Research Article

Molecular analysis of the first reported hereditary lymphedema-distichiasis case in Bangladesh

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ABSTRACT

Lymphedema-distichiasis syndrome (LD, OMIM 153400) is hereditary primary lymphedema with autosomal dominant nature of inheritance and variable expression. LD is characterized by late childhood or pubertal onset of lower limb lymphedema and an aberrant second row of eyelashes (distichiasis) arising from the meibomian glands. Among the molecular reasons behind this condition are the mutations in the FOXC2 gene, a forkhead transcription factor, that plays a role in the formation of lymphatic and vascular systems. In this study, we report the first case of LD from Bangladesh with classical lymphedema-distichiasis syndrome with an eight-base-pair deletion in the FOXC2 gene. ClinVar accession code for this deletion is RCV000007679.3. FOXC2 protein is 501 amino acids long. This deletion of 8 bp (ACGCCGCC) causes frameshift of codons after amino acid number 304. The frameshift creates an altered truncated protein with 154 new amino acids after codon 304. We assume that these changes in the protein may affect its function contributing to the disease manifestations. Further research may confirm these assumptions.

Introduction

Lymphedema-distichiasis syndrome (LD) [MIM 153400] is a form of hereditary lymphedema in which lymphedema, primarily of the limbs, with variable age at onset, is seen together with distichiasis, or double rows of eyelashes. The extra eyelashes grow from the meibomian glands and may protrude into the cornea, producing severe corneal abrasions. Various additional complications such as cleft palate, cardiac defects, abnormal curvature of the spine, droopy eyelids, etc (Szuba and Rockson, 1998) have also been observed.

The disease has been mapped to 16q24.3 (Mangion et al., 1999) where the forkhead/winged-helix transcription factor FOXC2 is located. Studies showed that dominant mutations in the FOXC2 gene (MIM602402), cause lymphedema with variable age of onset (range: 7-40 years), often associated with distichiasis (Fang et al., 2000). Key roles of FOXC2 include regulating differentiation of lymphatic endothelial cells, formation of smooth muscle cell layers, and morphogenesis of lymphatic valves. Along with VEGFR-3, FOXC2 acts to establish distinct features of the lymphatic vascular architecture (Sabine and Petrova, 2014).

In both humans and mice, FOXC2 is highly expressed in the developing lymphatic vessels, as well as in the adult lymphatic valves (Kriederman et al., 2003). Its critical role in lymphatic vascular development has been discovered by the manifestation of abnormal lymphatic patterning and the absence of proper lymphatic valves in Foxc2-deficient mice. LD patients develop similar defects characterized by lymph and venous reflux, indicating failure or absence of lymphatic and venous valves (Mellor et al., 2007).

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FOXC2 is transcribed to a 2.2 kb transcript containing a 1.5 kb single exon coding region. The FOXC2 protein contains 501 amino acids. The most characterized region in the gene (Fig. 1) is the fork-head DNA binding domain (FHD, amino acids 71 to 162). It also contains a nuclear localization signal (NLS1, amino acids 78-93). At the N-terminal, there is a transactivation domain 1 (AD-1) starting from the first amino acid until the FHD (amino acid 71). In the C-terminal, a second transactivation domain (AD-2, amino acids 395-494) and an inhibitory region (ID-2, amino acids 495-501) have been identified (Lam et al., 2013). In the central region of FOXC2 protein, after the nuclear localization signal 2 (NLS2, amino acids 168-176), some phosphorylation and SUMOylation conserved sites have been recently identified (Danciu et al., 2012; Ivanov et al., 2013).

In LD patients, almost 70 different FOXC2 mutations have been reported to date, scattered randomly along the whole coding sequence. The majority of FOXC2 mutations are small insertions or deletions and nonsense mutations causing truncated proteins (Brice et al., 2002). It probably creates a haploinsufficiency condition which explains the dominant nature of LD. The FOXC2- haploinsufficient state is associated with hyperplasia and distichiasis in mice (Kriederman et al., 2003). Mutations responsible for the disease greatly vary among affected families and the secondary phenotypes differ among families. Nonetheless, any strong correlation has not yet been established between allelic variants and phenotypes. The disease exhibits variable penetrance of among the family members carrying the same allelic variant.

