Mutations in transglutaminase 1 gene in autosomal recessive congenital ichthyosis in Egyptian families

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Abstract. Autosomal recessive congenital ichthyosis (ARCI) is a rare heterogeneous keratinization disorder of the skin. It is clinically divided into 2 subtypes, lamellar ichthyosis (LI) and congenital ichthyosiformis erythroderma (CIE). We investigated forty-three ARCI Egyptian individuals in 16 severe LI, and 10 CIE families. We identified 5 alleles in two Egyptian families as having intron-5/exon-6 splice acceptor mutation recognized by the \textit{MspI} restriction endonuclease. This promoted to a frequency of 9.6\% for this mutation (5 splice-mutation alleles/52 alleles tested). We extended our previous dataset to update the detection of R142H mutation in 4 CIE Egyptian families and one LI phenotype (frequency of 28.8\%; 15/52), whereas we still had no R141H among our Egyptian population. There was no correlation between phenotype and genotype in our study. Surprisingly, the mutant alleles detected in intron-5 acceptor splice-site were associated with the other extreme of CIE phenotypes rather than the severe LI form. We clearly demonstrated that the ARCI Egyptian families in Upper Egypt was ethnically pure and had a tendency not to be a hybrid with other populations in Lower Egypt, Delta zone and Cairo city.

1. Introduction

Lamellar ichthyosis (LI) is clinically and genetically heterogeneous hereditary keratinization disorder of the skin, which causes abnormalities of the stratum corneum, the upper most layer of the epidermis. In most cases, LI is inherited autosomal recessively. A rare form, however, of autosomal dominantly inherited LI has been described as well [8]. The estimated incidence of autosomal recessive congenital ichthyosis (ARCI) is 1:150,000–300,000 livebirths [26].

Depending on the finding of erythema, two major types of recessive LI were clinically discriminated lamellar ichthyosis (MIM #242300) and nonbullous congenital ichthyosiform erythroderma [(NCIE) MIM #242100] [19,24]. The erythematous forms are characterized by prominent erythroderma and fine white, superficial, semiadherent scales. A ninety percent presents at-birth as collodion babies. Patients suffer from palmoplantar keratoderma, often with painful fissures, digital contractures, and loss of pulp volume. In half of the cases, a nail dystrophy including ridging, subungual hyperkeratosis, or hypoplasia has been described. Ectropion, eclabion (turning outward of the eye lids and lip, respectively), scalp involvement, and loss of eye brows and lashes seem to be more frequent in NCIE than in lamellar ichthyosis [9].

In contrast, for a non-erythematous form presenting at-birth with a collodion-like membrane encasing the neonate, the skin later develops large, brown, plate-like scales covering the entire body. Patients may have palmar and plantar hyperkeratosis and significant tautness of the facial skin, which may be associated with ectropion and eclabion. If left untreated, severe ectropion can lead to blindness. Many affected persons exhibit scarring alopecia and secondary hypohidrosis [24]. Two loci for LI had been mapped: LI1, which is caused by mutations in the \textit{TGM1} gene on 14q11 [18] and LI2, which maps to 2q33–q35 [15]. Fischer et al. [12] identified 2 additional loci for autosomal recessive ichthyosis. One of these was LI3 mapping to chromosome 19 and the other was an erythrodermic form mapping to 3p21. He reported that linkage to 1 of the 4 loci could be demonstrated in more than half of 51 consanguineous families, most of them from the Mediterranean basin [12]. All 4 loci could be excluded in the others, implying further genetic heterogeneity.
NCIE can be caused by mutation in the TGM1 [5], arachidonate 12-lipoxygenase, R type; (ALOX12B) and the arachidonate lipoxygenase 3 (ALOXE3) [25] Jobard et al. [7] identified mutations in lipoxygenase-3 (ALOXE3) and 12R-lipoxygenase (ALOX12B) genes in NCIE linked to chromosome 17. A rare form of NCIE, Chanarin-Dorfman syndrome caused by mutation in the comparative gene identification 58 (CGI58) gene was identified the CGI58 gene in the critical region for triglyceride storage disease with impaired long-chain fatty acid oxidation on chromosome 3p21 [3].

