Albinism: modern molecular diagnosis

Susan M Carden, Raymond E Boissy, Pamela J Schoettker, William V Good

Albinism is no longer a clinical diagnosis. The past classification of albinism was predicated on phenotypic expression, but now molecular biology has defined the condition more accurately. With recent advances in molecular research, it is possible to diagnose many of the various albinism conditions on the basis of genetic causation. This article seeks to review the current state of knowledge of albinism and associated disorders of hypopigmentation.

The term albinism (L albus, white) encompasses genetically determined diseases that involve a disorder of the melanin system. Each condition of albinism is due to a genetic mutation on a different chromosome. The cutaneous hypopigmentation in albinism ranges from complete pigmentation of the skin and hair (as occurs in Waardenberg and Waardenberg–Hirschsprung syndrome where the melanocyte fails to populate certain areas during embryogenesis. Secondly, a mutation could affect seemingly unrelated cell types, including the melanocyte. Chediak–Higashi and Hermansky–Pudlak syndromes are disease examples.

Melanin physiology

Pigmented cells of the eye have one of two origins. The retinal pigment epithelium, the posterior and anterior iris epithelium, and the outer pigmented and inner non-pigmented (so-called) ciliary epithelium are all derived from the neuroectoderm of the primitive forebrain. On the other hand, melanocytes in the iris stroma, ciliary stroma, and choroid are derived from the neural crest. Melanoblasts from the neural crest migrate to the skin, brain, inner ear, and uveal tract to develop into melanocytes.4

The biochemical pathway converting tyrosine to melanin is shown in Figure 1. The pigment granule (melanosomes) are an intracellular organelle produced by melanocytes and into which melanin is confined. In the skin, the melanosome is later transferred from the melanocyte to the surrounding keratinocytes. The melanosome precursor arises from the smooth endoplasmic reticulum. Tyrosinase and other enzymes regulating melanin synthesis are produced in the rough endoplasmic reticulum, matured in the Golgi apparatus, and translocated to the melanosome where melanin biosynthesis occurs.

Tyrosinase is a copper containing, monophenol, mono-oxygenase enzyme that has long been known to have a critical role in melanogenesis.9 It catalyses three reactions in the melanin pathway. The rate limiting step is the hydroxylation of tyrosine into dihydroxyphenylalanine (DOPA) by tyrosinase, but tyrosinase does not act alone. There is a melanogenic complex in the melanocyte, functioning as an integrated unit, which consists of tyrosinase, tyrosinase related protein 1 (TRP-1), TRP-2, lysosome associated membrane protein 1, and melanocyte stimulating hormone receptors.5 TRP-1 and TRP-2 have multiple functions, one of which is to stabilise tyrosinase activity.6

A putative tyrosinase transporter is situated in the limiting membrane of the pre-melanosome. The P polyepitope is the product of the pink eye dilution locus (named from the mouse) and the counterpart on human chromosome 15 (15q11.2-13). It has been proposed,9 and refuted,9 that the P protein transports tyrosine into the melanosome.

Classification

See Table 1.

In the past, the classification of albinism has had a dichotomous foundation, based on whether the phenotype was oculocutaneous or simply ocular. Recently, investigations have become more sophisticated and some cases have been reclassified. For example, one third of cases initially labelled as OCA 2 can be reclassified as OCA 1B on the basis of molecular genetic studies.10 OCA 1 and OCA 2 were originally separated on the basis of the hair bulb test, which is now considered unreliable. We can now review the molecular classification of various albinism conditions.

Oculocutaneous albinism

“Oculocutaneous albinism” refers to a heterogenous group of autosomal recessive disorders in which melanin is reduced or absent. Pale skin and an increased risk of skin cancer, together with the previously mentioned ocular phenotype, are present.

