Treatment Of Congenital Ichthyosis With Acitretin
A Cakmak, F Baba, S Cakmak, K Shermatov, H Karazeybek

INTRODUCTION
Ichthyosis belongs to the group of genodermatoses, characterized by hyperkeratosis and desquamation of the epidermis. Clinical manifestation is heterogeneous and depends on the type of the disease (1). Congenital ichthyoses which affect the entire body in the perinatal period include several major subtypes; harlequin ichthyosis, lamellar ichthyosis (LI), and congenital ichthyosiform erythroderma (2,3,4). Severe congenital ichthyosis which shows intense hyperkeratosis on whole body of the neonate include several distinct types of ichthyosis with different pathogenesis (2,3,4).

At the terminal differentiation of keratinocytes, a variety of proteins specific for keratinization are expressed and unique subcellular structures are formed. Some of these structures and proteins are thought to be closely associated with the pathogenesis of severe congenital ichthyosis.

Topical treatment of ichthyosis classically includes generous and frequent applications of emollients and keratolytic agents such as retinoic acid (0.1% cream) lactic acid, urea, a mixture of lactic acid and propylene glycol, and calcipotriol (5,6,7,8,9). Unpredictable transcutaneous absorption of topically administered drugs and potential teratogenicity and toxicity of systemic drugs therefore make it mandatory to innovate more efficacious and less toxic drugs for the treatment of ichthyosis. We assessed the clinical efficacy of acitretin treatment of congenital ichthyosis, and hereby report the successful treatment of this disease without any side effects in a newborn.

CASE REPORT
A 2850 g male infant was born to a 24-year-old woman after a pregnancy of 40 weeks gestation. The baby was the first child of the family and parents were first degree relatives. There was no family history of congenital abnormalities.

The mother had neither regular obstetric follow-ups nor any ultrasound examinations during pregnancy because of financial problems. She had a normal spontaneous vaginal delivery at home with help of a midwife. Postpartum APGAR scores were not determined.

At delivery, the infant was noted to have a thick keratin-coated yellow skin, split by deep and extensive fissures into polygonal plates. He had severe bilateral ectropion, eclubium, underdeveloped nasal bridge, rudimentary ears and sparse scalp hair. He also had flexure contractures with hypoplasia of the fingers and toes (Figure 1). The baby was referred to our hospital’s neonatal intensive care unit. The baby’s routine laboratory tests and abdominal ultrasonographic evaluations were within normal limits. The baby was kept in the incubator and fed with nasogastric tube and intravenous fluids. Ophthalmic and dermatologic treatments, vaporisation of incubator, and wide-spectrum antibiotic treatments were started.

Phototherapy was applied because of hyperbilirubinemia. Acitretin (1mg/kg/day) treatment was started. Pretreatment skin punch biopsy revealed a moderate degree of hyperkeratosis with complete absence of granular cell layer in the epidermis. Hyperkeratosis extended into the hair
follicles forming keratotic follicular plugs. (Figure 2: 200X, HE)

The patient was clinically followed up regularly for six months. A post-treatment rebiopsy was performed and it revealed significant improvement of epidermal changes as continuous granular cell layer (Figure 3: 200 X, HE).

In accordance with histologic improvements, the patient's clinical appearance improved significantly as well (Figure 4).

**Figure 1**
Figure 1: Pretreatment appearance of the patient.

**Figure 2**
Figure 2: Pretreatment punch biopsy material showing moderate degree of hyperkeratosis with complete absence of granular cell layer. Hyperkeratosis extends into the hair follicle forming keratotic follicular plug. (200X, HE)

**Figure 3**
Figure 3: Posttreatment punch biopsy showing continuous granular cell layer. (200 X, HE)
DISCUSSION

Many surviving congenital ichthyosis patients have been treated with retinoids and survival is often attributed to early therapeutic intervention. However, the role of retinoids in improved survival still remains to be determined (11,12,13,14). Many perinatal congenital ichthyosis deaths are caused by interference of severe infections causing loss of vital functions and the late onset of retinoid action; therefore, other factors such as improved intensive care of severely compromised neonates, are likely to be important in survival of the patient (15).

The clinical features of infants with congenital ichthyosis are generally similar to our case, which is typical. No visceral malformation is reported, the disorder consisting exclusively of an error of epidermal keratinization. Respiratory movements are restricted; secondary infection and septicemia are frequent complications. Death usually occurs in the first few weeks of life.