A stable cell envelope formed of several components that are covalently bound to each other characterizes the horny cells of the stratum corneum. The components include a number of proteins, such as volucrin, loricrin, and the small proline-rich protein along with lipids on the outer surface of the cell envelope. Transglutaminases are calcium-dependent thiol enzymes that catalyze covalent cross-linking of proteins, thus enhancing the stability of biological structures. These enzymes are present in various tissues and body fluids, such as coagulation factor XII (FXII) in the blood stream, tissue transglutaminase in red blood cells, liver cells and chondrocytes and keratinocytes and epidermal transglutaminases in the skin [4]. Keratinocyte transglutaminase (TGK) is encoded by the transglutaminase 1 gene (TGM1) on chromosome 14q11 [10]. Moreover, epidermal transglutaminase, which is encoded by the transglutaminase3 gene (TGM3), is present in suprabasal keratinocytes. TGM3 has been localized to chromosome 20q11–q12 [22,27]. TGM1 catalyzes the formation of covalent ε-(γ-glutamyl)lysine cross-links between the precursor proteins of the cornified cell envelope (CE). The CE is a 15–20 nm-thick, highly insoluble structure that is formed, during the late phases of epidermal differentiation, on the inner side of the plasma membrane. Several proteins, such as volucrin, loricin, and the small proline-rich proteins, have been implicated as precursor proteins to the CE. Simultaneously with cross-linking of these precursor proteins, 1,6-hydroxyacylsphingosine lipids covalently bound to the outer surface of the protein CE and the plasma membrane are replaced by the CE [5]. In patients with autosomal recessive LI, a lack of TGK expression has been shown [1], and linkage of autosomal recessive LI to TGM1 has been described [18]. Subsequently, mutations have been identified in the TGM1 in patients with LI [14]. Normal transglutaminase activity and a normal TGM1 gene sequence are found in some patients with lamellar ichthyosis, indicating genetic heterogeneity, which was previously suggested on the basis of the ultrastructural analysis of skin biopsies [23]. The splice site mutation is the common TGM1 mutation in congenital recessive ichthyosis patients in the Norwegian population [21].

Our collaborative joint with dermatologists and molecular geneticists of the NIH skin staff since 1992 promoted fruitful results to map the disease locus for the severe autosomal recessive LI to chromosome 14q11 [17] and showed complete linkage of LI disease with TGM1 [18] Those studies had identified no mutations due to R141H in exon 3, whilst they had been identified the R142H-LI mutation in Egyptian population [17].

2. Subjects and methods

2.1. Selection and categorization of patients

Thirty patients and thirteen unaffected family members from twenty-six families were studied. The diagnosis of autosomal recessive congenital ichthyosis, on the clinical basis, was made at Medical Genetics Center, Ain Shams University Cairo Egypt following a medical and dermatological history. The 26 families were categorized as sixteen LI and ten CIE type. They were fifteen males and fifteen females. Their onset ages ranged from one day to six years. The 26 ARCI Egyptian families were geographically mapped along with the governorates of Egypt according to the parents’ birth origins.

2.2. Isolation of DNA

Genomic DNA was extracted from peripheral venous blood-EDTA using QIAamp DNA Mini kit (Qiagen) according to the blood and body fluid spin protocol. In some cases, DNA was prepared in situ by gentle scraping the buccal mucosa for 30 s using a cytobrush [2]. The cells obtained were treated directly with diluted NaOH solution, heated, and neutralized with Tris-Cl, pH 8.0 [17].

2.3. PCR amplification of genomic DNA

To investigate mutations in exon 3 of TGM1 gene, we amplified this exon using oligonucleotide primer pairs designed by Russell et al. [17] Both genomic DNA (200 ng) prepared from the whole blood or from buccal mucosa (5 µl), primers (100 nM each), were premixed with 200 µM dGTP, dATP, dCTP, dTTP in 10 nM Tris-
HCl, 50 mM KCl, 1.5 mM MgCl$_2$ in a total volume reaction of 30 µl. The Red Taq® enzyme 1.5 U was added to the reaction mixture and denatured at 94° for 2 min., then they are amplified at 94° for 60 s, 60° for 30 s, and 72° for 90 s, and an extension for 10 min after 35 cycles. Amplification products from each primary PCR (295-bp) was diluted 1:500, and 5 µl was re-amplified using the same primers and conditions in a 100-µl-reaction mixture. The secondary PCR amplicons were separated on 2% agarose gel (Promega) and purified, if necessary, with the Wizard PCR Prep kit (Promega).

2.4. Restriction enzyme analysis

The PCR amplicon was subjected to cutting with restriction enzymes ApaLI and AciI. If R141H mutation is present, the 295-bp product will be cleaved into 142 and 153 fragments, otherwise no cleavage will be the case. Using 10 units AciI endonuclease, the 295-bp product would be digested into four fragments 107, 100, 53 and 35 on a 2% MetaPhor (BMA) gel. The mutation removes two adjacent AciI sites so that digests of affected individuals reveal the presence of a 160-bp bands and loss of the fragments. Heterozygotes have both the 160-bp and 107-bp bands.