OCA 1

OCA 1 results from mutations in the tyrosinase gene affecting its synthesis and/or catalytic activity.11 Three steps in melanin biosynthesis are catalysed by tyrosinase.12 OCA 1 is further divided into OCA 1A and OCA 1B. Different types of mutations of the gene for tyrosinase explain the two types of OCA 1. The gene position is on chromosome...
Tyrosinase activity is completely absent and there is no melanin in the skin or eyes (fig 2). Visual acuity is decreased to 20/400. Nonsense, frameshift, and missense mutations account for OCA1A. OCA1B has a greatly diminished, but not absent, level of tyrosinase. OCA1B individuals may exhibit an increase in skin, hair, and eye pigment with age and do tan with sun exposure (Fig 2).

Fifty five mutations of the tyrosinase (TYR) gene have been found to date. The phenotype of OCA1B is indistinguishable from OCA1A at birth. Tyrosinase activity modulates pigmentation and causes a darker skinned phenotype during childhood. OCA1B was originally described in the Amish people, who are highly inbred. The condition was formerly referred to as “yellow mutant.”

### Table 1  Conditions of albinism

<table>
<thead>
<tr>
<th>Type of albinism</th>
<th>Gene position</th>
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<tbody>
<tr>
<td>OCA1</td>
<td>11q14-21</td>
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<tr>
<td>OCA1A</td>
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<tr>
<td>OCA1B</td>
<td></td>
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<tr>
<td>OCA2</td>
<td>15q11-13</td>
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<tr>
<td>OCA3</td>
<td>9p23</td>
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<tr>
<td>OA 1</td>
<td>Xp22.3-22.2</td>
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<tr>
<td>OA 2</td>
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<tr>
<td>Chediak-Higashi</td>
<td>1q42-44</td>
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<tr>
<td>Hermansky-Pudlak</td>
<td>10q23.1-23.3</td>
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<tr>
<td>Contiguous gene syndromes</td>
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<tr>
<td>Prader-Willi (OCA 2)</td>
<td>15q11-13</td>
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<tr>
<td>Angelman (OCA 2)</td>
<td>15q11-13</td>
</tr>
<tr>
<td>Kallmann syndrome (OA 1)</td>
<td>Xp22.3-22.4</td>
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<tr>
<td>Late onset sensorineural deafness</td>
<td>Xp22</td>
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<tr>
<td>(OA 1)</td>
<td>Xp22</td>
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<tr>
<td>X-linked ichthyosis and Kallmann syndrome (OA 1)</td>
<td>Xp22</td>
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<td>Microphthalmia and linear skin defects (OA 1)</td>
<td>Xp22</td>
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<tr>
<td>Aicardi</td>
<td>Xp22</td>
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<tr>
<td>Goltz</td>
<td>Xp22</td>
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<tr>
<td>Waardenburg 1</td>
<td>2q35-q37.3</td>
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<tr>
<td>Waardenburg 2</td>
<td>3p12</td>
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<tr>
<td>Waardenburg 3</td>
<td>2q35-q37.3</td>
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<tr>
<td>Waardenburg-Hirschsprung</td>
<td>13q22</td>
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</tbody>
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OCA 1 and 1B compound heterozygotes share phenotypic attributes of OCA1A and 1B. OCA1 phenotypes can result from homozygosity or compound heterozygosity of the alleles. In white people there is no one mutant TYR allele that is responsible for most of the mutations. Thus, if the parents are not related then the patient will probably be a compound heterozygote.

An OCA1A patient must have two type OCA1A alleles; an OCA1B can have two type OCA1B alleles or one type 1A and one type 1B allele. This explains why the phenotype of OCA1B has such a broad spectrum. Even a small amount of tyrosinase activity will alter the phenotypic expression.

OCA 2 has a prevalence of 1:15 000. Patients with OCA 2 were formerly referred to as “tyrosinase positive” oculocutaneous albinos because the mutation causing OCA 2 does not affect the tyrosinase gene, instead affecting the P polypeptide. The disease is also autosomal recessive, but coded on a different chromosome from OCA 1—that
is, 15q11-13. Patients with OCA 2 generally are not as severely affected as those with OCA 1, but the phenotypic spectrum is broad, ranging from virtually no apparent pigmentation to nearly normal pigmentation. Pigmentation is absent or poor at birth and develops during childhood. Vision in OCA 2 improves with age. In young childhood, it may be low (20/100–20/200) but with accumulation of pigment vision improves (20/40–20/70). Changes in acuity can occur up to adolescence, but whether improvement is specifically related to pigmentation is unknown. That delayed maturation of vision may occur in albinism is widely accepted. The mechanism of delay in albinism is unknown.

