetraopenazine
methoxsaalen
methylodopa
methylergonovine
methylprednisolone
methysergide
morphine (?)
naproxen
niacinamida (?)
nicotinic acid (?)
nicotinyl alcohol (?)
ofoxacin
opium
oral contraceptives
ouabain
oxygcn
oxyphenbutazone
paramethadionc
paramethasone
pentobarbital
perazine
pericyazine
perphenazine
phenobarbital
piperacetazine
prednisolone
prednisone
primdione
probarbital
prochlorperazine
promazine
promethazine
propiomazine
quinacrine
quinidine
quinine
radioactive iodides
secobarbital
sodium
sodium salicylate
streptomycin
sulfacetamide
sulfachlorpyridazine
sulfadiazine
sulfadimethoxine  sulfisoxazole  thyroid (?)
sulfamerazine   talbutal  trimcinolone
sulfameter  thiamylal  trichloroethylene
sulfamethizole  thiethylperazine  trifluoperazine
sulfamethoxazole  thiopental  triflupromazine
sulfamethoxypyridazine  thiopropazate  trimethadione
sulfanilamide  thioproperazine
sulfaphenazole  thioridazine  vinbarbital

D. Ethyl alcohol
E. Halogenated hydrocarbons-methyl chloride, methyl bromide, iodoform, trichloroethylene
F. Metals-lead, thallium (inorganic), arsenic
G. Methyl alcohol
H. Tobacco


**ENLARGEMENT OF BLIND SPOT**

1. Blind spot syndrome (multiple evanescent white-dot syndrome [MEWS])
2. Coloboma of the optic nerve
3. Drugs, including the following:
adrenal cortex injection  ergot  oxytetracycline
aldosterone  fludrocortisone  paramethasone
betamethasone  fluorometholone  prednisolone
carbon dioxide  fluprednisolone  prednisone
chlordetetracycline  hydrocortisone  quinacrine
cortisone  indomethacin (?)  tetracycline
demeclocycline  medrysone  triamcinolone
desoxycorticosterone  methacyclene  trichloroethylene
dexamethasone  methylprednisolone  vigabatrin
doxycycline  minocycline  vitamin A

3. Drusen of the optic nerve (see p. 559)
4. Glaucoma
5. Inferior conus
6. Inverted disc or nasally directed scleral canal
7. Juxtapapillary choroiditis
8. Medullated nerve fibers
9. Multifocal choroiditis
10. Multifocal evanescent white-dot syndrome
*11. Papilledema (pseudotumor cerebri) (see p. 593)
12. Papillitis (see p. 578)
13. Progressive myopia with a temporal crescent
14. Senility-senile halo
15. High-resistance instrument players


**ARCUATE (CUNEATE) SCOTOMA**

In this condition, scotoma follows the lines of the nerve fibers in the retina with the narrow end at the blind spot and broad end at horizontal raphe.

1. Acute bleeding episode
2. Branch artery occlusion
3. Branch vein occlusion
4. Chorioretinitis juxtapapillaris
5. Coloboma of the disc
6. Drusen of optic nerve
*7. Glaucoma
8. High myopia
9. Inferior conus
10. Ischemic optic neuropathy
11. Myelinated nerve fibers


**UNILATERAL SECTOR-SHAPED DEFECTS**

In this condition, the narrow end of scotoma characteristically touches the physiologic blind spot.
1. Optic disc involvement
   A. Glaucoma (early stages primarily on nasal side)
   B. Papillitis
   C. Secondary optic atrophy after choked disc (more on nasal side)

2. Retina
   A. Branch artery occlusion (see p. 468-472)
   B. Juxtapapillary chorioretinitis

3. Optic nerve-between disc and chiasm
   A. Aneurysm
   B. Drusen
   C. Tumor


**PERIPHERAL-FIELD CONTRACTION**

Central vision is present; the patient may complain of poor night vision.

1. Choroiditis-periphery of fundus
2. Chronic atrophic papilledema (pseudotumor cerebri)
3. Bilateral homonymous hemianopia (if the macular sparing in one homonymous hemianopia is larger than that in the other homonymous hemianopia, the spared central portion of the field has small vertical steps, above and below fixation, where the two areas of macular sparing do not quite coincide.)
   A. Cortical blindness with damage to occipital lobe and macular recovery
      1. Anoxia
      2. Carbon monoxide poisoning
      3. Cardiac arrest
      4. Cerebral angiography
      5. Exsanguination
      6. Trauma
   B. Stroke of infarction of occipital lobe
4. Drugs, including the following:
   acetophenazine  butalbital  chlorpromazine
   acridine  butallylonal  chlorpropamide (?)
   alcohol  butaperazine  clomiphene
   allobarbital  butethal  cobalt
   amobarbital  carbon dioxide  cortisone
   amodiaquine  carbon monoxide  cyclobarbital
   aprobarbital  carbromal  deslanoside
   arsenic  carisoprodol  desoxycorticosterone
   aspirin  carphenazine  dexamethasone
   barbital  chloramphenicol  diethazine
   bromisovalum  chloroquine  digitalis
<table>
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<tr>
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<td>compounds</td>
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5. Drusen of optic disc
6. Frontal-lobe tumors
7. General apathy in a lackadaisical subject
*8. Glaucoma
9. Hysteria and malingering
10. Many conditions in which night blindness occurs (see p. 656)
11. Optic atrophy (see p. 564)
12. Papillitis (see p. 578)
13. Post vitrectomy with fluid-air exchange
14. Retinitis-periphery of fundus
15. Retinitis pigmentosa
16. Unilateral concentric constriction, excluding diseased retina or glaucoma, suggests lesion of optic nerve and chiasm
   A. Meningioma of tuberculum sellae, sphenoid ridge, or the olfactory groove
   B. Tumor of optic nerve


ALTITUDINAL HEMIANOPIA

This condition comprises defective vision or blindness in the upper or lower horizontal half of the visual field. It may be unilateral or bilateral; unilateral field defect is prechiasmal.

1. Anemia - produces bilateral inferior altitudinal hemianopia
2. Anterior ischemic optic neuropathy
3. Bilateral branch retinal artery occlusion
4. Fusiform aneurysms (arteriosclerotic or congenital) - may produce inferior altitudinal hemianopia by pressure against the lateral halves of the optic chiasm or nerve
5. Herpes zoster
6. Lesion that presses the chiasm upward against the superior margin of the optic foramen
7. Occipital lobe lesions
   A. Hypoxia
   B. Stroke
8. Olfactory groove meningioma extending posteroinferior to compress the intracranial portion of the optic nerve
9. Optic-nerve lesion
   A. Anterior ischemia optic neuropathy
   B. Coloboma
   C. Glaucoma
   D. Optic neuritis
   E. Papilledema
   *F. Trauma
   G. Tumor
10. Sclerotic plaques of internal carotid artery or anterior cerebral arteries - pressure of plaques on optic nerve results in inferior altitudinal hemianopia
11. Following pars plana vitrectomy


BINASAL HEMIANOPIA

This condition comprises defects in the nasal half of visual fields, usually incomplete. This condition is due to lateral involvement of chiasm; it presupposes bilateral lesions.

1. Bilateral occipital lesion (thrombosis)
2. Chiasmic arachnoiditis, postneuritic optic atrophy, and bilateral retrobulbar neuritis of multiple sclerosis
3. Damage to chiasm
4. Drusen of optic nerve (see p. 559)
5. Fusiform aneurysms-arteriosclerotic or congenital-of internal carotid artery
6. Glaucoma
7. Meningiomas, especially from the lesser wing of the sphenoid bone
8. Nasal quadrant peripheral depression of glaucoma-bilateral and reasonably symmetrical
9. Pituitary tumor with third ventricle dilatation pushing laterally
10. Retinal damage
11. Severe exsanguination
12. Sclerotic plaques of internal carotid artery or anterior cerebral arteries
13. Symmetric lesions in the temporal halves of both retinas, such as severe retinal edema associated with diabetic retinopathy
14. Trauma


**BITEMPORAL HEMIANOPIA**

This condition involves defects in the temporal half of the visual field, usually incomplete; it is due to pressure on the optic chiasm.

1. Chiasmal lesions
   A. Congenital defect such as de Morsier syndrome (septooptic dysplasia); autosomal dominant
   B. Inflammatory lesions
      *(1) Basal meningitis, including: chronic chiasmal syphilitic, tuberculous, actinomycotic, and cysticercal arachnoiditis
      *(2) Chiasmal neuritis
   C. Tumors of the chiasm
      *(1) Primary tumors, including gliomas in childhood
      *(2) Secondary tumors (rare), including meningiomas, retinoblastoma, pinealoma, and ependymoma
   D. Vascular lesions
      (1) Arterial compression
      (2) Arteriosclerosis
      (3) Ectasia of the intracranial carotid arteries
      *(4) Intracranial aneurysms, such as congenital, endocardial emboli, traumatic, atheromatous, or syphilitic, especially intrasellar aneurysms
      (5) Thrombosis of the carotid artery

2. Perisellar lesions
   A. Parasellar tumors
      (1) Injuries to the chiasmal pathway, such as from trauma
*(2) Meningioma of the sphenoid ridge
(3) Migraine
(4) Sudden onset without apparent cause
   a. Arteriosclerotic or giant cell arteritic occlusion of nutrient vessels of the chiasm in older patients
   b. Disseminated sclerosis
(5) Tumors of the basal meninges
(6) Tumors of the sphenoid bone including osteochondroma, sarcoma, anaplastic carcinoma

B. Presellar tumors
*(1) Meningioma of the olfactory groove
(2) Neuroblastoma of the olfactory groove

C. Suprasellar tumors
(1) Chordoma
*(2) Craniopharyngioma—manifestations may include diabetes insipidus, infantilism, and calcification of hypophyseal—pituitary region
(3) Epidermoids
(4) Lymphoblastoma
(5) Pinealoma
*(6) Suprasellar meningioma
(7) Teratoma
(8) Tumors of the frontal lobe, including porencephaly (cystic cavity in brain substance) and glioma
(9) Tumors of the third ventricle and internal hydrocephalus, such as glioma and epidymoma

3. Pituitary lesions
   A. Pituitary hyperplasia
   B. Pituitary tumors
   (1) Adenoma
      a. Acidophilic adenoma—varies from gigantism to acromegaly
      b. Basophilic adenoma—hyperadrenalism (Cushing disease), Nelson syndrome
      c. Chromophobe adenoma—varies from no endocrine symptoms to panhypopituitarism; most common type of pituitary tumor, Fröhlich syndrome
   (2) Adenocarcinoma (rare)
   (3) Metastatic tumors as from breast (rare)


HOMONYMOUSQUADRANTANOPIA

In this condition, one quadrant is involved in upper or lower and right or left visual fields; etiology may include tumor, vascular lesion, or infection.

1. Superior homonymous quadrantanopia
   A. Inferior lip of the calcarine fissure-congruous
   B. Temporal lobe-incongruous
2. Inferior homonymous quadrantanopia
   A. Anton syndrome (denial-visual hallucination)
   B. Superior radiation in parietal lobe-incongruous
   C. Upper lip of the calcarine fissure in the occipital lobe-congruous


CROSSED QUADRANTANOPIA

In this condition, the upper quadrant of one visual field is along with the lower quadrant of opposite visual field.

