Hypomelanosis of Ito: A Rare Disorder with Rarer Presentation-Unilateral Retinal Detachment and Bilateral Glaucoma

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Abstract
Hypomelanosis of Ito, also known as Hypomelanosis of Achromia, Incontinentia pigmenti Achromians, pigmentary dysplasia or mosaicism, is a rare dermatological disease with a prevalence of 1 in 8000-10,000 live births predominantly affecting females. Disease is usually sporadic but familial cases have been reported. It is characterized by streaked/whorled patches of light colored skin. It may have neurological abnormalities (seizures being the most common), dental abnormalities (defective teeth implantation, cortical teeth, partial anodontia/hypodontia) and ophthalmic associations like strabismus [1-4]. We hereby report a case of a 13 years old female having Hypomelanosis of Ito with bilateral cataract and angle closure glaucoma complicated with unilateral retinal detachment which are rare associations. She underwent a cataract surgery as well as implantation of Glaucoma Drainage Device (GDD) in the right eye after a failed trabeculectomy.

Keywords: Angle closure glaucoma; Cataract; Hypomelanosis of Ito, Incontinentia pigmenti achromians; Retinal detachment

Case Report
A 13 years old female presented to our hospital in 2015 with complaints of sudden painful diminution of vision in left eye since the past one month. She had a history of recurrent episodes of left sided headache with nausea—about 5–6 such episodes per month over the past 1 year. These episodes were usually associated with diminution of vision in left eye and were relieved by taking analgesics or sleeping. Systemic, birth, family or personal history was non contributory. There was no history of any prolonged intake of steroids in any form (oral/topical/inhalational).

Upon clinical evaluation, hypopigmented streaks were present on both upper extremities (Figure 1a) along with scattered patches of alopecia over scalp (Figure 1b). She also had microdontic, abnormally shaped teeth along with polydactyly and syndactyly in left foot (Figure 1c).

Figure 1: Clinical photograph of the patient showing a) Hypopigmented patches/streaks on the skin of the arm, b) Alopecia patches on the scalp, c) Polydactyly and syndactyly of 2nd and 3rd toes as well as 5th and 6th toes of the left foot.

On ophthalmic evaluation, Best Corrected Visual Acuity (BCVA) was 6/24 in right eye and perception of light with inaccurate projection of rays in left eye. Horizontal nystagmus was present in both eyes. Iris was normal in color and pattern in right eye. However, left eye had Neovascularisation of Iris (NVI) with partial loss of pupillary ruff (Figure 2). Relative Afferent Pupillary Defect (RAPD) was noted in left eye. Cortical cataract as well as posterior subcapsular
cataract with diffuse pigment deposition on anterior lens capsule was observed in both eyes (Figure 2). Intraocular Pressures (IOP) as measured by Goldmann Applanation Tonometry were 43 and 6 mm Hg in right eye and left eye respectively. Central corneal thickness by ultrasonic pachymetry was 635 and 645 µm in right eye and left eye respectively. On gonioscopy, occludable angles were seen in Right Eye with steep iris configuration (Figure 3a) and angles were closed in left eye with 360 degrees Peripheral Anterior Synchiae (PAS) (Figure 3b). Fundus of RE showed tilted disc with cup to disc ratio of 0.6-0.7:1 and peripapillary hypopigmented streaks (Figure 4a). Total retinal detachment was seen in left eye which was also confirmed by Ultrasound B scan (Figure 4b).

On the basis of above typical dermatologic and dental findings, she was diagnosed as a case of Hypomelanosis of Ito by dermatologist and paediatrician. Vitrectinal surgery in left eye was avoided in view of poor visual prognosis. The patient underwent right eye phacoemulsification with intraocular lens implantation under general anesthesia. Cataract surgery was done for quick visual rehabilitation and combined surgery was avoided in view of patient being one eyed. Since, IOP was uncontrolled in right eye despite maximum medical therapy; the patient underwent trabeculectomy with 0.2% Mitomycin C under general anesthesia 3 months post her cataract surgery. The postoperative period was uneventful although IOP was still uncontrolled with a failed trabeculectomy, so the patient was advised GDD (Ahmed Glaucoma Valve-FP7 variant) implantation along with a scleral patch graft which she underwent 5 months post her trabeculectomy. Postoperatively, at the time of her 4 years follow up, her IOP was well controlled (14mm of Hg on 2 antiglaucoma medications, with well formed bleb and a patent GDD tube in anterior chamber) and BCVA of 6/9 in right eye.

