Lamellar Icthyosis: A Clinical Dilemma

Amar Verma¹, Rani Manisha², R. K. Narayan³

¹Professor, Department of Paediatrics & Neonatology, Rajendra Institute of Medical Sciences, ²Junior Resident (Academic), Rajendra Institute of Medical Sciences, ³Senior Resident, All India Institute of Medical Sciences.

Abstract

Congenital Autosomal recessive ichthyosis is a heterogeneous group of disorders that are present at birth with generalized involvement of skin and lack of other organ systems. This case report presents involvement of respiratory system and its management with outcome in a rather uncommon presentation of lamellar ichthyosis.

Keywords: Lamellar Icthyosis, Respiratory Distress, Ectropion, Epiphora.

Corresponding Author: Dr. Amar Verma, 6C Jyoti Vihar, Harmu Road, Ranchi, Jharkhand-834-001.

Received: August 2018
Accepted: September 2018

Introduction

Lamellar ichthyosis is a severe form of ichthyosis. Mostly it is autosomal recessive. Its incidence is less than 1 in 3 lakhs.[1] Other types of ichthyosis are Icthyosis vulgaris, Epidermolytic hyperkeratosis, Congenital echyosiform erythroderma, Sjogren Larsson syndrome, Rud’s syndrome, Refsum’s disease, Icthyosis linearis circumflexa etc. All types of ichthyosis presents from or shortly after birth. Diagnosis of lamellar ichthyosis is based on the history of colloidon membrane at birth which is shed soon after birth. The characteristic appearance of scales especially on shins.[2] Lamellar ichthyosis evolves into large, quadrilateral, dark scales that are free at the edges and adherent at the center. Scaling is often pronounced and involves the entire body surface, including flexural surfaces. The face is often markedly involved, including ectropion and small, crumpled ears. The palms and soles are generally hyperkeratotic. The hair may be sparse and fine, but the teeth and mucosal surfaces are normal.[3] The nails may be stippled, ridged and thickened with associated subungual hyperkeratosis. The condition is unremitting and persists throughout life.

Case Report

A 14 years old male patient reported in Department of Pediatrics, Rajendra Institute of Medical Sciences, Ranchi with complaint of difficulty in breathing since 7 days which was not associated with any history of fever or cough. Breathing difficulty was of sudden onset, slowly progressive, affected his daily activities, more in lying down position than sitting posture. There was no similar history of breathing difficulty in past. Detailed history revealed presence of dry scaly skin all over body since birth. The dryness improves in winter and worsens in summers. There was history of decreased sweating with mild pruritis. The patient was born out of non-consanguinous marriage. His growth and development milestones were normal. His parents and siblings are apparently healthy. History was suggestive of colloidon membrane present and premature rupture of membrane was present. Earlier patient was diagnosed in July, 2012 with lamellar ichthyosis at dermatology department, CMC Vellore.

Investigations done at CMC Vellore showed ---
Parathyroid hormone 249.9 pg/ml
Serum chloride 102 mmol/l, Serum sodium 134 mmol/l,
Serum potassium 4.3 mmol/l, Serum bicarbonate 21 mmol/l
Total bilirubin 0.5 mg%, direct bilirubin 0.2 %
Total Protein 7.4 g%, total albumin 4.2%
SGOT 17 U/L, SGPT 10 U/L, ALP 176 U/L
Serum Creatinine 0.64mg%
Total cholesterol 111 mg%, serum triglyceride 86 mg%,
HDL 33 %, LDL 60%
Hb 10.4 gm%, total WBC 12,600/mm³
N63%/L22%/M 10%/M 5%/B 0% with no nucleated RBCs
He did not report back to CMC Vellore for followup after
taking isotretiinoin for 15 days only.

Current patient reported at our Hospital with respiratory distress present even at rest (orthopnoea) without tachycardia and chest was clear. There was absence of organomegaly. CNS was within normal limits. Oxygen saturation was 96 percent at room air with no fever or lymphadenopathy. There was presence of dry skin with large scales all over body with minimal sparing of the perioral area.