2.5. Intron 5 splice acceptor mutation analysis

To assess the presence of the intron 5 splice acceptor site mutation in TGM1, genomic DNA (200 ng) prepared from the whole blood or from buccal mucosa (5 µl) was amplified with Red Taq enzyme (Sigma) using the oligonucleotide pair TGM1mutF and TGM1mutR [28]. PCR was performed for 35 cycles of 94° for 15 s, 60° for 30 s, and 72° for 45 s with extension at 72°C (PCRExpress, Hybaid). The products were analyzed on 2% agarose gel, and DNA fragments were cut out and purified, if necessary. The TGM1 intron 5 acceptor splice site mutation was confirmed in affected individuals and carriers by MspI digestion [20]. The mutation created an additional MspI restriction site, so that cleavage of the 467-bp PCR fragment generated from genomic DNA resulted in two specific fragments of 298-bp and 169-bp. The digested PCR products were electrophoresed on 2% MetaPhor agarose.

3. Results

3.1. Phenotypic expression

The twenty-six families with ARCI were investigated clinically and molecularly. The clinical picture of the patients analyzed was highly variable. Seven of them were born as collodion babies, three showed joint contractures and the last showed alopecia. Fourteen patients suffered from erythema accompanied in the more severely affected cases by ectropion, whereas sixteen patients from different families were presented with the non-erythrodermic form of LI. The type of scaling was highly variable, ranging from fine whitish scales to dark brownish scales. All but two showed positive consanguinity (3 were 3rd degree relatives and 21 were 2nd degree relatives) and fifteen showed positive family history. The clinical features of the affected ARCI were represented in Table 1.

3.2. Mutational analysis

3.2.1. R142H mutation in TGM1 gene

We detected a homozygous R142H-mutation in exon 3 detected by AciI recognition site in the LI-proband (I-59) of 2.5-year-old. The patient was the 1st offspring of consanguineous parents with 2 similarly affected females and one neonatal-death male with the same condition. The presenting symptoms started since birth.
with generalized erythroderma followed by scaling of skin, dark scales affecting the entire body with exaggeration over flexural and pressure areas with itching associated with flexion deformity of the fingers and deviation of hands. The erythroderma was attenuated on treatment with oral vitamin A and topical urea. The unaffected father (I-69) revealed a heterozygous R142H mutation.

A 2-year-proband classical CIE type (III-46) revealed a homozygous R142H mutation recognized with Acrl enzyme. He had had multiple skin ulcerations together with scales distributed all over the body, ectropion and eczema including scalp scarring (Fig. 1) with a collodion baby on birth. Treatment with oral vitamin A, topical use of urea, alpha- and poly-hydroxy acids, showed an excellent response on following up during one year. His mother (III-70) revealed a heterozygous R142H mutation towards the same R142H mutation.

A 12-year-old female CIE patient (VI-16) showed homoygosity towards the R142H mutation. She was presented with dry rough scaly skin covering the whole body with dark areas over the elbows and knees, starting the case since birth. Her affected mother (VI-17) showed a heterozygous R142H mutation manifesting mild scaling localized to the legs and arms and roughness of skin of palms and soles. They were treated with oral vitamin A, and topical hydroxy acids and urea showing a good response. A female patient of 20 years (VI-26) was birthed with a collodion membrane that shed off after one week leaving erythrodermatous surface followed by scales covering all over the body, more severely on the pressure areas (elbows & knees). In winter, the condition worsens where the scales are moderate in size, adherent and soft. The symptoms had followed with attacks of tonic-clonic convulsions and psychomotor epilepsy associated with xeroderma pigmentosum showing multiple brownish macules over the upper part of the neck, trunk and hands. This case was fairly attenuated with oral vitamin A. It was of interesting to note that the splice site mutation abnormally existed in family VI besides R142H mutation common in this family (Fig. 2).

Family (VII) showed an inclusion of homozygous R142H-mutation CIE patient (VII-20) with one-day age having a dry rough scaly skin all over the body with erythema and scaring with deformity of hands and feet, scarring the scalp, palm and soles. Also, he had not a developed ear, depressed nasal bridge and a severe ectropion. Treatment included oral vitamin A and topical keratolytics. The heterozygous mother (VII-21) was followed up in our dermatological clinic for genetic counseling for a neonatal-death female with a similar case (not examined).