* 1. Asymmetric homonymous hemianopia, such as vascular lesion of the upper lip of the calcarine area on one side and the lower lip of the opposite calcarine cortex
2. Chiasm compression from lesion below compressing it against contiguous arterial structure
3. Glaucoma
4. Inflammatory lesion, such as choroiditis juxtapapillaris


HOMONYMOUS HEMIANOPIA

This type of hemianopia affects the right or left halves of the visual fields; the lesion is posterior to the optic chiasm,

1. Optic tract lesions-visual conduction system posterior to optic chiasm and anterior to lateral geniculate body; lesion demonstrates incongruous field defect on side opposite to defect, often with decreased vision, optic atrophy and afferent pupil.
   A. Demyelinative disease-retrobulbar, multiple sclerosis, and Schilder disease
   B. Migraine
   C. Pituitary adenomas and craniopharyngiomas (most common); nasopharyngeal carcinomas, chordomas, infundibulomas, and gliomas (less common)
   D. Saccular aneurysms of internal carotid or posterior communicating artery
   E. Trauma
2. Temporoparietal lesions-temporal lobe lesions are manifest initially in the upper visual fields, whereas lesions of the parietal lobe are first manifest in the lower visual fields
   A. Diffuse demyelinative diseases
      (1) Krabbe type (Sturge-Weber-Krabbe syndrome)
      (2) Metachromatic leukoencephalopathy
      (3) Pelizaeus-Merzbacher type (aplasia axialis extracorticalis congenita)
      (4) Progressive multifocal leukoencephalopathy
      (5) Schilder type (encephalitis periaxialis diffusa)
      (6) Spongy degeneration of the brain (Canavan disease)
   B. Migraine
   C. Tumor-gradual onset of symptoms-lesions include intrinsic astrocytoma and glioblastoma, extrinsic meningioma, and lung metastasis
   D. Vascular lesions-sudden onset
      (1) Embolism-may be associated with rheumatic or arteriosclerotic heart disease, bacterial endocarditis, myocardial infarction, or septic focus in lungs
      (2) Occlusion-middle cerebral occlusion affects primarily the arm and face; anterior cerebral occlusion affects primarily the leg
      (3) Subdural hematoma-spontaneous or following trauma
      (4) Thrombosis-premonitory symptoms include unilateral blackouts in one eye
   E. Trauma (surgical)
3. Occipital lesions-congruous field defect and macular sparing most likely
   A. Demyelinative disease
      (1) Creutzfeldt-Jakob disease
      (2) Krabbe type (Sturge-Weber-Krabbe syndrome)
      (3) Metachromatic leukoencephalopathy
      (4) Pelizaeus-Merzbacher type (aplasia axialis extracorticalis congenita)
      (5) Progressive multifocal leukoencephalopathy
      (6) Schilder type (encephalitis periaxialis diffusa)
      (7) Spongy degeneration of the brain (Canavan disease)
   B. Migraine
   C. Poisons, such as carbon monoxide, digitalis, mescal, opium, lysergic acid diethylamide
   D. Trauma
      (1) Direct-penetrating missiles and depressed bone fragments
      (2) Indirect-general concussion syndrome
      (3) Temporal lobectomy
   E. Tumors-gradual onset of symptoms-lesions include intrinsic astrocytoma and glioblastoma, extrinsic meningioma, and lung metastasis
   F. Vascular lesion-sudden onset
      (1) Arteriovenous anomalies
      (2) Aneurysms (rare)
      (3) Occlusion of posterior cerebral artery-thrombotic or embolic
      (4) Subclavian steal syndrome, with reversal of blood flow through the vertebral artery


### SPIRAL-FIELD DEFECTS

*1. Hysteria
2. Radiation therapy in or about the retina, optic nerve, and anterior visual pathways*


### DOUBLE HOMONYMOUS HEMIANOPIA

This condition involves peripheral constriction with small vertical steps above and below fixation as a result of lesions of the occipital area and probable involvement of striate cortex of both occipital lobes.

1. Bilateral central retinal artery occlusion
2. Bilateral central retinal vein occlusion
3. Bilateral vascular lesions involving a calcarine fissure
4. Increased intracranial pressure with shift of uncal portion of temporal lobe down over edge of tentorium with compression of posterior cerebral arteries and infarction in calcarine cortex.
5. Partial recovery from cortical blindness (see p. 632) from trauma, anoxia, carbon monoxide poisoning, cerebral angiography, cardiac arrest, exsanguination, and other similar conditions
6. Severe end-stage glaucoma
7. Severe trauma with massive brain damage as in depressed fracture of occiput

PART II
General Signs and Symptoms
ACQUIRED MYOPIA *

This condition comprises an error of refraction in which parallel rays of light focus in front of the retina, usually producing blurred distant vision and clear near vision.

* 1. Conditions such as diabetes mellitus or nuclear sclerotic cataract in which there is increased index of refraction of lens
* 2. Refractive myopia-increased curvature of the refracting surfaces because of the following:
   A. Ciliary muscle spasm
      (1) Functional-adolescence, hysteria
      (2) Miotics such as carbachol, demecarium, echotothiophate, isofluorophate, neostigmine, and physostigmine
      (3) Trauma-ocular contusion or anterior dislocation of the lens
      (4) Mushroom (Amanita muscaria) poisoning
   B. Lens hydration changes-diabetes mellitus, dysentery, or toxemia of pregnancy
   C. Drug reaction-probably because of ciliary body edema, including the following:
      acetazolamide acetyphenazine adrenal cortex injection
alcohol
aldosterone
aspirin
beclomethasone
bendroflumethiazide
benzthiazide
betamethasone
betaxolol
butaperazine
carbachol
carphenazine
corticosterone
clofibrate
codeine
cortisone
cyclothiazide
demecarium
demeclocycline
desoxycorticosterone
dexamethasone
dichlorphenamide
diethazine
digoxin
digitalis (?)
doxycycline
droperidol (?)
edothing
ethopropazine
ethosuximide
ethoxolamide
etretinate
fludrocortisone
fluorometholone
fluphenazine
fluorphenolone
fluprednisolone
glibenclamide
haloperidol (?)
hyaluronic acid
hyaluronidase
hydrochlorothiazide
hydrocortisone
hydroflumethiazide
hydroxypropyl
ibuprofen
indapamide
isoflurophate
isosorbide dinitrate
isotretinoin
levobunolol
medrysone
meprednisone
mesoridazine
methacholine
methacycline
methazolamide
methdilazine
methotriepazine
methsuximide
methyclothiazide
methylcellulose
methylprednisolone
metolazone
minocycline
morphine
neostigmine
opium
oral contraceptives
oxygen
oxytetracycline
paramethasone
penicillamine
perazine
pericyazine
perphenazine
phenformin
phensuximide
physostigmine
pilocarpine
piperacetazine
polythiazide
prednisolone
prednisone
prochlorperazine
promazine
promethazine
propiomazine
quinethazone
quinine
sodium salicylate
spironolactone
sulfacetamide
sulfachlorpyridazine
sulfacycline
sulfadiazine
sulfadimethoxine
sulfamerazine
sulfameter
sulfamethazine
sulfamethizole
sulfamethoxazole
sulfamethoxypyridazine
sulfanilamide
sulfaphenazole
sulfapyridine
sulfasalazine
sulfathiazole
sulfisoxazole
tetracycline
thiethylperazine
thiopropazine
thioproperazine
thioridazine
timolol
triacminolone
trichlormethiazide
trifluoperazine
triflupromazine
trimeprazine

D. Elongated globe
E. Paralysis of accommodation for distance (sympathetic paralysis)-young patient with unilateral Homer syndrome or migraine
F. Retinopathy of prematurity (retrolental fibroplasia)
G. Congenital glaucoma
H. Albinism
I. Gyrate atrophy (ornithine ketoacid aminotransferase deficiency)
J. Hypoparathyroidism
K. Malaria
L. Inherited
   (1) Cochlear deafness with myopia and intellectual impairment-autosomal recessive
   (2) Epiphyseal dysplasia of femoral heads, myopia, deafness-autosomal recessive
   (3) Epiphyseal dysplasia, multiple, with myopia and conductive deafness-autosomal dominant
   (4) Microcornea and cataract-autosomal dominant
   (5) Microphthalmos with myopia and corectopia-autosomal dominant
   (6) Myopia-autosomal recessive or dominant or less often X-linked
   (7) Night blindness, congenital stationary with myopia (nyctalopia-myopia)-X-linked
   (8) Night blindness with high-grade myopia-autosomal recessive
   (9) Pinguecula blindness (total color blindness with myopia, achromatopsia with myopia)-autosomal recessive
M. With scleral buckling surgery

3. Syndromes associated with myopia
   A. Aberfeld syndrome (congenital blepharophimosis)
   B. Achard syndrome (Marfan syndrome with mandibulofacial dysostosis)
   C. Alport syndrome (hereditary familial congenital hemorrhagic nephritis)
   D. Bloch-Sulzberger syndrome
   E. Chromosome partial deletion (long-arm) syndrome
   F. Cohen syndrome
   G. Cri-du-chat syndrome
   H. de Lange syndrome (congenital muscular hypertrophy cerebral syndrome)
   I. Down syndrome (trisomy syndrome)
   J. Ehlers-Danlos syndrome (fibrodysplasia elastic a generalisata)
   K. Fetal alcohol syndrome
   L. Forsius-Eriksson syndrome (Aland disease)
   M. Gansslen syndrome (familial hemolytic icterus)
   N. Haney-Falls syndrome (congenital keratoconus posticus circumscriptus)
   O. Homocystinuria
   P. Hypomelanosis of Ito syndrome
   Q. Kartagener syndrome (sinusitis, bronchiectasis, situs inversus syndrome)
   R. Kniest syndrome
   S. Laurence-Moon-Bardet-Biedl syndrome (retinitis pigmentosa-polydactyly-adiposogenital syndrome)
   T. Marchesani syndrome (brachymorphy with spherophakia)
   U. Marfan syndrome (arachnodactyly dystrophia mesodermalis congenita)
   V. Marshall syndrome (atypical ectodermal dysplasia)
   W. Matsoukas syndrome (oculocerebroarticuloskeletal syndrome)
X. Myasthenia gravis (Erb-Goldflam syndrome)
Y. Noonan syndrome (male Turner syndrome)
Z. Obesity-cerebral-ocular-skeletal anomalies syndrome
AA. Oculodental syndrome (Peter syndrome)
BB. Pierre Robin syndrome (micrognathia-glossoptosis syndrome)
CC. Pigmentary ocular dispersion syndrome
DD. Rubinstein-Taybi syndrome (broad-thumbs syndrome)
EE. SED congenita (spondyloepiphyseal dysplasia, congenital type)-autosomal dominant
FF. Scheie syndrome
GG. Schwartz syndrome (glaucoma associated with retinal detachment)
HH. Siemens syndrome (hereditary ectodermal dysplasia syndrome)-autosomal recessive
II. Smith-Magenis syndrome
JJ. Stickler syndrome (hereditary progressive arthroophthalmopathy)-autosomal dominant
KK. Trisomy 20p syndrome
LL. Trisomy syndrome
MM. Tuumaala-Haapanen syndrome (unknown etiology, similar to pseudohypoparathyroidism)
NN. Van Bogaert-Hozoy syndrome (similar to Rubinstein-Taybi syndrome)
OO. Wagner syndrome (hyaloideoretinal degeneration)
PP. Weill-Marchesani syndrome (brachymorphy with spherophakia)
QQ. Wrinkly-skin syndrome
RR. XXXXY syndrome (hypogenitalism, limited elbow pronation, low dermal finger tip ridge count)

4. Transient myopia
   A. Chemical agents and disease
   *B. Diabetes
   *C. After surgery
   D. Trauma


ACQUIRED HYPEROPIA

This condition comprises far-sightedness and error of refraction in which parallel rays of light focus behind the retina, usually producing clear distant vision and blurred near vision.