OCA 3

In OCA 1, the gene mutation affects tyrosinase and, in OCA 2, the mutation affects the gene for P protein. OCA 3 is caused by a defect in tyrosine related protein 1 (TRP-1). TRP-1 is the product of the brown locus in the mouse. A mutant allele in the mouse at that position causes the fur to be brown rather than black. The human gene coding for TRP-1 is on the short arm of chromosome 9 (9p23). The phenotypic features of OCA 3 in black patients are light brown hair, light brown skin, blue/brown irides, nystagmus, and diminished visual acuity (Fig 3). OCA 3 has not been detected in white or Asian races, perhaps because the phenotype may not be obvious. It is autosomal recessive. Although the function of TRP-1 in human melanogenesis is not fully determined, it acts as a regulatory protein in the production of black melanin. Its mutation results in the disregulation of tyrosinase activity and the synthesis of brown rather than black melanin. In mice, TRP-1 catalytically functions as a dihydroxyindole carboxylic acid (DHICA) oxidase and may aid in stabilisation of tyrosinase and other melanosomal enzymatic processes. There may be an interaction with TRP-1 and tyrosinase to affect tyrosinase hydroxylase activity. TRP-1 may also upregulate the availability of L-DOPA during the onset of melanogenesis.

Ocular albinism

OCULAR ALBINISM 1

X linked recessive ocular albinism (Nettleship–Falls type) is categorised as ocular albinism type 1 (OA 1). It has a prevalence of 1:50 000 in the United States. The OA 1 locus is 5p22-3. Charles et al looked at 72 cases of OA 1 and found nystagmus (in all cases except one), reduced visual acuity (to 6/36 in most cases), refractive errors (especially astigmatism), fundus hypopigmentation, lack of foveal reflex, strabismus, iris translucency in all cases, and posterior embryotoxon in 30% (implying some dysgenesis of the anterior segment). The ocular phenotype of OA 1 is similar to OCA 1 and OCA 2. Affected patients have a normal skin phenotype, although it may be paler than that of first degree relatives. Patients with OA 1 may initially be diagnosed with congenital motor nystagmus. Examination of the mother usually shows signs of a carrier state—mud splattered fundus with hyperpigmented streaks in the periphery and marked iris translucency. Affected males exhibit the complete phenotype while female carriers have retinal and cutaneous signs. In particular, there is a mosaic pattern of pigmentation in the female carriers, in keeping with the hypothesis of X inactivation (lyonisation). It is, however, more distinct in the retinal periphery. Mild peripheral retinal changes are not definite indicators of carrier status, as 11% of age matched controls have similar changes. The other hand, definite changes in the peripheral retina identify at risk females as carriers quite sensitively. The pathogenesis of OA 1 is unknown. The presence of skin macromelanosomes helps diagnose OA 1 and they are seen in most affected adult males, but there is uncertainty about their presence in infants. One Australian family showed two males with OA 1 and one female carrier, none of whom had macromelanosomes, raising the possibility of different types of OA 1.

Macromelanosomes are formed putatively when premelanosomes do not separate from the endoplasmic reticulum-Golgi system. The accumulating proteins cause distension of the organelle. The OA 1 gene product could be a membrane protein necessary for the maturation of melanosomes. Skin biopsy for macromelanosomes also aids in determining the carrier state. Mutation in the OA 1 gene in Xp22.3 accounts for a third of the X linked albinism, suggesting that different gene defects on the X chromosome could cause the same phenotype—that is, locus heterogeneity could exist.