In this study, the FOXC2 gene of an individual diagnosed positive for lymphedema distichiasis was analyzed by targeted Sanger Sequencing. The whole exon of 1501 bp was amplified through primer walking using five sets of primers. Due to the high GC percentage of the gene, the primers were not efficient and the PCR reaction mixture needed the addition of certain additives and nested PCR was performed. It is the first molecular study of a Bangladeshi lymphedema distichiasis patient and a homozygous 8 base pair deletion has been identified.

Materials and methods

The case

A 32-year-old normotensive, non-diabetic, male presented with a history of bilateral below knee swelling since the age of twelve. He also noted foreign body sensation in both eyes for the last 10 years and the recent appearance of whitish spots on the left eye with impairment of vision of the same eye for about a month. Patient’s father and one of his younger brothers also showed the above-mentioned clinical features of eye lashes in the inner side of the eye lids and swelling of the lower limbs, albeit relatively milder. His sister is normal and does not exhibit any of the clinical problems. These characteristics indicated that the patient may be suffering from a certain type of hereditary lymphedema.

Mutational analysis of FOXC2 gene

Genomic DNA isolation

Genomic DNA was isolated from a 200 µl blood sample of the patient using GeneJet™ DNA extraction kit following the standard isolation protocol. Concentration and purity of DNA was sufficiently good for PCR and sequencing.

DNA amplification

Due to absence of any reported mutational hot spot and presence of only one exon in this gene, it was planned to sequence the whole gene using a method called primer walking. Five sets of primers were designed to amplify 5 products that overlap each other to cover the whole gene (Table 1, Fig. 5[c]).

All PCR reactions were performed using NEB PCR kit following standard reagent composition and reaction condition except template amount, annealing temperature and extension time. These were variable.
based on genomic DNA concentration, Tm of primer and product size of each PCR reaction.

It was also necessary to add DMSO (5% of volume) considering the high GC percentage of the target region. Detailed reagent composition and reaction condition of the PCR reactions are mentioned in Table: 2 and Table: 3. For retrieving full sequence, the following forward and reverse primer combinations were used: F1+R1, F2+R2, F3+R3, F4+R4, F5+R5, F1+R5, F2+R4. As the Tm value of each primer was very close, the annealing temperature of those primer combinations was set as 60 °C (Table 1, Table 3). PCR products were purified using ATP PCR/Gel DNA extraction kit following standard protocol.

**Sanger sequencing of PCR products**

At first, cycle sequencing was performed with the purified PCR amplified DNA. Big Dye Terminatortv1.1 Cycle Sequencing Kit and the forward or reverse primer of the respective product were utilized. Conditions for this cycle sequencing on thermocycler were: 1 minute for an initial denaturation of the DNA at 96 °C, followed by 35 cycles of a 10-second denaturation at 96 °C, variable annealing temperature based on the primers used for 5 seconds, and the extension step at 60 °C for 4 minutes. After this, capillary electrophoresis was performed on the 3130 Genetic Analyzer. The full process of sequencing was conducted in Advanced Molecular Biotechnology Laboratory, Department of Genetic Engineering and Biotechnology of University of Dhaka.

**Results and Discussion**

**Clinical features and diagnosis**

The patient was admitted with characteristics indicating a certain type of hereditary lymphedema. All the parameters of the general examination including the vitals were normal. Examination of lower limbs revealed bilateral edema of legs and feet (Fig. 2[a]) with dryness and hyperpigmentation of the overlying skin. Examination of the eye revealed partial ptosis on the left side, an extra partial set of eyelashes on both eyes (Fig. 2[c] and Fig. 2[d]), and left corneal opacity with reduced visual acuity on the left eye. Slit-lamp examination of both eyes showed bilateral distichiasis, epithelial and stromal opacity in the center of the cornea, and superficial corneal vascularization on the left eye. Results of other systemic examinations were normal. Isotope Lymphoscintigraphy (Fig. 2[b]) of both lower limbs revealed grade-1 and grade-2 lymphedema on the left and right sides respectively. Findings of Serum Creatinine, ALT, Prothrombin Time, Urine R/M/E, ECG, Echocardiogram, and CXR were normal. ICT for filaria was also negative. A Duplex study of both lower limbs found no evidence of deep vein thrombosis and no evidence of arterial insufficiency.