Prenatal diagnosis is now possible, using sonography, which shows ectropion, eclairbium, nasal or aural abnormalities, and hypoplasia of the extremities. However, a precise diagnosis needs fetal skin biopsy made between 20 and 22 weeks of gestation (17). If the number and size of the biopsy specimens are small, it may be possible to misdiagnose an affected fetus as normal (17). Prenatal diagnosis is vital in genetically inherited lethal diseases. According to the prenatal diagnosis, the patients should thoroughly be evaluated and the families be informed about the disease (18).

At light microscopic examination, a thick compact stratum corneum with parakeratosis, hypogranulosus and hyperkeratosis of the hair canals is seen (19). On ultrastructural examination, the granular layer keratinocytes present abnormalities of lamellar granules. Between the cornified cell and the granular cell layers, extracellular lamellar structures are lacking (20). In our case histopathologic evaluation revealed moderate degree of hyperkeratosis and complete absence of granular cell layer in the epidermis with keratotic follicular plugs.

Control biopsy performed after a acitretin (1 mg/kg/day/ 6 month) therapy showed that the prior findings had completely ceased.

Some rare long-term survivals of congenital ichthyosis are reported. This survival is often attributed to early treatment with retinoids. The clinical presentation then evolves with age into a viable form of generalized ichthyosis (21). However, the persisting dermatosis remains severe and the quality of life of affected children is poor.

El-Ramly et al (22) presented eight cases of ichthyosis treated with retinoids. They reported that aromatic retinoid treatment resulted in a satisfactory improvement of skin condition. Tamayo et al (23) also recommended oral retinoid treatment after their experience in eight children with lamellar ichthyosis.

Gicquel et al (24) reported bilateral ectropion treatment in an infant with severe lamellar ichthyosis by N-acetylcysteine applied directly to the skin with association of oral acitretin. Topical N-acetylcysteine has been proved to have an antiproliferative effect on keratinocytes in vitro and in vivo. It may be useful in the treatment of major forms of ectropion in children with lamellar ichthyosis. Its association with conventional acitretin treatment may prevent unnecessary surgery. In accordance with the prior studies in literature, our patient showed dramatic improvement in skin lesions and ectropion with oral acitretin treatment.

Verfaille et al. (25) investigated the efficacy, tolerability and safety of oral liarozole, a retinoic acid metabolism blocking agent, with acitretin in patients with ichthyosis. Between-group comparisons for efficacy and tolerability revealed no statistically significant differences except for scaling on the trunk at baseline which was significantly worse in the liarozole group and showed a more pronounced
improvement in this group than in the acitretin-treated patients. The expected retinoic acid-related adverse events were mostly mild to moderate and tended to occur less frequently in the liarozole group. The results of this trial warrant further clinical trials to confirm efficacy and safety of liarozole as an orphan drug in ichthyosis. Saracoglu et al. (25) treated two newborn infants with ichthyosis with acitretin (1 mg/kg/day). Clinical improvement was achieved shortly after treatment. They concluded that early management of severe ichthyosis cases could prevent life-threatening events.

Furthermore, long-term retinoid therapy is responsible for toxicity. Long-term therapy with oral retinoids provokes anxiety and shows adverse effects on bone mineralization. The most commonly reported effect is the production of osteophytes and calcification of ligaments (26). Paige et al (26) observed no evidence of skeletal toxicity in 42 children treated over an 11-year period. No complication due to oral acitretin treatment was observed in our patient.

In conclusion, early management of severe ichthyosis cases can prevent life-threatening events. In severe congenital ichthyosis cases, mortality due to the complications, is high in the neonatal period. Therefore, acitretin treatment seems to be the most efficient treatment in severe congenital ichthyosis with no major side effects and easy oral application.

CORRESPONDENCE TO
Alpay Cakmak, MD Harran University School of Medicine Department of Pediatrics TR-63100, Sanliurfa, TURKEY.
Tel: +90 414 3141170 Mobile Phone : +90 5326235956 e-mail: alpaycakmak@gmail.com Fax: +90 414 3151181

References
Author Information

Alpay Cakmak
Assistant Professor, Department of Pediatrics, Harran University School of Medicine

Fusun Baba
Assistant Professor, Department of Pathology, Harran University School of Medicine

Sevin Cakmak
Associate Professor, School of Medicine, Dicle University Department of Ophthalmology

Kabil Shermatov
Associate Professor, Department of Pediatrics, Harran University School of Medicine

Himmet Karazeybek
Professor, Department of Pediatrics, Harran University School of Medicine