AUTOSOMAL RECESSIVE OCULAR ALBINISM?

Patients diagnosed with autosomal recessive ocular albinism (AROA) have appeared similar to mild cases of OCA 1 or OCA 2. However, genetic analysis has shown that some have abnormalities of the tyrosinase gene and so are OCA 1, while others have abnormalities of the P gene, thus categorising them as OCA 2. In fact, AROA may not be a distinct entity: 14% of those previously diagnosed with AROA have an abnormality of the tyrosinase gene, making them OCA 1 (chromosome 11); 36% have an abnormality of the P gene, relabelling them OCA 2 (chromosome 15);
disposition, bursts of laughter, hyperactive behaviour, microcephaly, brachycephaly, tongue protrusion, widely spaced teeth, wide based gait, and stiff movements. PWS is also a multisystem disorder, characterised by obesity, hypotonia, hypogonadism, intellectual impairment, short stature, and dysmorphic facial features.

Both AS and PWS are caused by the same, cytogenetically indistinguishable, chromosomal deletion. Yet, two separate phenotypes occur because of the phenomenon of genomic imprinting. If the microdeletion of 15q11-13 occurs on the paternally derived chromosome, then PWS occurs. Maternally derived chromosomal deletions of 15q11-13 cause AS. A difference in the imprint of the chromosome occurs, depending on whether it is paternal or maternal in origin, even though both chromosomes have the same DNA sequence. The cause of imprinting is unknown. One theory holds that genes from the mother may differ in their level of methylation from paternally inherited genes. This may influence the level of expression of the genes. Highly methylated genes seem to be less likely to be transcribed into messenger RNA.

**CONDITIONS LINKED TO OCA2**

Many patients with contiguous gene syndromes involving the Xp22.3 region have been described and have albino phenotypes if the deleted region includes the OA1 gene. X linked ichthyosis, Kallmann syndrome (hypogonadotropic hypogonadism with anosmia due to olfactory lobe agenesis) and X linked recessive chondrodysplasia punctata have been linked to the OA1 gene.

A patient has been described with impaired visual acuity, nystagmus, hypopigmentation of the retina, iris translucency, ichthyosis, hypogonadism, anosmia, and neurological abnormalities. A contiguous gene syndrome, which includes three X linked recessive disorders—that is, X linked ocular albinism, X linked ichthyosis, and Kallmann syndrome, is proposed.

X linked ocular albinism and late onset sensorineural deafness (OASD) causes visual impairment and deafness by middle age. OA1 and OASD are caused either by allelic variants or by contiguous gene defects. OASD differs from X linked albinism deafness syndrome by the presence of patchy, cutaneous hypopigmentation and hyperpigmentation and the absence of significant eye involvement in the latter disorder. These two conditions may be pleiotropic manifestations of the OA1 gene or a result of a contiguous gene defects.

The loci for OA1 and “microphthalmia with linear skin defects” (MLS) are closely situated on the X chromosome. MLS is characterised by microphthalmia, corneal opacities, and patches of erythematous skin in the head and neck in female patients.

A review of 16 patients with a deletion of the terminal end of the Xp arm, including Xp22 (Xpter-Xp22), was undertaken by Wapenaar et al. Two male patients had OA1 based on the presence of nystagmus, iris translucency, pale fundi, and poor visual acuity. One female patient has areas of hypopigmentation and hypopigmentation in her fundus, indicating that she might be a carrier of OA1. MLS features were found in eight of nine female patients. When other features in the female patients were looked for, chorioretinal abnormalities, agenesis of the corpus callosum, costovertebral abnormalities, mental retardation, and seizures were found. These findings overlap with two other conditions—Aicardi and Goltz syndromes. Both of the syndromes are X linked dominant and lethal in males. Aicardi syndrome consists of retinal lacunae, absence of the corpus callosum, and seizures. Goltz syndrome is characterised by microphthalmia, focal dermal hypoplasia, and skeletal abnormalities. It is hypothesised that these condi-