1. Aarskog syndrome (facial-digital-genital syndrome)
2. Adie syndrome (tonic pupil)
3. *Aphakia
4. Best syndrome (vitelliform dystrophy)
5. *Diabetes mellitus (poorly controlled to controlled)
6. Down syndrome (mongolism)
7. Drugs, including the following:
   - antihistamines
   - cannabis
   - chloroquine
   - ergot
   - imipramine
   - meprobamate
   - parasympatholytic drugs
   - penicillamine
   - phenothiazine
   - sulfachloropyridazine (?)
   - sulfadiazine (?)
   - sulfadimethoxine (?)
   - sulfamerazine
   - sulfameter (?)
   - sulfamethazine
   - sulfamethizole
   - sulfamethoxazole
   - sulfamethoxypyridazine
   - sulfamethoxypyridazine (?)
   - sulfanilamide (?)
   - sulfaphenazole (?)
   - sulfapyridine (?)
   - sulfasalazine (?)
   - sulfathiazole (?)
   - sulfisoxazole (?)
   - tolbutamide (?)

8. Flat cornea
9. Gorlin-Chaudhry-Moss syndrome (multiple basal cell nevi syndrome)
10. Hyperopia-refractive or axial
11. Hypoglycemia
12. Kenny syndrome (nanophthalmos with hyperopia)
13. Leber congenital amaurosis
14. Lesions causing internal ophthalmoplegia with paralysis of accommodation
15. Macular edema
16. Orbital tumor with extraocular globe pressure and retinal striae
17. Postsurgical correction of myopia (retinal keratotomy, automated lamella keratoplasty, photoreactive keratectomy)
18. Presbyopia
19. Rubinstein-Taybi syndrome
20. Sorsby syndrome (hereditary macular coloboma syndrome)
21. Toxin of Clostridium botulinum

DYSMEGALOPSIA-OPTICAL ILLUSIONS OF SIZE

1. Macropsia (objects appear larger)
   *A. Miotics
   *B. Spasm of accommodation (see p. 416-417)
   C. Use of excessive plus lenses
2. Metamorphopsia (objects appear distorted)
   A. Cerebral
      (1) Drug intoxications
      (2) Epilepsy
      (3) Focal lesions such as thrombosis of right middle cerebral artery
      *(4) Migraine
      (5) Parietal lobe lesion, including tumor and vascular lesion
      (6) Schizophrenia
   B. Hysteria
   C. Ocular
      (1) Astigmatism
      *(2) Macular lesions, including orbital tumor with macular striae and macular edema, inflammation, heterotopia or hole
      (3) Posterior vitreous separation and residual vitreoretinal macular traction
      (4) Retinal detachment
   D. Paget disease (osteitis deformans)
3. Micropsia (objects appear smaller)
   A. Accommodative paralysis and subnormal accommodation
   B. Atropinization
   C. Botulism
   D. Diphtheria
   *E. Presbyopia
   F. Use of excessive minus lenses
   G. Use of scopolamine
4. Teleopsia (objects appear farther away than they actually are)
   A. Bilateral parietal lesion
   B. Parietal lesion in nondominant hemisphere


**BILATERAL TRANSIENT LOSS OF VISION (TRANSIENT DARKENING OF VISION)**

* 1. Circulatory disturbances when bending over or straining (postural hypotension)
2. Essential hypotension
   A. Arteriosclerosis
   B. Chronic hypotension
   *C. Fatigue
   D. Hormonal disorders
   E. Hunger
   F. Vitamin deficiency
3. Fainting with vasomotor collapse
4. Heart failure
5. Transurethral resection of the prostate


**AMAUROSIS FUGAX (TRANSIENT MONOCULAR BLACKOUT OF VISION)**

1. Amaurosis fugax syndrome
2. Arteriosclerosis, hypertension, and hypertensive crisis
3. Canalis opticus syndrome: functional-hysteria, neurasthenia
*4. Cerebrovascular insufficiency
   A. Arterial aneurysms
   B. Congenital or acquired arteriovenous malformations
   C. Fibromuscular hyperplasia
   D. Post-traumatic acute and chronic arterial occlusion
   E. Takayasu syndrome (pulseless disease)
*F. Unilateral occlusive carotid disease
5. Compressive optic neuropathy
6. Corneal surface problems
7. Functional-hysteria, neurasthenia
8. Hematologic causes
   A. Emboli
1. Infective, such as subacute bacterial endocarditis
2. Gas in dysbarism
B. Idiopathic thrombocytosis
C. Multiple myeloma (Kahler disease)
D. Polycythemia (Vaquez disease)
E. Severe anemia
F. Sickle cell disease (Herrick syndrome)

9. Hypotension of fundus
A. Cardiac arrhythmia
B. Glaucoma, narrow angle, pigment dispersion
C. Gaze-evoked amaurosis (transient monocular loss of vision occurring in a particular direction of eccentric gaze)
   *(1) Bone fragment adjacent to optic nerve following orbital fracture
   *(2) Central retinal artery occlusion
   *(3) Optic nerve sheath meningiomas
   *(4) Orbital cavernous hemangiomas
   *(5) Orbital osteoma
B. Impending vascular occlusion, retinal vasospasm associated with systemic vasospastic disease (migraine)
C. Increased intracranial pressure, such as from intracranial tumors that interfere with vascular supply to the optic nerve
D. Increased venous pressure
   (1) Impending central retinal vein occlusion (see p. 468-472)
   (2) Intermittent elevation of intraocular pressure (glaucoma)
E. Negative G-force in pilots-circular maneuver with head toward the center of the circle
F. Ocular ischemic syndrome
G. Orbital vascular insufficiency with giant cell arteritis
H. Papilledema—lasts for to seconds (see p. 593)
I. Positive G-force in pilots-circular maneuver with feet toward center of circle
J. Vasospasm including temporal arteritis, polyarteritis nodosa, eosinophilic vasculitis, migraine and cluster headache

10. Large vitreous floater
11. Optic disc anomalies (congenital)
12. Ornithine transcarbamoylase deficiency
13. Pituitary tumor
14. Polymyalgia rheumatica
15. Quinine poisoning
16. Raynaud disease (paroxysmal digital cyanosis)
17. Retrobulbar anesthesia
18. Schleral buckling procedure
19. Spontaneous bleeding from a normal-appearing iris or intraocular lens iris touch
20. Taveras syndrome (progressive intracranial arterial occlusion syndrome)
21. Uhthoff symptom—vision decreased with exercise or ocular hyperthermia can occur with:
   A. Friedreich ataxia
B. Insufficiency of posterior cerebral arteries
C. Intrasellar and parasellar tumor
D. Multiple sclerosis (disseminated sclerosis)

22. Uremic amaurosis-with eclampsia
23. Vasospasm
24. Wasp sting


**SUDDEN PAINLESS LOSS OF VISUAL ACUITY-ONE EYE**

1. Acute keratoconus
2. Complication of retrobulbar block
3. Injury to the optic nerve
4. Meningeal carcinomatosis
5. Occlusion of central retinal artery
*6. Retinal detachment
7. Vitreous or retinal hemorrhage


**POSTTRAUMATIC LOSS OF VISION**

*1. Acute (angle-closure) glaucoma precipitated by emotional trauma of recent accident or from intumescent lens capsular trauma or other blunt trauma
2. Avulsion of optic nerve by lateral orbital wall trauma or contrecoup blow to head
3. Central retinal artery or vein occlusion (from markedly increased orbital pressure or embolus)*
4. Cortical blindness from hematoma, ischemia, or anoxia (patient may be unaware of blindness)
5. Hyphema, vitreous hemorrhage
6. Hysteria
7. Indirect trauma to optic nerves or chiasm
8. Intracranial interruption of visual pathways (hemorrhage, foreign body)
   *9. Lid swelling, blood or foreign material covering cornea, corneal damage
10. Malingering
   *11. Retinal detachment
12. Traumatic cataract, luxation of the lens (see p. 401)
13. Traumatic retinal edema and hemorrhages of retina from direct or contrecoup blows


**DECREASED VISUAL ACUITY**

1. Achromatopsia
2. Amblyopia ex anopsia-disuse
   A. Anisometropia-difference in refractive error between the eyes
   B. Monocular occlusion
   C. Strabismus-esotropia, exotropia, or hypertropia
   D. Unilateral atropinization
3. Anomalous elevation of optic disc with hyperplastic glial tissue and anomalous retinal vessels
4. Apparently normal eye with central fixation with poorer visual acuity in one than other eye-anisometropia
5. Apparently normal eye with normal fixation with disparity between near and distance vision-amblyopia, hysteria, malingering, retrobulbar neuritis, presbyopia, and micronystagmus
6. Apparently normal eye with normal fixation with poor distance and near vision astigmatism, amblyopia, hyperopia in older persons
7. Drugs, including the following:
   - acebutolol
   - aceclidine
   - acetaminophen
   - acetanilid
   - acetazolamide
   - acetophenazine
   - acetyldigoxin
   - acid bismuth sodium tartrate
   - acyclovir
   - adiphenine
   - adrenal cortex injection
   - albuterol
   - alcohol
   - aldosterone
   - alkaavervir
   - allobarbital
   - acetohexamide
   - allopurinol
   - alprazolam
   - alseroxylon
   - aluminum nicotinate
   - amantadine
   - amphotericin B
   - amiodarone
   - amitriptyline
   - amobarbital
   - aminosalicylic acid
   - amiphenine
   - amphotericin B
   - amyl nitrite
   - anisindione
   - anisotropine
   - antazoline
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psilocybin
pyridostigmine
pyrilamine
quinacrine
quinethazone
quinidine
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rabies vaccine
radioactive iodides
ranitidine
rauwolfia serpentina
rescinnamine
reserpine
rifampin
rubella and mumps virus vaccine (live)
rubella virus vaccine (live)
scopolamine
secobarbital
semustine
silver nitrate
silver protein
smallpox vaccine
sodium antimonylgluconate
sodium salicylate
spironolactone
stibocaptate
stibogluconate
stibophen
streptomycin
streptozocin
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sulfachlorpyridazine
sulfadiazine
sulfadimethoxine
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 uracil mustard
 urea
 urethan
 verapamil
 veratrum viride
 alkaloïds
 vinbarbital
 vinblastine
 warfarin
8. Hysteria
9. Irregular astigmatism-distortions in the anterior corneal surface (scarring, ectasia, edema, ulcer, postinflammatory processes)
10. Macular pathology (including edema, hemorrhage, or scar tissue)
11. Malingering
*12. Myopia
13. Myotonic dystrophy-exertional vision loss
14. Nystagmus
15. Opacities of cornea, lens, or vitreous precluding good vision
16. Optic neuritis-retrobulbar and papillitis, including toxic causes such as those due to tobacco, alcohol, and quinine (see p. 564)
17. Sphenoid sinus mucocele
18. Transient refractive errors
   *A. Hyperopia
   B. Myopia-diabetes


**BILATERAL BLURRING OF VISION**

1. Drug-induced (see paresis of accommodation, p. 631-632)
2. Intracranial hypertension and advanced papilledema (see p. 593)
*3. Migraine-attacks last 15 to 20 minutes
4. Narcolepsy
5. Refractive error (myopia, hyperopia, presbyopia)
6. Retinal "blackout" experienced by pilots
7. Severe systemic hypertension
8. Systemic hypotension
9. Vertebrobasilar insufficiency


**CORTICAL BLINDNESS (CEREBRAL BLINDNESS)**
This condition involves a complete loss of all visual sensation, including all appreciation of light and dark and a loss of reflex lid closure to bright illumination and to threatening gestures; retention of pupil constriction to light and accommodation; normal ophthalmoscopic examination; and normal motility. It may be associated with hemiplegia, sensory disorders, aphasia, and disorientation.

