Hermansky-Pudlak syndrome (HPS) is a rare autosomal recessive disease, characterized by a triad of oculocutaneous albinism (OCA), bleeding diathesis due to deficiency of dense bodies in platelets, and lysosomal accumulation of ceroid lipofuscin. Seven genetic subtypes of HPS have been identified in humans; HPS1 on chromosome 10q23 is the most common and represents a founder effect in northwest Puerto Rico. The clinical features of HPS and its ophthalmic involvement are well documented, but no ocular histopathology has been published (to our knowledge). Herein, we describe the ocular histopathology of an adult patient with HPS type 1 (HPS-1).

**Ocular Pathologic Features of Hermansky-Pudlak Syndrome Type 1 in an Adult**

Hermansky-Pudlak syndrome (HPS) is a rare autosomal recessive disease, characterized by triad of oculocutaneous albinism (OCA), bleeding diathesis due to deficiency of dense bodies in platelets, and lysosomal accumulation of ceroid lipofuscin. Seven genetic subtypes of HPS have been identified in humans; HPS1 on chromosome 10q23 is the most common and represents a founder effect in northwest Puerto Rico. The clinical features of HPS and its ophthalmic involvement are well documented, but no ocular histopathology has been published (to our knowledge). Herein, we describe the ocular histopathology of an adult patient with HPS type 1 (HPS-1).

**Report of a Case.** This study was approved by the National Human Genome Research Institute and National Eye Institute institutional review boards for human subjects, and informed consent was obtained from the patient. A 43-year-old Puerto Rican man was seen in March 2004. The diagnosis of HPS-1 was confirmed by demonstrating homozgyosity for the 16–base pair duplication in exon 15 of HPS1. Systemic manifestations included OCA, colitis, nasal and gum bleeding, basal cell carcinoma, and severe pulmonary fibrosis.

Findings from an ophthalmic examination revealed horizontal infantile-onset jerk nystagmus with a torsional component and intermittent exotropia. Best-corrected visual acuity was 20/160 OU (Early Treatment of Diabetic Retinopathy Study), and hyperopic astigmatism was present in both eyes. Posterior embroyotoxon, marked iris transillumination, and macular transparancy were noted (Figure 1). The absence of foveal pits and light reflexes indicated foveal hypoplasia, which was confirmed with optical coherence tomography (Figure 2). The patient died of pulmonary fibrosis in April 2005.

**Pathologic Findings.** Macroscopically, the right globe measured 24 × 25 × 22 mm and the left globe measured 23 × 24 × 23 mm. The cornea, anterior chambers, and optic nerves were normal. The uvea and retinal pigment epithelium displayed marked hypopigmentation. No macula lutea was visible. A focal hemorrhage was noted in the conjunctiva of the right eye.

Microscopically, hemorrhage was present in the temporal conjunctiva of the right eye. A small cluster of mesenchymal cells was adherent to the enlarged and anteriorly located Schwalbe line (posterior embryotoxon) bilaterally (Figure 3A and B). The scleral spur of the right eye was hypoplastic. Only small aggregates of large melanin granules were visible in the pupillary margin (Figure 3C and D). Moderate hyalination of the ciliary body was observed in both eyes (Figure 3E and F). There was marked depigmentation in the entire uvea (Figure 4). A few fine melanin granules remained in ocular pigment epithelial layers. The fovea showed a lack of differentiation, and the retinal pigment epithelium contained sparse melanin granules (Figure 4B). A few hemorrhages and platelet aggregates were seen in the optic nerve head of the right eye.

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**Figure 1.** Patient with Hermansky-Pudlak syndrome type 1 described herein. A, The right iris is light brown with marked transillumination such that the edge of the crystalline lens is clearly visible. B, The fundus of the left eye shows clear choroidal vasculature with residual pigmentation in the posterior pole.
Ultrastructurally, the uveal tissues contained a few melanocytes with sparse immature melanosomes of incomplete round and oblong-shaped melanized granules (Figure 5A and B). Some melanocytes had electron-dense halos and electron-lucent cores. Most melanocytes were swollen and devoid of melanosomes. The nuclei had condensed chromatin, and some had a folded nuclear membrane. The iris pigmented epithelial cells were enlarged, with few cytoplasmic microorganelles (mitochondria, ribosomes, endoplasmic reticulum, and Golgi complex) and thickened basement membranes (Figure 5C and D). The morphologic structure of the choroid was similar to that of the iris (Figure 5E). Only rarely did choroidal melanocytes contain stage IV melanosomes (Figure 5F).