**Conditions associated with albinism: close linkages**

**CONCLUSIONS LINKED TO OCA1**

Prader–Willi syndrome (PWS) and Angelman syndrome (AS) are both due to a microdeletion on the human chromosome 15q11-13. The P gene defective in OCA2 is also coded in the region q11-13 on chromosome 15, adjacent to the area commonly deleted in PWS or AS syndrome. One per cent of patients with PWS or AS have OCA2. Therefore, if a patient has a deletion of one allele, as in Prader–Willi syndrome, and the fellow allele has a mutation for OCA2, the patients will be albinos. Angelman syndrome is characterised by mental retardation and absence of speech. Affected patients have a happy disposition, bursts of laughter, hyperactive behaviour, microcephaly, brachycephaly, tongue protrusion, widely spaced teeth, wide based gait, and stiff movements. AS is also a multisystem disorder, characterised by obesity, hypotonia, hypogonadism, intellectual impairment, short stature, and dysmorphic facial features.

Both AS and PWS are caused by the same, cytogenetically indistinguishable, chromosomal deletion. Yet, two separate phenotypes occur because of the phenomenon of genomic imprinting. If the microdeletion of 15q11-13 occurs on the paternally derived chromosome, then PWS occurs. Maternally derived chromosomal deletions of 15q11-13 cause AS. A difference in the imprint of the chromosome occurs, depending on whether it is paternal or maternal in origin, even though both chromosomes have the same DNA sequence. The cause of imprinting is unknown. One theory holds that genes from the mother may differ in their level of methylation from paternally inherited genes. This may influence the level of expression of the genes. Highly methylated genes seem to be less likely to be transcribed into messenger RNA.
Conditions associated with albinism and not due to close linkage

CHEDIAK–HIGASHI SYNDROME

Chediak–Higashi syndrome (CHS) is an autosomal recessive disease characterised by oculocutaneous albinism and severe immunological deficiency. An increased susceptibility to infections and deficient natural killer cell activity occurs. Giant intracytoplasmic inclusion bodies are seen in most granulated cells, granulocytes, lymphocytes, mast cells, histiocytes, platelets, melanocytes, Schwann cells, neurons, fibroblasts, and renal tubular epithelium. CHS has recently been mapped to chromosome 14q21–q24. and the gene has been cloned and sequenced, but the CHS gene product is unknown. Patients have defective vesicular transport to and from the lysosome and late endosome. Without a bone marrow transplant, CHS patients die during childhood from organ infiltration or haemorrhage. The oldest documented patient with CHS is aged 39 years. Seizures, mental retardation, cranial nerve palsies, clumsiness, gait abnormalities, and peripheral neuropathy develop with time. 42

Thus, CHS patients have hypopigmentation due to abnormal melanosomes; the hypopigmentation is caused by a different mechanism from other albinoid disorders. The presence of giant melanosomes indicates decreased pigmentation, producing hypopigmentation of the hair, skin, and ocular fundus. The hair in CHS appears a metallic frosted grey colour (Fig 4). 43

Cats with CHS have misrouting of the optic nerve fibres at the chiasma, disorganisation of the dorsal lateral geniculate nuclei, and an abnormality of the superior olivary complex. CHS cats, other albino cats, and human albinos have anomalies in the brain stem auditory structures involved in transmitting information to the contralateral portion of the brain stem. 44–46 The fact that CHS cats have neurological abnormalities similar to other albino subjects suggests that the abnormal pigmentation itself may be the cause.