1. Degenerative conditions
   A. Alper progressive gray matter
   B. Cerebral dysgenesis associated with dementia
   C. Creutzfeldt-Jakob disease (corticostriatospinal degeneration)
   D. Cytomegalic inclusion disease (rare)
   E. Galactosemia
   F. Hodgkin disease
   G. Infantile neuroaxonal dystrophy
   H. Krabbe syndrome
   I. Phenylpyruvic oligophrenia
   J. Pompe disease (generalized glycogenosis)
   K. Porencephaly
   L. Renal failure
   M. Schilder disease (encephalitis periaxialis diffusa)
   N. Scholz subacute cerebral sclerosis
   O. Spongy degeneration of the brain
   P. Subacute sclerosing panencephalitis
   Q. Tay-Sachs disease (familial amaurotic idiocy)
   R. Toxoplasmosis (rare)

2. Drugs, including the following:
   - alcohol
   - bendroflumethiazide (?)
   - benzhiazide (?)
   - carbon dioxide
   - carbon monoxide
   - chloroform (?)
   - chlorothiazide
   - chlorothalidone
   - cisplatin
   - corticotropin
   - cyclosporin
   - cyclothiazide (?)
   - diatrizoate meglumine or sodium estradiol
   - ether (?)
   - etoposide
   - etretinate
   - FK506
   - glycine
   - hexamethonium chloride
   - hydrochlorothiazide (?)
   - hydroflumethiazide (?)
   - indapamid (?)
   - iothalamate meglumine or sulfamethazine
   - indapamid (?)
   - iopamidol
   - iothalamate meglumine or sodium iothalanuc
   - ketamine (?)
   - lead poisoning
   - meglumine
   - metohalamide
   - methadone
   - methylergonovine (?)
   - metolazine (?)
   - methyclothiazide (?)
   - methylergonovine (?)
   - metrizamide
   - nifedipine
   - nitroglycerin
   - nitrous oxide (7)
   - polythiazide (7)
   - quinethazone
   - sulfachlorpyridazine
   - sulfacytine
   - sulfadiazine
   - sulfadiazine
   - sulfamerazine
   - sodium iothalanuc
   - sulfameter
   - sulfamethazine
   - sulfamethizole
   - sulfamethoxazole
   - sulfamethoxypyridazine
   - sulfanilamide
   - sulfaphenazonel
   - sulfapyridine
   - sulfasalazine
   - sulfathiazole
   - sulfisoxazole
   - tansy poisoning
   - thiopental (?)
   - trichlormethiazide (?)
   - vinblastine
*3. Inflammatory lesions
   A. Bacterial endocarditis
   B. Encephalitis (including that due to measles and pertussis) and subacute
      sclerosing panencephalitis
   C. Influenza
   D. Meningococcal meningitis
   E. Mumps
   F. Pneumococcal meningitis
   G. Syphilitic meningitis

4. Space-taking lesions, such as tumors, gummas, abscesses, and cysts

*5. Trauma
   A. Birth trauma, including heart dysfunction, postictal, and vertebral artery injury
   B. Chiropractic manipulation of the neck and odontotic subluxation
   C. Posthypoxic syndrome
   D. Subdural hematoma with cerebral edema
   E. Occipital region
   F. Ventriculography and ventriculoatrial shunt operation

*6. Vascular lesions
   A. Air embolism
   B. Angioma of occipital region
   C. Angiospastic lesions, including hypertension, nephritis, eclampsia, uremia, and
      chronic lead poisoning (saturnism)
   D. Anoxia from chronic respiratory insufficiency
   E. Anoxia from high altitude
   F. Basilar artery thrombosis
   G. Bilateral posterior cerebral artery occlusion
   H. Blood loss syndrome (acute cerebral hypotension)
   I. Blood transfusion reaction
   J. Cardiac arrest
   K. Cerebral hemorrhage
   L. Electroshock
   M. Following burns and sunstroke
   N. Following cardiac, cerebral or vertebral angiography
   O. Hemorrhage in spastic paralysis
   P. Herniation of hippocampal gyrus associated with subdural hematoma
   Q. Hydrocephalus and microcephaly
   R. Malaria
   S. Obstruction of the local venous sinus, such as from septic thrombosis of
      superior longitudinal sinus
   T. Periarteritis nodosa
   U. Subarachnoid hemorrhage
   V. "Subclavian steal syndrome" with reversal of blood flow through the vertebral
      artery
W. Thrombotic thrombocytopenic purpura


Extracted Table Cortical Blindness

**BLINDNESS IN CHILDHOOD**

1. Cornea
   A. Hereditary dystrophies
   B. Inflammations such as varicella, rubeola, vaccinia, and gonorrhea, ophthalmia neonatorum, and pemphigus
   C. Trauma (abrasion or laceration)
2. Cortical blindness (see p. 632)
3. Globe
   A. Anophthalmos (see p. 228)
   B. Buphthalmos (see p. 222)
   C. Congenital, primary infantile, or secondary glaucoma (see p. 305)
   D. Hydrophthalmos
   E. Microphthalmos (see p. 220)
4. Lens
   A. Aphakia
   B. Congenital cataracts (see syndromes associated with cataracts, p. 410-416)
5. Optic nerve
   A. Aplasia
   B. Asphyxia at birth
   C. Associated with widespread disease such as mental deficiency, cerebral palsy, or epilepsy
   D. Atrophy (hereditary or secondary) (see p. 564)
   E. Cavernous sinus thrombosis (Foix syndrome)
   F. Cerebral hemorrhage (associated with major brain damage from accidental trauma, abuse or birth trauma)
   G. Crouzon syndrome (craniofacial dysostosis)
   H. Hydrocephalus
   I. Inflammatory damage-encephalomyelitis, encephalitis, tuberculosis
J. Osteopetrosis (Albers-Schönberg syndrome)
K. Subdural hematoma
L. Trauma-fracture at the orbital canal
M. Tumors

6. Psychic blindness
   A. Agnosia
   B. Alexia

7. Retina
   A. Achromatopsia
   B. Albinism
   C. Coats disease (retinal telangiectasia)
   D. Early chorioretinal heredodegenerations, including Stargardt disease and pigmentary retinopathy (see pseudoretinitis pigmentosa, p. 497-499)
   E. Embryopathies, including rubella, toxoplasmosis, and syphilis
   F. High myopia
   G. Infantile macular degeneration
   H. Pseudoretinitis pigmentosa (see p. 497-499)
   I. Reese retinal dysplasia
   J. Retinal detachment
   K. Retinoblastoma
   L. Retinoschisis
   M. Retinopathy of prematurity
   N. Tapetoretinal degeneration

8. Syndromes associated with amaurosis or blindness
   A. Adie syndrome
   B. Davidoff single ventricle
   C. Laurence-Moon-Bardet-Biedl syndrome (retinitis-pigmentosa-polydactyly-adiposogenital syndrome)
   D. Malformative syndrome with cryptophthalmos
   E. Marfan syndrome (arachnodactyly dystrophia mesodermalis congenita)
   F. Metachromatic leukodystrophy (arylsulfatase A deficiency syndrome)
   G. Niemann-Pick syndrome (essential lipoid histiocytosis)
   H. Sandhoff disease
   I. Schilder disease (encephalitis periaxialis diffusa)

9. Uveal tract
   A. Chorioretinitis
   B. Congenital coloboma
   C. Iridocyclitis

10. Vitreous
    A. Persistence of primary vitreous
    B. Pseudoglioma


**BINOCULAR DIPLOPIA (DOUblE VISION USING BOTH EYES)**

1. Intractable postoperative diplopia
   - Anomalous retinal correspondence with or without amblyopia (common), which is called *paradoxical diplopia*
   - Cyclotropia due to oblique muscle operation
   - Following surgical treatment of retinal detachment because of symblepharon or limitation of extraocular movement
   - "Horror fusionis" (rare)- congenital or developmental deficiency of fusion (i.e., absence of sensory correspondence between two eyes (not the same as abnormal retinal correspondence, because visual directions are normal in these cases)
   - Large surgical overcorrection
2. Other
   - Aniseikonia, including association with macular disease
   - Divergence paresis
   - Heterophoria-due to lesions such as orbital tumor and cellulitis
   - Narcolepsy
   - Physiologic diplopia
   - Psychogenic causes
3. Paralysis of one or more extraocular muscles
   - Fourth-nerve palsy (rare) (see p. 158)
   - Sixth-nerve palsy-has no localizing value (see p. 161)
   - Third-nerve palsy-with isolated muscle paralysis one must suspect a nuclear lesion (hemorrhage, syphilis, multiple sclerosis) or myasthenia gravis (see p. 153)


**BINOCULAR TRIPOPIA (UNIOcular DIPLOPIA)**

1. Abnormal retinal correspondence with single image given two associations of direction so that the abnormal retinal point is brought into consciousness at the same time as the macula image
2. Central uniocular diplopia (rare)-systemic or neurologic causes include cerebral aneurysm, abscess or gross degenerative lesions, encephalitis lethargica, postencephalitis, multiple sclerosis, basal meningitis, cerebellar tumor, and vertebrobasilar insufficiency

*3. Malingering, hysteria, or psychogenic causes

4. Optical causes external to the eye
   A. Double or single prism placed in center of pupil before one eye
   *B. Improper correction of a high astigmatism
   *C. Looking through the edge of a bifocal or margin of lens

5. Optical causes in the eye
   A. Air bubbles or transparent foreign bodies in aqueous or vitreous
   B. Complete or partial contraction of the eyelids in which the eyelids impinge on the cornea (de Schweintz)
   C. Dislocation of the lens or misalignment of corneal and lenticular optical axis
   D. Double pupil
   E. High myopia, probably because of irregular astigmatism
   F. Irregular astigmatism, such as pressure on the globe
   G. Irregular spasm of the ciliary muscle
   H. Keratoconus (see p. 288)
   I. Lens abnormalities, such as fluid clefts or incipient cataract
   J. Looking through edge of intraocular lens
   *K. Map-dot fingerprint dystrophy
   L. Megalocornea (see p. 255)
   M. Migration of filtering bleb into the cornea
   N. Multifocal intraocular lens
   O. Post iridectomy
   *P. Refractive surgery
   Q. Retinal detachment
   R. Spherophakia (see p. 400-401)


**DIPLOPIA FOLLOWING HEAD TRAUMA**

1. Avulsion, contusion, or transection of extraocular muscles
2. Avulsion of the pulley of the superior oblique
3. Decompensation of a preexisting ocular phoria, becoming a tropia
4. Edema or detachment of the macula (monocular diplopia)
*5. Hematoma in the orbit or the ocular muscles
*6. Orbital fracture (particularly blowout fracture of the floor, causing restricted function of inferior rectus and inferior oblique muscles)
7. Subluxation of the lens (monocular diplopia)
8. Third, fourth, or sixth cranial nerve palsies (orbital or intracranial) (see pp. 158, 161, 153)
9. "Whiplash" injury and the diplopias of obscure origin


**ECCENTRIC VISION**

In this condition, vision is best when the individual is not looking directly at object of regard.

1. Central scotoma
2. Craniopharyngioma
3. Eccentric fixation with amblyopia
4. Ectopic macula, such as macula displaced by retinal scarring or fibrous strands, often a result of retinopathy of prematurity
5. Glaucoma-late with only eccentric field remaining
6. Homonymous hemianopia with macular involvement (see p. 613)
*7. Macular scar, such as with age-related macular degeneration.