A CIE-family showed a proband (IX-28) of 1.5-year-old that is the first sibling of two consanguineous parents. At birth, he was covered with a collodion membrane that shed off leaving scales all over the body with generalized erythroderma. The scales were small, dark and adherent. The patient was treated with topical glycolic acid, lactic acid and urea. The unaffected mother (IX-29) has a heterozygous R142H mutation.

### 3.2.2. Intron 5 splice acceptor site mutation

Two individuals with autosomal congenital recessive ichthyosis (ARCI) and a mutation in the intron 5/exon 6 canonical splice site in TGM1 were identified in two families.

Family VI was intensively investigated where it included 5 phenotypically CIE patients and their mild affected mother (VI-17) living in the far south of Upper Egypt. We detected here only one homozygous sibling (VI-64) as having intron 5 splice acceptor mutation. She is a one-year-old female CIE patient being the 4th offspring of proband sibship of consanguineous

### Table 1

<table>
<thead>
<tr>
<th>Case</th>
<th>Sex</th>
<th>Age</th>
<th>Skin at birth</th>
<th>Scales</th>
<th>Phenotypes</th>
<th>Severity of TGM1 disease</th>
<th>Treatment history</th>
</tr>
</thead>
<tbody>
<tr>
<td>I-59</td>
<td>Male</td>
<td>2.5y</td>
<td>? +</td>
<td>LI</td>
<td>+</td>
<td>+</td>
<td>VA, U</td>
</tr>
<tr>
<td>III-46</td>
<td>Male</td>
<td>2y</td>
<td>Collodion +</td>
<td>CIE</td>
<td>+</td>
<td>+</td>
<td>SPB, VA, AHA, PG</td>
</tr>
<tr>
<td>VI-16</td>
<td>Female</td>
<td>12y</td>
<td>Collodion +</td>
<td>CIE</td>
<td>+</td>
<td>+</td>
<td>VA, U, AHA</td>
</tr>
<tr>
<td>VI-17</td>
<td>Female</td>
<td>30y</td>
<td>-</td>
<td>CIE</td>
<td>-</td>
<td>+</td>
<td>U, AHA</td>
</tr>
<tr>
<td>VI-26</td>
<td>Female</td>
<td>20y</td>
<td>? +</td>
<td>CIE + XPb</td>
<td>-</td>
<td>+</td>
<td>VA, PG</td>
</tr>
<tr>
<td>VI-64</td>
<td>Female</td>
<td>1y</td>
<td>-</td>
<td>CIE</td>
<td>-</td>
<td>-</td>
<td>U, PG, LA</td>
</tr>
<tr>
<td>VII-20</td>
<td>Male</td>
<td>1d</td>
<td>-</td>
<td>CIE</td>
<td>+</td>
<td>+</td>
<td>VA, PG, LA</td>
</tr>
<tr>
<td>IX-28</td>
<td>Male</td>
<td>3.5y</td>
<td>Collodion +</td>
<td>CIE</td>
<td>-</td>
<td>+</td>
<td>AHA, PG, U</td>
</tr>
<tr>
<td>X-40</td>
<td>Female</td>
<td>3.5y</td>
<td>-</td>
<td>CIE</td>
<td>-</td>
<td>-</td>
<td>AHA</td>
</tr>
</tbody>
</table>

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a Age at time of diagnosis.
b Xeroderma pigmentosum associated with convulsions.

VA = vitamin A, U = urea, AHA = alpha-hydroxy acids, PG = polyethylene glycol, SPB = sodium phenobarbione, LA = lactic acid.
parents. She was presented with generalized small fine white scales all over the body. The condition was started at a 3-day-age by roughness and dry scaly skin over the wrist joint then became generalized. The condition was associated with tonic-clonic convulsions started at the age of 10 m. The proband was fairly improved with a phenobarbitone salt as an anticonvulsant, and topical urea. Another 3.5-year-old CIE female (X-40) showed the same splice-site mutation in intron 5/exon 6 manifesting a homozygous mutation. Her condition started to be noticed at birth by dry rough scaly skin all over the body associated with itching and erythema. Her mother (X-44) showed a heterozygous allele manifestation.