HERMANSKY–PUDLAK SYNDROME

The Hermansky–Pudlak syndrome (HPS) is an autosomal recessive inherited disease characterised by albinism, a bleeding diathesis due to storage pool deficiency of platelets, and a lysosomal ceroid storage disorder. Three organelles are affected—melanosomes, platelet dense bodies, and lysosomes. Late complications of HPS include interstitial pulmonary fibrosis, inflammatory bowel disease, renal failure, and cardiomyopathy due to ceroid-like material deposition. Pulmonary fibrosis is the most common cause of death, but haemorrhage and granulomatous colitis can be fatal. 47–50 The carrier frequency in Puerto Rico is estimated at 1:21. HPS represents the most frequent type of albinism in the Puerto Rican population, with a frequency of 1:1800. Five out of six albinos in Puerto Rico have HPS. It is thought that a founder effect may be operating. The original mutation may have occurred after attacks by mutant British soldiers in 1598, Dutch attackers in 1625, the presence of aboriginal tribes, or slave traders. HPS is also over-represented in an isolated Swiss village. 51

The HPS gene is localised to 10q23.3–q23.4 and has recently been cloned. 52 Cutaneous pigmentation varies from none to almost normal, with the majority developing ocular signs of nystagmus, strabismus, reduced acuity, foveal hypoplasia, and decreased retinal pigmentation. 53 The melanosomes are decreased in number, have a reduced amount of melanin, and have an abnormal morphology. Absence of dense bodies in platelets (these are storage granules for adenine nucleotides and other compounds for secondary aggregation) predispose to bleeding and bruising. It is not known how all the components of HPS are related.

Congenital hypopigmentation resembling albinism

WAARDENBURG SYNDROME

Waardenburg syndrome is characterised by heterochromic irides and congenital hypopigmentation of the remainder of the uveal tract. Other findings include lateral displacement of the medial canthi, dystopia of the lacrimal puncta, blepharophimosis, synophrys, white forehead, pale skin, premature greying, defective pigmentation, congenital deafness, and a broad root of the nose. There are various genetic forms of Waardenburg syndrome that express subtle phenotype distinctions. Waardenburg syndrome types one and three result from mutations in the PAX 3 gene encoding a transcription factor and localised to chromosome 2q35–q37. 53–55 Waardenburg syndrome type 2 results from mutations in the MITF (microphthalmia transcription factor) located at 3p12. 56–58 A fourth type, Waardenburg–Hirschsprung disease, results from mutations in the ENDGB gene at 13q22 that encodes a G protein–coupled plasma membrane receptor that can bind endothelin 1, 2, and 3. Patients present primarily with congenital aganglionic megacolon, resulting from the absence of neural crest derived Meissner’s and Auerbach’s autonomicplexuses in the intestinal smooth muscle wall. Hirschsprung syndrome is also associated with hypopigmentation of varying extent, heterochromic irides, and deafness. The hypopigmentation is thought to result from the inability of a subset of neural crest cells—that is, melanoblasts and enteric ganglia, to reach their target site during embryogenesis. 59–62

Conditions of possible relevance

CONGENITAL MOTOR NYSTAGMUS

Congenital motor nystagmus (CMN) is diagnosed by the presence of rhythmic horizontal nystagmus. Most children diagnosed with CMN have X-linked albinism. Kriss et al 61 found that some males diagnosed with CMN, in fact, have XLOA.

Mechanisms of impaired vision in albinism

Mechanisms of visual impairment in albinism are multifactorial and undoubtedly related to pigmentation. In foveal hypoplasia, as occurs in albinism, the central cones are spaced apart and so the central visual acuity is decreased while the peripheral visual acuity is normal. Refractive errors are more common, especially astigmatism. Some degree of amblyopia may, thus, be
present. It is noted that the refractive error in many infants is due to poor visual acuity rather than its course. The albino has a high risk of strabismus. Stereopsis is absent as a result of misrouting of the optic pathways. Intraocular light scatter will also degrade the retinal image.

**Neurological ramifications of disordered pigmentation**

Melanin plays an important role in the developing visual system; a critical level of pigment production is required for the normal development of visual pathways. Lower levels lead to a stereotypic set of defects of neuronal migration, vision impairment, nystagmus, and strabismus.

An increased number of decussating axons at the chiasm is a regular feature of all types of albinism. At the cellular level, decussation is caused by a complex interaction between ganglion cell and optic chiasm embryonic neurons. Embryonic neurons of the optic chiasm apparently control the orientation and decussation of ganglion cells via the expression of surface cell molecules, L1 and CD44. When monoclonal antibody and complement are used to ablate embryonic chiasmal neurons, the chiasm does not form and the retinal ganglion cells are truncated at the position where they would be expected to enter the chiasm.

