Congenital Icthyosis: A case report

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Abstract
Congenital Ichyosis is relatively uncommon disease but is the most common cause of Collodian baby. Early diagnosis is required for immediate management and prevention of fluid-electrolyte balance as well as prevention of secondary infections. Definitive diagnosis is essential for long-term management as well as counselling to the parents of affected infant if they are considering having children together.

Case report
A full term male infant was born by spontaneous vaginal delivery to a 35 y/o, G5P4 woman. Her prenatal course, including prenatal ultrasound, was unremarkable.

Examination at birth: He had normal anthropometry with height, weight and head circumference being between 25th to 75th percentiles. Skin: Diffuse erythematous skin lesion with collodion membrane covering the chest and back, tight skin on upper and lower extremities, fissures on areas with skin folds, eversion of the eyelids with inability for complete eye examination, eversion of the lips, and malformation of the auricle.

Tone and turgor: within normal limit.

Heent: Anterior fontanelle: soft and flat, normal shape and size.

Eyes: Eversion of the eyelids.

Ears: External malformation, unable to assess if secondary to skin lesions.

Nose: Nares and septum appear within normal limit.

His cardiovascular, abdominal, spine, back, extremities and external genitile examinations were normal.

On further evaluation, family history was not significant for any medical and dermatological conditions. There was no history of consanguinity. The patient has a 2 y/o healthy full sister. Mother was carrier for Cystic Fibrosis. He has two half-brothers (13 y/o and 10 y/o) from maternal side who has eczema, otherwise healthy; two half-sisters and one brother from paternal side- all reportedly healthy (Figure 1 and 2).

Discussion
Collodion baby is a severe form of Congenital Ichthyosis. Congenital Ichthyosis is heterogeneous group of disorders of cornification involving all or most of integumentary system [1] which is characterized by a generalized scaling of the skin of varying severity. Mucosal surface and visceral organs are generally spared. Cornification is process of terminal keratinocyte differentiation. At this stage, a lipid envelope is extruded into the intercellular space from lamellar bodies to form lipid sheets composed of ceramides, cholesterol, and free fatty acids [2].

Normal skin turnover
Skin has dermis and epidermis layers. Epidermis has four layers-stratum basale, spinosum, granulosum, and corneum from base to top. During normal skin turnover of 28 days, site-relevant proteins and lipids are synthesized in different layers of epidermis.

Revised nomenclature and classification of inherited ichthyoses
The First Ichthyosis Consensus Conference in Sorèze 2009 proposed the following pathogenesis of inherited Ichthyosis [1] which is based on disorders involved in different stages of protein and lipid metabolism and on other mechanism.

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lifelong complication of various degrees to most affected individuals widely with complete resolution within few weeks in ten percent, of keratinized epithelium [3]. Clinical course of collodion baby varies membrane is transient, transparent membrane present at birth, made of “acquired Icthyosis”, primarily seen adults as a manifestation of diseases. (D) Disorders of DNA transcription/ repair.

(A) Disorders of keratinocyte protein (“bricks”) metabolism–cytoskeleton, cornified lipid- cell envelope, protease/protease inhibitors, that lead to weakening of cytoskeleton and decreased mechanical stability/cytotoxic effects, premature loss and defective processing).

(B) Disorders of lipid (“mortar”) metabolism- defective assembly or transport of lipids including steroid sulfatase deficiency, lipid metabolism defects including liposomal and neutral lipid storage diseases.

(C) Disorders of tight and gap junctions and

(D) Disorders of DNA transcription/ repair.

Same consensus nomenclature classified inherited Icthyosis as following (Table 1):

Table 1. Clinico-genetic classification of inherited Icthyosis. Abbreviations: ARCI, autosomal recessive congenital ichthyosis; CNS, central nervous system; KI, keratinopathic ichthyoses; IV Icthyosis Vulgaris; RXLI Recessive X Linked Icthyosis; HI Harlequin Icthyosis; LI Lamellar Icthyosis; CIE: congenital ichthyosiform erythroderma; EI epidermolytic ichthyosis; SEI superficial epidermolytic ichthyosis [1].

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<tr>
<th>Non-syndromic ichthyoses</th>
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<td>1. Common ichthyoses: IV, RXLI</td>
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<td>2. ARCI: HI, LL, CIE</td>
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<td>3. KE- EI, SEI</td>
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<td>4. Other forms of non-syndromic ichthyosis</td>
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<th>Syndromic ichthyoses</th>
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<td>5. X-linked ichthyosis syndromes</td>
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<td>6. Other ichthyosis syndromes (CNS signs, fatal disease course, hair abnormalities and/or other associated signs)</td>
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We sent genetic testing for TMG1 gene. There was significant for first five days of life, as it was almost impossible to get peripheral IV access. We kept newborn in tertiary level NICU. He was in Isolette with high humidity and milk/ intravenous fluid intake is adjusted to avoid dehydration. CB also have impaired sweating mechanism so clinical must be vigilant to avoid and aggressively treat hyperthermia and hypohydrosis. Daily weight and electrolytes help in maintaining fluid and electrolyte balance.

Routine use of emollients is controversial as it helps decreasing insensible water loss but increases the risk of infection [11,12]. Defecting skin barrier promotes bacterial and fungal infection. Deep skin crease may delay early diagnosis. Daily skin examination including skin fissures is recommended. Recommended hand wash cannot be over emphasized. Early diagnosis and aggressive treatment of both local and systemic infection is required for better outcome. Eye care should be coordinated with ophthalmologist and bland lubricant is recommended to prevent development of keratitis and conjunctivitis [6,13]. For severe scaling, oral retinoids, such as acitretin and isotretinoin, are an important therapeutic option [14].

Our case

We kept newborn in tertiary level NICU. He was in Isolette with humidity of 70-80 %. He was given feeds in demand and IV fluid for first 2 days. Intravenous Ampicillin and gentamycin was given for 5 days. We monitored daily weight and electrolytes, strict input and output. There was no significant weight change and no electrolyte balance. There was no significant weight change and no electrolyte abnormalities. We used topical Vaseline twice daily and looked for signs and symptoms of cutaneous and systemic infections daily. He did not have any infection. He got eye lubricants 3-4 times a day to avoid dry eye and its complications. Baby had umbilical venous line for first five days of life, as it was almost impossible to get peripheral IV access. We sent genetic testing for TMG1 gene. There was significant improvement in skin by early 2nd week of life- erythema decreased and collodion membrane shed. Skin was almost normal looking by end of 2
weeks. Ectropion of eyelid and eclabion improved as membrane shed. Genetic testing were negative. He got afebrile, superficial skin infection, treated with systemic antibiotics at 6 weeks of age. Child is doing well in 15 months of life. We discussed the inheritance of congenital Ichthyosis. The majority of cases of congenital Icthyosis are inherited in an autosomal recessive manner. This means that for both male and female offspring, the risk of recurrence is 25% with each pregnancy.

References