4. Discussion

4.1. Geographical analysis

The ARCI twenty-six patient families (43 individuals) were selected from the genodermatosis database.
records in ASMGC. Those families were categorized as having severe LI and CIE the other extreme of the disease. The demographic distribution within the regional map of Egypt revealed that the frequency of the ARCI families in Upper Egypt was lower than in Nile Delta zone, Cairo, and Lower Egypt. This might be due to the tendency of Upper Egyptians to be isolated from other governorates of Egypt due to restricted habits and social rules. This was clearly confirmed by the existence of only one CIE family (VI) living in the far south to Aswan (Fig. 2). Unfortunately, there are no TGM1 mutational studies in the Arabs’ area, except those in Northern Africa due to TGM2 gene on chromosome 2q34, (15, 16) though there is a strong observation for an inclusion of ARCI in the Gulf area.

4.2. Mutational analysis

We screened both LI and CIE individuals for the intron 5 splice acceptor mutation previously described by Huber et al. [20] Surprisingly, we have investigated and identified three-CIE-patient-Egyptian individuals having splice acceptor mutation in the TGM1 due to a change of A to G at position 56 in genomic DNA. Therefore, this indicated that that mutation has not been limited to the Norwegian and German populations [28]. Of 26 families from whom DNA were available, we identified 2 families who carried the intron 5 G>A acceptor-site mutation revealing 2 homozygous and 1 heterozygous alleles. This gives a frequency of 9.6% for the splice mutation allele in our data (5 splice-mutation alleles/52 alleles tested). This is a clinically high heterogeneous group of families and no conclusions about the frequency of this mutation in ARCI can be derived directly from this dataset.

After identifying the splice-site mutation in Egyptian ARCI patient’ families, we moreover investigated R141H and R142H in exon 3 of the TGM1. Extra genomic DNA screening for R141H of the TGM1 revealed the absence of this mutation in Egyptian population which confirmed the previous pilot study [17]. An extension to the previous results discussed by the ASMGC-NIH joint staffs was carried out using the newly banked genomic DNA samples. We identified 5 homozygous probands carrying the R142H mutation and 5 heterozygous carriers with a frequency of 28.8% (15 mutant R142H alleles/52 alleles tested). The mutation in each of the five families segregated with the disease. In those families G to A transitions in CGC codons in exon 3 results in single amino acid substitutions either R141H or R142H. Again, deamination of 5-methylcytosine associated with methylation of CpG dinucleotides has been suggested to interpret the high frequency of \( C > T \) or \( G > A \) transitions causing the genetic disease [11].

4.3. Genotype/phenotype correlation

Autosomal recessive LI is characterized by remarkable clinical heterogeneity. An erythematous and a nonerythematous type of LI were clinically distinguished [19,24] Moreover, patients might exhibit different clinical patterns and colors of the scales, they might or might not have an inclusion of palmoplantar hyperkeratosis, and not all of them were born as colloidion babies. The whole spectrum of LI variants, however, was also seen in TGM1 LI patients (Table 1) in family IV (Fig. 2). Seeking a genotype/phenotype correlation for specific mutations in TGM1 gene, we were able to compare two patients who carry the same TGM1 genotype. Both III-46 and X-40 were homozygous for the splice 5 intron acceptor site mutation, but they differed in their clinical picture despite having erythematous skin scales as illustrated in Table 1. Hence, there were no conclusive criteria for the clinical differentiation of the two extremes of LI and CIE phenotypes. The phenotype/genotype correlation and heterogeneity of the ARCI may be identified and interpreted by extending studies to other loci linked to the disease.

4.4. ARCI treatment

In this study, we obtained detailed clinical information about each of the patients, including topical treatment and oral medication history particularly the oral vitamin A, topical alpha- and poly-hydroxy acids and ureas. We have observed for the initiation of the effective therapy, 50 to 100 mg vitamin A have to be administered orally which is the upper limit of toxic effects. The oral dose may be optimally reduced to 20 mg twice daily and after 4 weeks to about 10 mg twice daily. Stopping the treatment and reappearance of ichthyotic lesions, oral vitamin A at a lower dose 10 to 20 is well tolerated by most patients with satisfactory results. It was shown that urea strongly increased water-binding capacity of the scales several days after the last treatment and the transepidermal water loss was reduced [13]. It is noteworthy that the high concentration of urea in a basic cream represents highly potent agent than other keratolytics for treating ichthyosis vulgaris-like phenotypes. Moreover, alpha- and poly-hydroxy acids restore toward normal the hyperkera-
totic skin of the ichthyoses. The sequence of the histological and clinical changes in lamellar ichthyosis suggests that the effect of those hydroxy compounds on the skin are mediated by means of their influence at the level of underlying epidermis in so far as after several days of topical treatment a normal skin surface abruptly appears clinically, consequent to sheet-like desquamation of the entire thickened stratum corneum without dissolution [6].

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References