**DECREASED DARK ADAPTATION (NYCTALOPIA; NIGHT BLINDNESS)**

1. Choroideremia
2. Congenital night blindness
*3. May be due to drugs, including the following:
   - alcohol
   - amodiaquine
   - cantanidin
   - carbon dioxide
   - chloroquine
   - colloidal silver
   - deferoxamine
   - dronabinol
   - ergonovine
   - ergotamine
   - etretinate
   - hashish
   - hydroxychloroquine
   - indomethacin (?)
   - isotretinoin
   - lithium carbonate
   - lysergic acid
   - diethylamide (LSD)
   - lysergide
   - marihuana
   - mescaline
   - methysergide
   - oxygen
   - pilocarpine
   - psilocybin
   - silver nitrate
   - silver protein
   - tetrahydrocannabinol
   - TI-iC
   - vinblastine
   - vincristine
4. Progressive cone-rod dystrophy
5. Refsum syndrome (heredopathia atactica polyneuritiformis syndrome)
6. Retinitis pigmentosa (see p. 497)


**ASTIGMATISM**

In this condition, the refractive power of the eye varies along different meridians; its steepest meridian is vertical in "with the rule" (corrected with plus cylinder at 90 degrees) and horizontal in “against the rule."  

1. Adnexal masses
2. Anterior segment surgery for cornea, lens, or glaucoma
   *3. Chalazion
   *4. Contact lens wear
   *5. Corneal scars
   *6. Following refractive surgery
7. Keratoconus (see p. 288)
8. May be dominant inheritance with incomplete penetrance
9. Nuclear cataract with coloboma of lens, iris, and choroid
10. Oversized, rigid, anterior chamber, intraocular lens implant
11. Physiologic—about 0.5 diopters of "with the rule"
12. Retinal detachment procedures
13. Scleral infolding
14. Tilted intraocular lens


**VISUAL ALLESTHESIA**

This condition involves displacement of image to opposite half of the visual field.

1. Parietooccipital lobe disease
   A. Neoplasm
   B. Vascular insufficiency
   C. Trauma
   D. Seizure activity

2. Occipital- lobe disease
   A. Neoplasm
   B. Vascular insufficiency
   C. Trauma
   D. Seizure activity


**VISUAL ACUITY LOSS AFTER GLAUCOMA SURGERY**

1. Cystoid macular edema
*2. Hypotony maculopathy
3. Intraocular pressure spike
*4. Lens opacification
5. Postoperative capsule opacity
6. Retinal detachment
7. Suprachoroidal hemorrhage
8. Unknown
9. Vitreous hemorrhage
10. Wipeout (loss of central fixation)


**SUDDEN PAINFUL LOSS OF VISION**
1. Acute-angle closure glaucoma
2. Fracture of the lesser wing of the sphenoid bone
3. Keratoconus
4. Optic neuritis
5. Temporal arteritis
6. Uveitis


**SUDDEN PAINLESS LOSS OF VISION-BOTH EYES**

1. Brain injury
2. Brainstem arteriovenous malformations
3. Meningeal carcinomatosis
4. Quinine poisoning
5. Wood alcohol poisoning (methyl)


**GRADUAL PAINLESS LOSS OF VISION**

1. Age-related macular degeneration
2. *Bartonella henselae*
3. Behcet disease
4. Cardiolipin antibody syndrome
5. Cataract
6. Chronic corneal disease
7. Diabetic retinopathy
8. Eales disease
9. Glaucoma, open-angle
10. Herpetic viral infection
11. Idiopathic causes
12. Optic neuropathy/atrophy
13. Refractive error
14. Retinal disease, chronic
15. Systemic lupus erythematosus
16. Wegener granulomatosis

Visual Complaint

CONTENTS

Photopsia 644
Hallucinations 645
"Spots" before eyes 648
Colored halos around lights 648
Light streaks 649
Photophobia 650
Asthenopia 652
Dazzling or glare discomfort 652
Chromatopsia 653
Heightened color perception 656
Nyctalopia (night blindness) 656
Hemeralopia (day blindness) 657
Oscillopsia 658
Color blindness 658
Palinopsia 661
Vertical reading 662
Visual agnosia 662
Ocular lateropulsion 662
Pain in and about eye 662
Headache 663
Pulfrich phenomenon 664

PHOTOPSIA (SCINTILLATIONS, SPARKS, OR FLASHES OF LIGHT BEFORE THE EYES)

1. Associated with arteriovenous aneurysm
2. Auditory-visual synesthetic phenomena-optic nerve lesion; usually demyelinative
3. Brain concussion
4. Clomiphene citrate (Clomid)
5. Focal lesions of occipital region
6. Glaucoma
7. Idiopathic thrombocytosis
8. Impending retinal detachment
9. Lyme borreliosis
*10. Migraine and epilepsy
11. Moore lightning streak-traction of a partially liquefied vitreous on the retina
12. Oculodigital phenomenon (entopic phenomenon)
13. Paraneoplastic retinopathy
14. Phosphene of quick eye motion (Flick phosphene)
15. Retinal microembolization
16. Retinitis
17. Vertebral basilar insufficiency


**HALLUCINATIONS** *(FORMED IMAGES)*

1. Blind persons (central or peripheral visual field loss)
2. Bilateral eye covering—such as may be required after an eye operation, especially in older patients
3. Ocular lesions, such as retinal hemorrhage, glaucoma, optic atrophy of tertiary syphilis, and choroidal neovascularization.
4. Psychoses
5. Central nervous system lesions
   A. Alzheimer disease
   B. Diffuse irritative lesion of parietotemporal area, including uncinate seizures of the temporal lobe, stimulation of superior colliculus, and optic radiation
   C. Encephalitis
   D. Hippocampus lesions
   E. Hypophyseal duct tumors
   F. Measles
   G. Medulloblastoma
   H. Myxedema
   I. Narcolepsy
   J. Occipital lobe seizures—moving lights and colors, visual and complex hallucinations with formed images
   K. Papilledema (see p. 593)
   L. Peduncular hallucinations with midbrain lesions from vascular, encephalitic, and mass lesions
   M. Pellagra
   N. Pituitary and optic chiasmal lesion
   O. Vertebrobasilar insufficiency/basilar artery migraine
6. Chronic mountain sickness (Monge syndrome)
7. Malignant melanoma
8. Poisonings, such as mushroom, psilocin, cannabis, hashish, hemp, camphor, mescaline from peyote, myristica (nutmeg), gasoline, mullet (Hawaiian fish), and ololiuqui (morning-glory seeds)
9. Drugs, including the following:
   - acebutolol
   - acetaminophen
   - acetanilid
   - acetophenazine
   - acid bismuth sodium
   - acyclovir
   - adrenal cortex injection
   - albuterol
   - titrate
   - alcohol
   - aldosterone
   - allobarbital
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procyclidine  sulfamethizole  tobramycin
promazine  sulfamethoxazole  tocainide
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psilocybin  sulfathiazole  trifluoperazine
pyrilamine  sulfisoxazole  trifluperidol
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quinethazone  talbutal  trihexyphenidyl
quinidine  temazepam  trimeprazine
quinine  tetanus immune globulin  trimipramine
radioactive iodides  tetanus toxoid  tripleennamine
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secobarbital  thiabendazole  urea
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sulfachlorpyridazine  thiopental  verapamil
sulfacytine  thiopropazate  vidarabine
sulfadiazine  thioproprazine  vinbarbital
sulfadimethoxine  thioridazine  vinblastine
sulfamerazine  thyroglobulin  vincristine
sulfameter  thyroid  vitamin D

10. Exercise induced with occipital lobe tumor
11. Patients in seclusion


"SPOTS" BEFORE EYES (DOTS OR FILAMENTS THAT MOVE WITH MOVEMENT OF EYE)
1. Vitreous opacities-muscae volitantes; associated with preretinal hemorrhage, myopia, posterior vitreous detachment, or intraocular inflammations
2. Scotomatous defects
   A. Retinal lesions
   B. Myopia
3. Corneal foreign-body reflection/corneal opacity
4. Carbon tetrachloride poisoning
5. Migraine


**COLORED HALOS AROUND LIGHTS (BLUE AND VIOLET ARE NEXT TO THE STIMULATING LIGHT AND RED OUTERMOST)**

1. Glaucoma
   A. Acute-angle closure with stretching of corneal lamellae
   B. Open-angle glaucoma-halo noted on awakening (intraocular pressure is highest in the morning)
2. Mucus on the cornea
3. Corneal scar/corneal edema
4. Krukenberg spindle
5. Lens opacities
6. Vitreous opacities (see p. 429-430)
7. Any haze of ocular media
8. Drugs probably affecting corneal epithelium, including the following:
   acetophenazine  ethopropazine  paramethadione
   acetyldigitoxin  ethylene diamine  perazine
   amiodarone  fluorometholone  pericyazine
   amodiaquine  fluphenazine  perphenazine
   amyl nitrite  gitalin  piperacetic
   butaperazine  hydrocortisone  prednisolone
   carphenazine  hydroxy chloroquine  prochlorperazine
   chlorine dioxide  lanatoside C  promazine
   chloroquine  medrysone  promethazine
   chlorpromazine  mesoridazine  propiomazine
   cortisone  methdilazine  quinacrine
   deslanoside  methotrimetazine  sildenafil
   dexamethasone  methylprednisolone  thiethylperazine
   diethazine  nitroglycerin  thiopropazate
   digitalis  nitronaphthalene  thioproperazine
   digitoxin  oral contraceptives  thioridazine
   digoxin  ouabain  trifluoperazine
triflupromazine  trimethadione
trimeprazine  water (sterile)

9. Physiologic halos—most common when lens acts as diffracting gradient
10. Too intense exposure to light, as in snow blindness
11. Asymmetric placement of the intraocular lens in relation to the pupillary aperture


LIGHT STreaks

1. Cataracts
2. Contact lenses
3. Excessive tear meniscus
4. Intraocular lens scratches
5. Lashes
*6. Migraine
7. Posterior capsules-lens fibers and debris-filled corrugations
8. Rapid eye movements (especially in a dark environment)
9. Reflection off edge of intraocular lens
10. Reflection off manipulation holes of intraocular lens
*11. Retinal break/tear or detachment
12. Spectacles
13. Windshields, windows


PHOTOPHOBIA (PAINFUL INTOLERANCE OF THE EYES TO LIGHT)

1. Aniridia
*2. Ocular, including conjunctivitis, keratitis, iritis, iridocyclitis, uveitis, and corneal, lenticular, and vitreous opacities, Angelucci syndrome (critical allergic conjunctivitis), acute hemorrhagic conjunctiva, and cone-dysfunction syndrome
3. Albinism
4. Total color blindness (achromatopsia)
5. Patients with corneal lesions having diseases characterized by photosensitization (xeroderma pigmentosa, hydroa vacciniforme, and smallpox)

6. Systemic diseases, including botulism, cystinosis, erythropoietic porphyria, hypoparathyroidism, rabies, psittacosis, and schistosomiasis