The level of pigmentation and the amount of tyrosinase activity is related to the amount of ipsilateral innervation. In some cases, the tyrosinase gene may play a role in controlling ganglion cell growth. An elegant study performed by Jeffer ey et al found that introduction of a tyrosinase gene in transgenic mice corrected the abnormal pathway that occurred in the albino mice strain investigated. The amount of tyrosinase activity appears closely related to the number of uncrossed optic chiasm axons, adding further evidence to the theory that some factor in the retinal axon development plays a pivotal role in the direction of ganglionic axons in the regions of optic chiasmata.

The manner in which albinism affects ganglionic axon guidance through the optic chiasm is controlled at the level of the retina and not at the level of the chiasm. Albinism changes, at the chiasm, the number of ganglion cells specified to respond to cues for decussating. Albinism does not alter signals from the chiasm to the ganglion axons.

**Steps in assessing an albino suspect**

In order to define more precisely the type of albinism a patient has, the following steps can be undertaken.

1. **In assessing the phenotype of the patient, if the newborn, and especially the adult, completely lacks pigment in the skin and hair, the individual probably has OCA 1A. If minimal pigment is apparent, the individual could be classified as either OCA 1B, OCA 2, or OCA 3. If the individual has silvery coloured hair and neutrophils with large inclusions, as observed by light microscopy of a blood smear, the diagnosis would be Chediak-Higashi syndrome. If the individual has moderate to minimal hypopigmentation and exhibits reduced blood clotting upon testing, the diagnosis would be Hermansky-Pudlak syndrome.**

2. **A hair bulb assay could be performed to identify OCA 1A more definitively. In this test, hair bulbs are plucked from the scalp and the catalytic activity of tyrosinase is determined either (a) by a histochemical procedure in which the samples are incubated in the substrate DOPA and the consequent induction of melanin determined by visual inspection, or (b) by a radioactive biochemical assay in which the samples are incubated with a radiolabelled tyrosine precursor and the amount of radiolabel released after enzymatic conversion quantified spectrophotometrically. A negative result with either assay indicates OCA 1A, a positive result indicates OCA 1B, OCA 2, or OCA 3. OCA 1 will also be positive.**

3. **To differentiate between OCA 1B, OCA 2, OCA 3, sequence analysis of the genes encoding tyrosinase, P protein, and TRP-1, respectively, must be performed. This is not a routine laboratory procedure and currently can be done in only a few research laboratories. Alternatively, a 5–8 mm shave skin biopsy can be obtained from the patient. Cultures of melanocytes can be established in order to assess the expression and/or function of tyrosinase, P protein, and TRP-1. Again, this is not a routine laboratory procedure and currently can be done in only a few research laboratories.**

4. **If X linked ocular albinism (OA 1) is suspected after examining the ocular phenotype, female family members need to be assessed for the presence of a mud splattered fundus, indicating carrier status. A skin biopsy can be sent to a histology laboratory to look for macromelanosomes in the epidermis. A few laboratories will attempt to sequence the gene, but that mutation has not been found to account for all patients.**

We thank Dr J Nordlund, Department of Dermatology, University of Cincinnati, Cincinnati, Ohio, USA.

SUSAN M CARDEN

Royal Children’s Hospital, Melbourne, Australia and Smith-Kettlewell Eye Research Institute, San Francisco, California, USA.

RAYMOND E BOISSY

Department of Dermatology, University of Cincinnati Medical Center, Cincinnati, Ohio, USA

PAMELA J SCHEETTKER

Department of Ophthalmology, University of Cincinnati Medical Center, Cincinnati, Ohio, USA

WILLIAM V GOOD

Smith-Kettlewell Eye Research Institute, 2232 Webster Street, San Francisco, CA 94115, USA

Correspondence to: William V Good, MD.
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