The scales were dark brown in colour, polygonal in shape, large in size, thick and adherent to skin prominently over shins. Palmoplantar keratoderma was present. External
auditory meatus was hyperpigmentation and scaling was present. He was not able to stretch his fingers completely and there was minimal flexor contracture. Patellar and leuconychia was present. Diffuse scaling was present over scalp. Eyelashes of lower eyelids were decreased and there was decrease in eyebrows and scalp hairs. Discoloration of teeth was present.

Ocular examination shows scaly lids. Adnexa was found with bilateral ectropion accompanied with epiphora. There was incomplete closure of eyelids due to contracture along with presence of bell’s phenomenon. Visual acuity was also decreased with 6/18 in left and 6/12 in right.

Patient was admitted and treated with oxygenation, intravenous antibiotics, nebulisation with levosalbutamol and budesonide.

Patient was further investigated to rule out any cardiac or pulmonary causes of breathing difficulty. Investigations included complete blood count and urine examination was within normal limits. Mantoux test was negative. Spirometry showed grade III obstruction. Chest x-ray reported nothing significant. Electrocardiogram and echocardiography was within normal limit. Conservative treatment led to relief of respiratory distress. Patient was discharged with emollients, keratolytics, antihistaminic, vitamin A, appropriate eye drops with advice for follow-up. Patient was followed up after 21 days & reported with similar episode of breathing difficulty of 20 min duration precipitated by cycling a distance of around 100 meters, relieved on taking rest. Patient was advised the same treatment to continue and advised for regular follow-up.

![Figure 1: 14 years old boy with generalized ichthyosis](image1)

![Figure 2: Ectropion of both eyes with epiphora in 14 years old boy with lamellar ichthyosis](image2)

**Discussion & Conclusion**

The term ichthyosis is derived from the Greek word ‘ichthys’ meaning “fish” and refers to the similarity in appearance of the skin to fish scales. Early reports of ichthyosis in the Indian and Chinese literature dates back to several hundred years.\(^6\)

Six genes have been identified that cause non-HI ARCI: TGM (the gene encoding transglutaminase), ABCA12, NIPAL4 (also known as ICHTHYIN), CYP4F22, and the lipoxigenase genes ALOX12B and ALOX15. Transglutaminase mutations lead to abnormalities in the cornified envelope, whereas defects in ABCA12 cause abnormal lipid transport and those in CYP4F22 produce abnormal lamellar granules. The lipoxigenases are likely to play a role in epidermal barrier formation by affecting lipid metabolism.\(^2\)

Epidermal kinetic studies show increased epidermal mitotic activity and a rapid epidermal cell transit time. Scales are due to increased proliferation of epidermal cells.
Histopathology shows marked hyperkeratosis, hypergranulosis and moderate acanthosis with prominent rete ridges. Mitotic figures in the epidermis are frequent. A mild perivascular infiltrate is noted in the upper dermis. The clinical features of lamellar ichthyosis are typical & easily clinically identifiable. Interprofessional coordinated management of dermatologist, paediatrician, geneticist, opthamologist and physiotherapist play an important role in managing such cases. The management of a typical case is aimed at decreasing symptoms and include emollients (petrolatum, coconut oil, alpha hydroxyl acetic acid), Keratolytics containing salicylates with propylene glycol and local and systemic retinoids.[5]

The present case is unique in having lamellar ichthyosis with respiratory systemic involvement as evidenced by grade III obstruction of airways in Spirometry. Such case should be managed with focus on relieving respiratory symptoms with supportive treatment followed by regular monitoring of respiratory functions. Chest infections should be prevented with proper immunization and should be promptly managed with antibiotics if occurs. Nutritional supplements especially vitamin A should be provided. Efforts should be made to prevent respiratory complications and respiratory failure leading to morbidity and mortality.

References