7. Toxic causes, including mercury poisoning

8. Drugs, including the following:

- acetohexamide
- acetophenazine
- adiphenine
- adrenal cortex injection
- allobarbital
- alprazolam
- ambutonium
- amiodarone
- amitriptyline
- amobarbital
- amodiaquine
- anisotropine
- antazoline
- atropine
- auranofin
- aurothioglycanide
- beclomethasone
- belladonna
- bendroflumethiazide
- benzthiazide
- betamethasone
- botulinum A toxin
- brimonidine tartrate
- brinzolamide
- bromide
- brompheniramine
- butaperazine
- capecitabine
- captopril (?)
- carbon dioxide
- carphenazine
- chlordiazepoxide
- chloroquine
- chlorpromazine
- chlorpropamide
- chloretetracycline
- cimetidine
- clidinium
- clomiphene
- clonazepam
- clorazepate
- dapiprazole
- dehydrochloride
- deferoxamine
- demeclocycline
- desipramine
- dextran
- dextrothyroxine
- diacetylmorphine
- diazepam (?)
- dicyclomine
- diethazine
- dipivalyl epinephrine
- diethazine
- dipivalyl epinephrine
- (DPE)
- dipivefrin
- disopyramide
- doxepin
- doxycycline
- dronabinol
- edrophonium
- enalapril
- ethambutol
- ethionamide
- ethosuximide
- ethotoin
- ferrocholinate
- ferrous fumarate
- ferrous gluconate
- ferrous succinate
- ferrous sulfate
- flecainide
- fleroxidine
- fludrocortisone
- fluoromethylone
- fluorouracil
- fluoxetine hydrochloride
- fluphenazine
- fluprednisolone
- flurazepam
- fluvoxamine maleate
- gold Au 198
- gold sodium thiomalate
- gold sodium thiosulfate
- guanethidine
- halazepam
- hashish
- hexethal
- hexobarbital
- hexocyclium
- homatropine
- hydralazine
- hydroxyamphetamine
- hydroxychloroquine
- ibuprofen
- imipramine
- indomethacin
- iron dextran
- iron sorbitex
- isocarboxazid
- isoniazid
- isopropamide
- labetalol
- latanoprost
- levaterenol
- levothyroxine
- liothyronine
- liotrix
- lithium carbonate
- lorazepam
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9. Normal ocular findings with photophobia

A. Trigeminal neuralgia (Charlin syndrome)
B. Migraine
C. Neurasthenia
D. Meningitis
E. Subarachnoid hemorrhage
F. Acromegaly
G. Associated with hypophyseal tumor and craniopharyngioma
H. During and following retrobulbar neuritis
I. Acrodynia (Feer syndrome)
J. Following severe head injury
K. Hypoparathyroidism
L. Lesions of gasserian ganglion
M. Tumors of ophthalmic branch of the trigeminal nerve, such as neuroma, middle fossa tumor, and posterior fossa tumor, such as meningioma or acoustic neuroma
N. Increased intracranial pressure, including subdural hematomas

10. Acrodermatitis chronic a atrophicans
11. Avitaminosis B (pellagra)
12. Chédiak-Higashi syndrome (anomalous leukocytic inclusions with constitutional stigmata)
13. Danbolt-Closs syndrome (acrodermatitis enteropathica)
14. Elschnig syndrome (meibomian conjunctivitis)
15. Feer syndrome (acrodynia)
16. Falling syndrome (phenylketonuria)
17. Following refractive surgery
18. Gradensigo syndrome (temporal syndrome)
19. Hanhart syndrome (pseudoherpetic keratitis)
20. Hartnup syndrome (niacin deficiency)
21. Hysteria
22. Infantile globoid cell leukodystrophy (Krabbe disease)
23. Keratodermia palmaris et plantaris
24. Photosensitivity and sunburn
25. Reiter syndrome (polyarthritis enterica)


ASTHENOPIA (UNCOMFORTABLE OCULAR SENSATION OR EYE ACHE)

1. Dazzling from bright light
2. Episcleritis or scleritis
3. Glaucoma
4. Iritis or iridocyclitis
5. Neurasthenia or hysteria
6. Passive congestion
7. Phoria or tropia
8. Retrobulbar neuritis
9. Sinus disease
10. Spasm from muscles held too long in a restricted position
11. Subclinical open-angle glaucoma
12. Uncorrected refractive errors, especially hyperopia or astigmatism
13. Unknown
14. Use of miotics
15. Weak accommodation


DAZZLING OR GLARE DISCOMFORT

1. Altered pupillary response
2. Asymmetric placement of the intraocular lens in relation to the pupillary aperture
3. Corneal scars or foreign bodies
4. Drugs, such as chloroquine, acetazolamide, or trimethadione (Tridione)
5. Emotional disorders
6. Following refractive surgery
7. Idiopathic
8. Lenticular changes


CHROMATOPSISIA (COLORED VISION, OBJECTS ARE ABNORMALLY COLORED)

1. Blue color (cyanopsia)
   A. Drugs, including the following:
      acetyldigitoxin deslanoside methylene blue
      alcohol digitalis nalidixic acid
      amodiaquine digitoxin oral contraceptives
      amphetamine digoxin ouabain
      chloroquine gitalin quinacrine
      combination products of hydroxyamphetamine sildenafil citrate
         estrogens and hydroxychloroquine
         progestogens lanatoside C

   B. Pseudophakia
   C. Optic atrophy of tertiary syphilis

2. Red color (erythropsia)
   A. Drugs, including the following:
      acetyldigitoxin gitalin sulfachlorpyridazine
      atropine homatropine sulfacytine
      belladonna lanatoside C sulfadiazine
      deslanoside methylergonovine sulfadimethoxine
      digitalis methysergide sulfamerazine
      digitoxin naproxen sulfameter
      digoxin ouabain sulfamethazine
      ergonovine quinine sulfamethizole
      ergotamine sulfacetamide sulfamethoxazole
sulfamethoxypyridazine  sulfapyridine  sulfisoxazole
sulfanilamide  sulfasalazine  sulthiame
sulfaphenazole  sulfathiazole

B. Hysteria
C. Optic atrophy of tertiary syphilis
D. Vitreous or retinal hemorrhage (see p. 473-478)
E. Pseudophakia and aphakia
F. Snow blindness or blindness following electric shock
G. After working with green monochrome video display terminals
H. Welding arc maculopathy

3. Yellow color (xanthopsia)
   A. Drugs, including the following:
   acetaminophen  deserpidine  mild silver protein
   acetanilid  deslanoside  nalidixic acid
   acetophenazine  diethazone  nitrofurantoin (?)
   acetyldigitoxin  digitalis  ouabain
   allobarbital  digitoxin  pentobarbital
   alseroxylon  digoxin  pentylenetetrazol
   amobarbital  dronabinol  perazine
   amodiaquine  ethopropazine  pericyazine
   amyl nitrite  fluorescein  perphenazine
   aprobarbital  fluphenazine  phenacetin
   aspirin  furosemide  phenobarbital
   barbital  gitalin  piperoxetine
   bendroflumethiazide  hashish  polythiazide
   benzthiazide  heptabarbital  primidone
   butabarbital  hexethal  pro barbital
   butalbital  hexobarbital  prochlorperazine
   butallylonal  hydrochlorothiazide  promazine
   butaperazine  hydroflumethiazide  promethazine
   butethal  hydroxychloroquine  propiomazine
   carbachol (?)  indapamide  quinethazone
   carbon dioxide  lanatoside C  quinacrine
   carphenazine  manjuana  rauwolfia serpentina
   chloramphenicol  mepobarbital  rescinnamine
   chloroquine  mesoridazine  reserpine
   chlorothiazide  methaqualone  secobarbital
   chlorpromazine  metharbital  silver nitrate
   chlortetracycline  methazolamide  silver protein
   chlorthalidone  methdilazine  sodium salicylate
   cimetidine (?)  methitural  streptomycin
   colloidal silver  methohexitol  sulfacetamide
   cyclobarbital  methotrimepazine  sulfachlorpyridazine
   cyclopentobarbital  methyclothiazide  sulfacytine
   cyclothiazide  metolazone  sulfadiazine
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<thead>
<tr>
<th>Drug Name</th>
<th>Drug Name</th>
<th>Drug Name</th>
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<tbody>
<tr>
<td>sulfadimethoxine</td>
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<td>thiopropazine</td>
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<td>sulfamerazine</td>
<td>sulfisoxazole</td>
<td>thioproperazine</td>
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<tr>
<td>sulfameter</td>
<td>syrosingopine</td>
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<td>sulfamethazine</td>
<td>talbutal</td>
<td>trichlonnethiazide</td>
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<td>trifluoperazine</td>
</tr>
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<td>sulfamethoxazole</td>
<td>tetrahydrocannabinol</td>
<td>triflupromazine</td>
</tr>
<tr>
<td>sulfamethoxypyridazine</td>
<td>(THC)</td>
<td></td>
</tr>
<tr>
<td>sulfanilamide</td>
<td>thiabendazole</td>
<td>vinbarbital</td>
</tr>
<tr>
<td>sulfaphenazole</td>
<td>thiamylal</td>
<td>vitamin A</td>
</tr>
<tr>
<td>sulfapyridine</td>
<td>thiethylperazine</td>
<td></td>
</tr>
<tr>
<td>sulfasalazine</td>
<td>thioptal</td>
<td></td>
</tr>
</tbody>
</table>

B. Lenticular change
C. Aphakia
D. Poisons, including aconite, dichlorodiphenyl trichloroethane, carbon disulfide, chromic acid, methyl salicylate, aspidium (Felix mas), santonin, picric acid
E. Jaundice
F. Hysteria

4. Green color (chloropsia)
A. Drugs, including the following:

<table>
<thead>
<tr>
<th>Drug Name</th>
<th>Drug Name</th>
<th>Drug Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>acetyldigitoxin</td>
<td>digoxin</td>
<td>nalidixic acid</td>
</tr>
<tr>
<td>allobarbital</td>
<td>epinephrine</td>
<td>naproxen</td>
</tr>
<tr>
<td>amobarbital</td>
<td>gitalin</td>
<td>ouabain</td>
</tr>
<tr>
<td>amodiaquine</td>
<td>griseofulvin</td>
<td>pentobarbital</td>
</tr>
<tr>
<td>aprobarbital</td>
<td>heptabarbital</td>
<td>phenobarbital</td>
</tr>
<tr>
<td>barbital</td>
<td>hexethal</td>
<td>primidone</td>
</tr>
<tr>
<td>butabarbital</td>
<td>hexobarbital</td>
<td>probarbital</td>
</tr>
<tr>
<td>butalbital</td>
<td>hydroxychloroquine</td>
<td>quinacrine</td>
</tr>
<tr>
<td>butallylonal</td>
<td>iodide and iodine</td>
<td>quinine</td>
</tr>
<tr>
<td>butethal</td>
<td>solutions and</td>
<td>radioactive iodides</td>
</tr>
<tr>
<td>chloroquine</td>
<td>compounds</td>
<td>secobarbital</td>
</tr>
<tr>
<td>cyclobarbital</td>
<td>lanatoside C</td>
<td>talbutal</td>
</tr>
<tr>
<td>cyclopentobarbital</td>
<td>mephobarbital</td>
<td>thiamylal</td>
</tr>
<tr>
<td>deslanoside</td>
<td>metharbital</td>
<td>thiopental</td>
</tr>
<tr>
<td>digitalis</td>
<td>methitural</td>
<td>vinbarbital</td>
</tr>
<tr>
<td>digitoxin</td>
<td>methohexital</td>
<td></td>
</tr>
</tbody>
</table>

B. Poisons such as santonin

5. Violet color (ianthinopsia)
A. Drugs, including the following:

<table>
<thead>
<tr>
<th>Drug Name</th>
<th>Drug Name</th>
<th>Drug Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>dronabinol</td>
<td>marihuana</td>
<td>quinacrine</td>
</tr>
<tr>
<td>hashish</td>
<td>nalidixic acid</td>
<td>tetrahydrocannabinol</td>
</tr>
</tbody>
</table>