Appropriate genetic counseling along with evaluation of other siblings was undertaken although all the other siblings and parents were asymptomatic. Patient was advised to be on regular follow up every 6 monthly.

Discussion

Hypomelanosis of Ito is a rare dermatological disease with a reported prevalence of 1 in 8000-10,000 and it predominantly affects females. Disease is usually sporadic but familial cases have also been reported with inheritance patterns being autosomal dominant, autosomal recessive as well as X linked. Its cause is unknown and various chromosomal abnormalities have been identified on chromosomes- 9q33, 15q11-q13, Xp11, X p21.2 [1,2]. However, in many patients the condition arises from genetic irregularities that are present in some cells of the body, but not in others (mosaicism). Some researchers believe that it does not represent a distinct disorder but rather a symptom, common to a group of disorders involving genetic mosaicism [5]. It is characterized by dermatologic manifestations include changes in hair color, diffuse alopecia on scalp, trichorrhexis nodosa, polydactyly, coarse facial features. Hypomelanosis may present as hypopigmented patches, streaks or whorled areas in any part of the body. The skin lesions usually appear during the first year of life and remain unchanged through childhood, but may fade or darken in adulthood. Skin lesions are not associated with inflammation or a premalignancy. Neurological findings such as infantile seizures resistant to therapy, cognitive impairment, developmental delays and musculoskeletal symptoms such as scoliosis are commonly associated with this condition. Dental manifestations include defective implantation, cortical teeth, partial anodontia, hypodontia [3-6]. The symptoms usually become apparent during the first or second year of life. Ophthalmic associations can be variable like strabismus, aplasia/hypoplasia of iris, heterochromia iridis, corneal opacity, hypertelorism, myopia, uveal coloboma or microphthalmos. Rare ophthalmic associations include nystagmus and cataract [3,4]. Presence of bilateral cataract in this condition is very rare as was seen in our case. Furthermore, we did not find any reported association of glaucoma with this condition. Thus ours is a rare case of Hypomelanosis of Ito which was diagnosed on the basis of characteristic clinical features, with bilateral angle closure glaucoma and cataract and unilateral retinal detachment. However, we did share same findings like cataract and retinal detachment with a previous...
reported case [7] although this patient had cataract in one eye and retinal detachment in the contralateral eye. Another case report of a 4 month female baby with incontinenti pigmenti had ophthalmic features in the form of bilateral retinal detachment is also documented [8].

We believe that the chronically raised IOP (>40mm of Hg) in the left eye led to traction on the retina leading to retinal detachment and also the chronic ischaemia due to raised IOP lead to NVI. There was no similar history in family or in other siblings thus highlighting that there was a sporadic mutation affecting our patient.

This case highlights the importance of a good general physical examination, especially in cases with atypical findings. Patients with Hypomelanosis of Ito should be investigated for associated sight-threatening ocular pathologies to prevent visual morbidity.

**Conclusion**

This case highlights the importance of a good general physical examination, especially in cases with atypical findings. Appropriate referrals to paediatrician/dermatologist should be done when faced with atypical features in clinical practice. Patients with Hypomelanosis of Ito should be investigated for associated sight-threatening ocular pathologies. It also highlights the role of an ophthalmologist among these patients because timely and appropriate management can significantly reduce visual morbidity. An inquisitive mind with detailed systemic evaluation is required on part of ophthalmologist to reach a correct diagnosis in cases of unusual ophthalmic findings.

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