Comment. Hermansky-Pudlak syndrome represents a disorder of the formation, sorting, or trafficking of intracellular vesicles, including melanosomes in melanocytes and dense bodies in platelets. As a consequence, patients with HPS-1 exhibit albinism, bleeding, and (for unknown reasons) pulmonary fibrosis and ceroid lipofuscinosis. The eyes of our patient had no ceroid accumulation, but their choroidal melanocytes contained numerous aberrant membranous structures, both U- and ring-shaped (Figure 5), as previously reported in HPS skin melanocytes. This is consistent with dysregulated targeting or fusion of Golgi-derived vesicles and with impaired targeting of melanocyte-specific proteins (eg, tyrosinase-related protein 1 and granulophysin) to premelanosomes.

Other major ocular pathologic findings consisted of posterior embryotoxon with a small cluster of adherent mesenchymal cells, moderate hyalinization of the ciliary body, foveal hypoplasia, multiple ocular hemorrhages, and marked hypopigmentation. The iris pigment epithelium displayed the most severe ultrastructural pigmentary defect. Uveal melanocytes of the ocular tissue contained mainly stage III melanosomes; the existence of stage IV melanosomes in rare choroidal melanocytes indicates that this patient with HPS-1 was still able to produce morphologically mature melanosomes, albeit in an extremely reduced quantity.
Hyalinization of the ciliary body occurs in aging ciliary processes, longstanding glaucoma, or heterochromic iridocyclitis, implying an atrophic tendency. We propose that the hemorrhagic diathesis may have contributed to this complication. Axenfeld anomaly and foveal hypoplasia, also present in our patient, are common in OCA.\(^8\) Posterior embryotoxon was clinically apparent in 5 of 20 previously described patients with HPS, with only 1 manifesting iris process adhesion.\(^3\) In our patient, the mesenchymal cells, not the iris processes, were adherent to posterior embryotoxon. Such a subtle abnormality is difficult to detect clinically, especially in the presence of the underlying nystagmus of HPS.

Our patient’s ocular melanocytes contained mainly stage III and rarely stage IV melanosomes, consistent with previous reports of OCA.\(^9\) This finding differs, however, from descriptions of HPS skin melanocytes, which contain exclusively premature melanosomes, mostly stage I and II.\(^6,9\) Natsuga et al\(^7\) identified stage IV melanosomes in the melanocytic nevus of a patient with HPS-1 and hypothesized that this reflected the high melanin production of the nevus cells. The replacement of melanosomes with multiple vacuoles and a marked decrease in microgranules of the iris pigmented epithelia have not been previously reported in OCA, to our knowledge. Although the occurrence of a giant melanosome in OCA and HPS is not unusual,\(^7,9\) no ultrastructural giant melanosomes were apparent in this case. Instead, large melanin granules were present in the pupillary margin microscopically, which could be pigmented macrophages.

The mouse pale ear (ep) mutation is the homologue of human HPS1.\(^10\) These mice exhibit abnormalities in melanosomes and platelet-dense granules that are similar to those in patients with HPS. Similar to our patient, ep mice initially have markedly decreased and unevenly pigmented melanosomes, lightly pigmented intermediate structures, and stage IV melanosomes. However, ep mice display macromelanosomes, which were absent in our patient.\(^10\)

This case offers the first example (to our knowledge) of ocular histopathologic features in HPS-1. Similar descriptions are needed for the other HPS subtypes.

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Outbreak of Fusarium Keratitis in Soft Contact Lens Wearers in San Francisco

We report a cluster of 4 cases of soft contact lens–associated *Fusarium* keratitis seen at the University of California, San Francisco (UCSF), during a 5-week span in early 2006 and compare this cluster with the number of previous cases of culture-positive *Fusarium* keratitis seen at UCSF during the prior 30 years. This cluster represents part of a larger outbreak of *Fusarium* keratitis currently under investigation by public health authorities in Singapore and the United States. As in these outbreaks under investigation, soft contact lens wear and use of ReNu with MoistureLoc or ReNu MultiPlus (Bausch & Lomb, Roch-

Figure 5. Transmission electron micrographs. A, A few melanocytes that contained sparse immature stage III melanosomes (0.2-0.6 µm in diameter) with incomplete round and oblong melanized granules (arrows). B, Rare membrane-limited lipofuscinlike material with fat globules (open arrow) is noted. C and D, The iris pigmented epithelial cells are enlarged, contain a decreased number of cytoplasmic microorganelles, and have thickened basement membranes. E and F, In the choroids, many melanocytes were swollen and degenerated. Some melanocytes contained sparse, small, immature melanosomes, and some contained an aggregation of stage IV melanosomes (arrow). U-shaped structures were occasionally seen (open arrow) (scale bar, 2 µm).