B. Pseudophakia and aphakia
C. Intracameral air

6. Brown color
### A. Drugs, including the following:

<table>
<thead>
<tr>
<th>Acetophenazine</th>
<th>Methdilazine</th>
<th>THC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Butaperazine</td>
<td>Perazine</td>
<td>Thiethylperazine</td>
</tr>
<tr>
<td>Carphenazine</td>
<td>Pericyazine</td>
<td>Thiopropazate</td>
</tr>
<tr>
<td>Chlorpromazine</td>
<td>Perphenazine</td>
<td>Thioproperazine</td>
</tr>
<tr>
<td>Diethazine</td>
<td>Piperacetazine</td>
<td>Thioridazine</td>
</tr>
<tr>
<td>Ethopmpazine</td>
<td>Prochlorperazine</td>
<td>Trifluoperazine</td>
</tr>
<tr>
<td>Fluphenazine</td>
<td>Promazine</td>
<td>Triflupromazine</td>
</tr>
<tr>
<td>Inethotrimeprazine</td>
<td>Promethazine</td>
<td>Trimeprazine</td>
</tr>
<tr>
<td>Mesoridazine</td>
<td>Propiomazine</td>
<td></td>
</tr>
</tbody>
</table>

### B. Lenticular change

#### 7. White color

<table>
<thead>
<tr>
<th>Capreomycin</th>
<th>Paramethadione</th>
<th>Phenytin</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diphenylhydantoin</td>
<td>Trimethadione</td>
<td></td>
</tr>
</tbody>
</table>

### B. Pseudophakia and aphakia


### HEIGHTENED COLOR PERCEPTION

1. Heightened color perception is due to drugs, including the following:

<table>
<thead>
<tr>
<th>Dronabinol</th>
<th>Lysergide</th>
<th>Psilocybin</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ethionamide</td>
<td>Marihuana</td>
<td>Tetrahydrocannabinol</td>
</tr>
<tr>
<td>Hashish</td>
<td>Mescaline</td>
<td>THC</td>
</tr>
<tr>
<td>LSD</td>
<td>Oxygen</td>
<td></td>
</tr>
</tbody>
</table>


### NYCTALOPIA (NIGHT BLINDNESS)

1. Anemia
2. Carbon monoxide poisoning
3. Congenital high myopia
4. Diffuse opacities of media including corneal edema, keratitis and nuclear cataract
5. Following refractive surgery
6. Glaucoma—especially open-angle and angle-closure glaucoma
7. Paraneoplastic retinopathy including melanoma-associated retinopathy and cancer-associated retinopathy
8. Psychologic causes-malingering or psychoses
9. Optic atrophy
10. Refsum syndrome (phytanic acid oxidase deficiency)
11. Tapetoretinal degenerations
    A. Choroideremia
    B. Congenital night blindness
       (1) Dominant form
       (2) Recessive form
       (3) Recessive, sex-linked
    C. Detachment of retina, including malignant melanoma
    D. Drugs, including the following:
       acetophenazine mesoridazine propiomazine
       amodiaquine methdilazine quinidine
       butaperazine methotrimedazine quinine
       carphenazine paramethadione thiethylperazine
       chloroquine perazine thiopropazate
       chlorpromazine pericyazine thioproperazine
       diethazine perphenazine thioridazine
       ethopropazine piperacetazine trifluoperazine
       fluphena zine prochlorperazine triflupromazine
       hydroxychloroquine promazine trimeprazine
       indomethacin promethazine trimethadione
    E. Drusen (familial)-minimal
    F. Fleck retina-nonprogressive, congenital, rare
    G. Fundus flavimaculatus-minimal
    H. General choroidal sclerosis
    I. Gyrate atrophy
    J. Retinitis pigmentosa
    K. Retinitis punctata albescens
    L. Minar nystagmus
    M. Oguchi disease-may be abnormal
12. Visual field defects
13. Vitamin A deficiency
    A. Dietary deficiencies, including malnutrition, alcoholism and cystic fibrosis
    B. Digestive tract disturbance
       (1) Colitis and enteritis
       (2) Crohn disease
       (3) Jejunoileal bypass surgery
       (4) In pancreas-such as chronic pancreatitis
       (5) In stomach-achlorhydria, chronic gastritis or diarrhea, peptic ulcer
       (6) Abetalipoproteinemia
    C. Liver disease, such as chronic cirrhosis
D. Malaria
E. Pregnancy
F. Pulmonary tuberculosis
G. Skin disorders, such as pityriasis rubra pilaris
H. Thyroid gland disorders, such as hyperthyroidism

14. Vitreous opacities, including hemorrhage
15. Vitreotapetoretinal degeneration-sex-linked recessive and autosomal recessive


**HEMERALOPIA**

This condition involves day blindness, that is, an inability to see as distinctly in a bright light as in a dim one.

1. Adie pupil
2. Albinism
3. Aniridia
4. Central opacities of the lens-nuclear or perinuclear cataracts
5. Central scotoma
6. Congenital-autosomal recessive trait usually associated with amblyopia and color deficiency
7. Hereditary retinoschisis
8. Intraocular iron
9. Partial occlusion of the central retinal artery (see p. 457-461)
10. Refsum syndrome (phytanic acid oxidase deficiency)
11. Total color blindness


**OSCILLOPSIA**

This condition involves illusionary movement of the environment; it may be unilateral or bilateral and usually occurs because of acquired nystagmus.
1. Drugs, including the following:
   - Alcohol
   - Allobarbital
   - Amobarbital
   - Aprobarbital
   - Barbital
   - Butabarbital
   - Butalbital
   - Butallylonal
   - Butethal
   - Carbamazepine
   - Cyclobarbital
   - Diphenylhydantoin
   - Gentamicin
   - Heptabarbital
   - Hexothal
   - Hexobarbital
   - Mepobarbital
   - Metharbital
   - Methitural
   - Methohexital
   - Pentobarbital
   - Phenobarbital
   - Primidone
   - Probabtal
   - Secobarbital
   - Talbutal
   - Thiamylal
   - Thiopental
   - Valproate sodium
   - Valproic acid
   - Vinbarbital

2. Fixation and voluntary nystagmus
3. Defective vestibuloocular reflex/vestibular pathway lesion occurs during movement of the head or body
   - A. Sectioning of vestibular (VIII) nerve for vertigo
   - B. Streptomycin toxicity
   - C. Spontaneous loss
4. Head trauma/seizures
5. Intermittent exotropia
6. Involvement of medial longitudinal fasciculus affecting ipsilateral medial rectus in internuclear ophthalmoplegia-monocular oscillopsia
7. Myokymia of the eyelid
8. Opsoclonus and ocular flutter
9. Vertebral artery dissection


**COLOR BLINDNESS**

1. Inherited-stable defect, affecting both eyes
   - A. Bassen-Kornzweig syndrome (abetalipoproteinemia)
   - B. Congenital dyslexia syndrome (developmental dyslexia syndrome)
C. Down syndrome (mongolism)
D. Duane retraction syndrome (Stilling syndrome)
E. Duchenne muscular dystrophy
F. Glucose-6-phosphate dehydrogenase deficiency (glycogen storage disease type I)
G. Guillain-Barré syndrome (acute infectious neuritis)
H. Hemophilia
I. "Intrinsic" defect
   (1) Dichromat—two colors mixed to see white
      a. Deuteranope—green deficiency
      b. Protanope—red deficiency
      c. Tritanope—blue deficiency
   (2) Monochromat—one color mixed to see white
      a. Cone deficient
      b. Rod deficient
   (3) Trichromat—three colors mixed to see white
      a. Deuteranomaly—green anomaly
      b. Protanomaly—red anomaly
      c. Tritanomaly—blue anomaly
J. Kallman syndrome (hypogonadotrophic hypogonadism—anosmia syndrome)
K. Klinefelter syndrome (XXY) (gynecomastia-aspermogenesis syndrome)
L. Turner syndrome (XO) (gonadal dysgenesis)
2. Acquired-defect can increase or decrease; may affect only one eye; impairment of other visual function; often characterized by chromatopsia; hue discrimination primarily affected; yellow-blue defects more common in retinal disease; red-green defects in optic nerve disease
   A. Advanced hypertensive retinopathy
   B. Albinism
   C. Amblyopia
   D. Blue-yellow defect with retinal disorders from drugs, including the following:
      acetophenazine clofazimine methotrimeprazine
      amiodarone (?) clonidine (?) minoxidil (?)
      amodiaquine cobalt (?) mitotane
      azathioprine cycrimine (?) naproxen (?)
      benztropine (?) deeroxamine penicillamine
      biperiden (?) diethazine perazine
      butaperazine diethylcarbamazine pericyazine
      carbamazepine ethambutol perphenazine
      carphenazine ethopropazine piperacetazine
      cephaloridine (?) fluphenazine prazosin (?)
      chloramphenicol hydroxychloroquine prochlorperazine
      chloroquine indomethacin (?) procyclidine (?)
      chlorphenoxamine (?) ketoprofen (?) promazine
      chlorpromazine mesoridazine promethazine
      chlorprothixene methdilazine propiomazine
      cisplatin methotrexate quinacrine (?)
quinine thiopropazate trifluoperazine
sulindac (?) thiopeperazine triflupromazine
tamoxifen thioridazine trihexyphenidyl (?)
thiethylperazine thiothixene trimeprazine

E. Chorioretinitis
F. Color anomia-inability to name colors; may be associated with homonymous hemianopia resulting from infarct of posterior parietal and corpus callosum
G. Diabetic retinitis
H. Dominantly inherited juvenile optic atrophy
I. Drugs and chemical substances causing optic neuropathy with red-green defect, including the following:

acetophenazine chloroprocaine (?) hexethal
alcohol chloroquine hexobarbital
allobarbital chlorpromazine hydrocortisone
alseroxylon (?) cholecalciferol hydroxychloroquine
aminosalicylate (?) clindamycin iodide and iodine
aminosalicylic acid (?) cobalt (?) solutions and compounds
amobarbital cortisone
amodiaquine cyclobarbital iodochlorhydroxyquin
antimony lithium cyclopentobarbital iodoquinol
thiomalate cycloserine (?) iron dextran (?)
antimony potassium dapsone iron sorbitex (?)
tartrate deferoxamine isoniazid
antimony sodium deserp dine (?) levothyroxine (?)
tartrate dexamethasone lidocaine (?)
antimony sodium dextrothyroxine (?) liothyronine(?)
thioglycollate diethazine liotrix (?)
antipyrine ergocalciferol medrysone
aprobarbital ergonovine (?) mephobarbital
aspirin ergot (?) mepivacaine (?)
barbital ergotamine (?) mesoridazine
betamethasone ethambutol metharbital
bromide (?) ethopropazine methdilazine
bromisovalum etidocaine (?) methitural
broxyquinoline ferrocholinate (?) methohexital
bupivacaine (?) ferrous fumarate (?) methotrexate (?)
butabarbital ferrous gluconate (?) methotrimetrazine
butalbital ferrous succinate (?) methyl alcohol
butallylonal ferrous sulfate (?) methylene blue
butaperazine fluorometholone methylergonovine
butethal fluphenazine methylergyste (?)
calcitriol gentamicin nitroglycerin (?)
carbromal heptabarbital oxyphenbutazone
carphenazine hexachlorophene pentobarbital
chloramphenicol hexamethonium perazine
<table>
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<tr>
<th>Drug Name</th>
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<th>Drug Name</th>
</tr>
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<tr>
<td>pericyazine</td>
<td>sodium</td>
<td>sulfisoxazole (?)</td>
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<td>perphenazine</td>
<td>antimonylgluconate</td>
<td>suramin</td>
</tr>
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<td>phenobarbital</td>
<td>sodium salicylate</td>
<td>syrosingopine (?)</td>
</tr>
<tr>
<td>phenylbutazone</td>
<td>stibocaptate</td>
<td>talbutal</td>
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<td>piperacetazine</td>
<td>stibogluconate</td>
<td>thiamylal</td>
</tr>
<tr>
<td>polysaccharide-iron complex (?)</td>
<td>stibophen</td>
<td>thiethylperazine</td>
</tr>
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<td>prenisolone</td>
<td>streptomycin</td>
<td>thiopental</td>
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<td>prilocaine (?)</td>
<td>sulfacetamide (?)</td>
<td>thiopropazate</td>
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<td>primidone (?)</td>
<td>sulfachlorpyridazine (?)</td>
<td>thiopropazine</td>
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<td>probarbital</td>
<td>sulfadiazine (?)</td>
<td>thioridazine</td>
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<tr>
<td>procaïne (?)</td>
<td>sulfadimethoxine (?)</td>
<td>thyroglobulin (?)</td>
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<td>promazine</td>
<td>sulfameter (?)</td>
<td>trichloroethylene</td>
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<td>quinine</td>
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<td>sulfasalazine (?)</td>
<td>vitamin D</td>
</tr>
<tr>
<td>secobarbital</td>
<td>sulfathiazole (?)</td>
<td></td>
</tr>
</tbody>
</table>

J. Friedreich ataxia
K. Glaucoma, including narrow and open angle
L. Hepatic cirrhosis
M. Hystria
N. Macular lesions, including juvenile degeneration, senile degeneration dystrophy, and edema
O. Night blindness
P. Occlusion of retinal vessels
Q. Oguchi disease
R. Open-angle glaucoma
S. Ophthalmologist who use argon blue-green lasers or operating microscopes
T. Optic atrophy
U. Optic pathways, including brain tumor
V. Papillitis
W. Peripheral chorioretinal degeneration
X. Retinal detachment
Y. Retinitis pigmentosa
Z. Retrobulbar optic neuritis
AA. Snow blindness
PALINOPSIA

This condition involves persistence or recurrence of visual images after exciting stimulus object has been removed; the patient has a hemianopic field defect. Polyopia (visual trailing effect with movement) may be present.

1. Acute migraine
2. Demyelinative optic neuritis
3. Encephalitis
4. Epilepsy
5. Intoxications, such as mescal delirium, LSD, trazodone-induced and clomiphene citrate.
6. Kartagener syndrome
7. Laser treatment of diabetic macular edema
8. Leber hereditary optic neuropathy
9. Temporal-parietal-occipital lesion
   A. Degenerative
   B. Neoplastic
   C. Traumatic
   D. Vascular
10. Schizophrenia
11. Drug, such as nefazodone (akinetopsia-persistence of moving objects)


**VERTICAL READING (PATIENT READS FROM ABOVE DOWNWARD)**

1. Astigmatism—high error of refraction
2. Homonymous hemianopia (see p. 613)
3. Voluntary as oriental script written vertically


**VISUAL AGNOSIA**

This condition involves a failure to recognize objects by sight for animate and inanimate objects, but it does not interfere with recognition of language symbols.

1. Drugs
2. Kluver-Bucy syndrome (temporal lobectomy behavior syndrome)
3. Lesion of Brodmann area 18


**OCULAR LATEROPULSION**

The eyes feel as though they are being drawn toward one side, but this problem can be overcome with conscious effort; full range of extraocular muscle movements is maintained.

1. Lateral medullary disease, including infarction of lateral medullary plate, acoustic neurinoma, posterior fossa meningioma, or multiple sclerosis
2. Peripheral vestibular disease


**PAIN IN AND ABOUT EYE**

1. Ocular
   A. Angle-closure glaucoma
   B. Chronic ocular hypoxia, carotid occlusive disease
   C. Dry-eye and tear-deficiency syndrome
   D. Local lid, conjunctival, and anterior segment disease
   E. Ocular inflammation including lyme borreliosis
2. Ophthalmic division
   A. Herpes zoster
   B. Migraine, cluster headache
   C. Painful ophthalmoplegia syndrome
   D. Raeder paratrigeminal neuralgia
   E. Referred (dural) pain, including occipital infarction
   F. Sinusitis
   G. Tic douloureux (infrequent in VI)

3. Mandibular division
   A. Dental disease
   B. Tic douloureux

4. Maxillary division
   A. Dental disease
   B. Nasopharyngeal carcinoma
   C. Sinusitis
   D. Temporomandibular syndrome
   E. Tic douloureux

5. Miscellaneous
   A. Atypical facial neuralgias
   B. Cranial arteritis
   C. Pain with medullary lesions
   D. Trigeminal tumors


HEADACHE

1. Vascular headache of migraine type
   A. Cephalalgia migraine (migraine equivalent)-migraine aura without headache
   B. Classic migraine-migraine with aura
   C. Common migraine-migraine without aura
   D. Complicated migraine-hemiplegic migraine and ophthalmoplegia migraine
   E. Cluster headache
   F. Lower-half headache

2. Muscle-contraction headache

3. Combined (skeletal vascular)

4. Headache of nasal vasomotor reaction

5. Headache of delusional, conversion, or hypochondriacal states

6. Nonmigraine vascular headaches
   A. Primary or metastatic tumors of meninges, vessels, or brain
   B. Hematomas (epidural, subdural, or parenchymal)
   C. Abscesses (epidural, subdural, or parenchymal)
   D. Post lumbar puncture headache (leakage, headache)
   E. Pseudomotor cerebri and various causes of brain swelling
7. Headache due to overt cranial inflammation
   A. Intracranial disorders
      (1) Mass
      (2) Meningitis
      (3) Subarachnoid hemorrhage
   B. Extracranial disorders (temporal arteritis)
8. Headache because of diseases of ocular structures
9. Headache because of diseases of aural structures
10. Headache because of diseases of the nasal and sinus structures
11. Headache because of diseases of dental structures
12. Headache because of diseases of other cranial and neck structures
13. Cranial neuritides
14. Cranial neuralgia
   A. Glossopharyngeal neuralgia
   B. Trigeminal neuralgia
15. Analgesic/ergotamine rebound headache


**PULFRICH PHENOMENON**

This condition involves a three-dimensional illusion in which a moving object that is viewed binocularly with a light attenuating filter in front of one eye appears to transcribe an anomalous pathway.

1. Age-related macular degeneration
2. Anisocoria
   A. Induced
   B. Traumatic
3. Anisometropic amblyopia
4. Cataract
5. Central serous retinopathy
6. Corneal Opacity
7. Hemianopia
8. Multiple Sclerosis
9. Optic Neuritis
10. Postretinal detachment repair


HEAD TURN (FACE TURN)

1. Head turned toward right (gaze left)
   A. Left Brown syndrome
   B. Left inferior oblique muscle palsy
   C. Left medial rectus muscle palsy
   D. Left superior oblique muscle palsy
   E. Right Duane syndrome
   F. Right jerk nystagmus
   G. Right inferior rectus muscle palsy
   H. Right lateral rectus muscle palsy
   I. Right superior rectus muscle palsy
   J. Right supranuclear gaze paresis

2. Head turned toward left (gaze right)
   A. Left Duane syndrome
   B. Left jerk nystagmus
   C. Left inferior rectus muscle palsy
   D. Left lateral rectus muscle palsy
   E. Left superior rectus muscle palsy
   F. Left supranuclear gaze paresis
   G. Right Brown syndrome
   H. Right inferior rectus muscle palsy
   I. Right medial rectus muscle palsy
   J. Right superior oblique muscle palsy

3. Head turned toward either left or right
   A. Congenital jerk nystagmus-head turned away from field with least amplitude of nystagmus (i.e., left jerk nystagmus improves in right gaze; left head turn)
   B. Esotropia-head turned in the direction of convergent eye (cross fixation)
   C. Hearing defect
   D. One blind eye-head turn away affected side (good eye fixates in adduction)
   E. Photophobia (see p. 650)
   F. Progressive intracranial arterial occlusion syndrome (Taveras syndrome)
   G. Strabismus fixus (general fibrosis syndrome)
   H. Under corrected myope


**HEAD TILT (HEAD TILTED TOWARD EITHER SHOULDER OR AROUND AN ANTEROPOSTERIOR AXIS)**

1. Head tilted toward right
   - A. Left superior oblique muscle palsy
   - B. Left superior rectus muscle palsy
   - C. Right inferior oblique muscle palsy
   - D. Right inferior rectus muscle palsy

2. Head tilted toward left
   - A. Left inferior oblique muscle palsy
   - B. Left inferior rectus muscle palsy
   - C. Right superior oblique muscle palsy
   - D. Right superior rectus muscle palsy

3. Head tilted toward either right or left
   - A. Astigmatism
   - B. Beckwith-Wiedemann syndrome
   - C. Blowout fracture
   - D. Incorrectly aligned cylinder axis
   - E. Monocular torticollis-patching of eyes does not eliminate problem;
     roentgenogram may help
       - (1) Congenital malformation of fracture of cervical spine or vertebral processes
       - (2) Fracture of clavicle
       - (3) Functional habit and hysteria
       - (4) Pain from infection
         - a. Adenitis
         - b. Arthritis
         - c. Mastoiditis
         - d. Synovitis
       - (5) Paralysis of absent muscles on opposite side of head tilt
       - (6) Sandifer syndrome (hiatus hernia-torticollis syndrome)
       - (7) Spasm of sternocleidomastoid or contracture of sternocleidomastoid muscle on side of head tilt
       - (8) Vestibular defect
         - a. Acoustic neuroma
         - b. Labyrinthitis
c. Otitis media
   F. Nystagmus—turned away from field with least amplitude of nystagmus
   G. Superior oblique tendon sheath syndrome (Brown syndrome)

Greenberg MF, Pollard ZF. Ocular plagiocephaly: ocular torticellis with skull and facial

Kattah JC, Dagi TF. Compensatory head tilt in upbeating nystagmus. *J Clin Neuro-


**CHIN ELEVATION**

1. Adaptive symptom of contact lens wearer
2. A—esotropia with fusion in downward gaze
3. Blowout fracture of orbit
4. Brown syndrome (superior oblique tendon sheath syndrome)
5. Double elevator palsy
6. General fibrosis syndrome (strabismus fixus)
7. Incomplete bilateral ptosis
8. Inferior oblique muscle palsy
9. Parinaud syndrome (dorsal midbrain syndrome)
10. Superior rectus muscle palsy
11. Supranuclear lesion (upgaze palsy)
12. Thyroidectomy
13. "V" pattern exotropia with fusion in downward gaze


Hiatt RL, Cope-Troupe C. Abnormal head positions due to ocular problems. *Ann

**CHIN DEPRESSION**

1. "A" pattern exotropia with fusion in upward gaze (A pattern)
2. Inferior rectus muscle palsy
3. Photophobia
4. Progressive supranuclear palsy
5. Superior oblique muscle palsy (bilateral)
6. Supranuclear lesion (down gaze palsy)
7. Uncorrected myope of low degree
8. "V" pattern esotropia with fusion in upward gaze
1. Benign or familial tremor
2. Bobble head doll syndrome-to-and-fro bobbing of the head and trunk, at 2- to 3-second intervals because of cyst of third ventricle
3. Congenital nystagmus
4. Extrapyramidal dysfunction, such as paralysis agitans (Parkinson syndrome)
5. Habit spasm
6. Spasms nutans


**HEAD TREMOR**

1. Cerebellar system afflictions (benign essential senile tremor)-most common
2. Extrapyramidal disorder
3. Hereditary postural tremor (familial tremor)
4. Postural tremor


Klawans HL. Rhythmic head tremor. JAMA 1982; 248:1510.

**HEAD THRUST**

1. Oculomotor apraxia-defect or horizontal voluntary movements
2. Ataxia-telangiectasia syndrome
3. Isolated
4. Male predominance
5. Oral-facial-digital syndrome type II

* = most important