<table>
<thead>
<tr>
<th>Primers</th>
<th>sequence</th>
<th>length</th>
<th>Tm(°C)</th>
<th>Size(bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>F1</td>
<td>GAGCCGTCTCGGAAGCAG</td>
<td>18</td>
<td>60.66</td>
<td>401</td>
</tr>
<tr>
<td>R1</td>
<td>TCGTTGACGAGAGTTG</td>
<td>20</td>
<td>60.32</td>
<td></td>
</tr>
<tr>
<td>F2</td>
<td>TTCATCATGGAGCCCTTCCC</td>
<td>20</td>
<td>60.11</td>
<td></td>
</tr>
<tr>
<td>R2</td>
<td>ATGTTCACGCTGAAGCC</td>
<td>20</td>
<td>60.47</td>
<td>524</td>
</tr>
<tr>
<td>F3</td>
<td>CAAGGGAGCCGAGAGAAG</td>
<td>19</td>
<td>60.1</td>
<td>564</td>
</tr>
<tr>
<td>R3</td>
<td>GTGGGTGCTGTTGTTGTTG</td>
<td>18</td>
<td>62.7</td>
<td></td>
</tr>
<tr>
<td>F4</td>
<td>ATCATGACCCCTGCGAAG</td>
<td>18</td>
<td>60.7</td>
<td>598</td>
</tr>
<tr>
<td>R4</td>
<td>TGCCACTCACCTGGGACT</td>
<td>18</td>
<td>60.3</td>
<td></td>
</tr>
<tr>
<td>F5</td>
<td>GCCTCCTGGTATCTCAAAC</td>
<td>20</td>
<td>60.1</td>
<td></td>
</tr>
<tr>
<td>R5</td>
<td>TCTCTGCAGCCCTTAAA</td>
<td>20</td>
<td>60.3</td>
<td>328</td>
</tr>
</tbody>
</table>
Based on these clinical features and the patient statement of his father and younger brother having the similar illness, he was diagnosed with a rare hereditary disease, Lymphedema distichiasis syndrome. After necessary documentation, blood sample from the patient was collected to test for the presence of FOXC2 mutations.

### Table 2. PCR reagent composition

<table>
<thead>
<tr>
<th>Reagents</th>
<th>Volume (µl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>10X Reaction Buffer</td>
<td>2.5</td>
</tr>
<tr>
<td>dNTP solution</td>
<td>0.5</td>
</tr>
<tr>
<td>Forward primer</td>
<td>0.5</td>
</tr>
<tr>
<td>Reverse primer</td>
<td>0.5</td>
</tr>
<tr>
<td>Taq Polymerase (5U/µl)</td>
<td>0.125</td>
</tr>
<tr>
<td>DMSO (5%)</td>
<td>1.25</td>
</tr>
<tr>
<td>Nuclease free H₂O</td>
<td>19.625</td>
</tr>
<tr>
<td></td>
<td>25 µl</td>
</tr>
<tr>
<td>Total</td>
<td>25</td>
</tr>
</tbody>
</table>

### Table 3. PCR reaction conditions

<table>
<thead>
<tr>
<th>Step</th>
<th>Temperature (°C)</th>
<th>Duration</th>
<th>Cycle</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial denaturatn</td>
<td>95</td>
<td>4 min</td>
<td>1</td>
</tr>
<tr>
<td>Denaturation</td>
<td>95</td>
<td>30 sec</td>
<td>35</td>
</tr>
<tr>
<td>Primer annealing</td>
<td>60</td>
<td>30 sec</td>
<td></td>
</tr>
<tr>
<td>Extension</td>
<td>72</td>
<td>Variable</td>
<td></td>
</tr>
<tr>
<td>(1 min for 1kb)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Final Extension</td>
<td>72</td>
<td>5 min</td>
<td>1</td>
</tr>
</tbody>
</table>

PCR results of FOXC2 gene

PCR amplification of the 5 primer sets showed specific bands only for pair numbers 1 and 5 (Fig. 3[a]). Regions covered by primer pairs one and five were sequenced (Fig. 4).

To reveal the sequence between, several modifications were tried. Interestingly, gel electrophoresis indicated amplification from primer combination of F1 and R5 with 5% DMSO (Fig. 3[b]). Annealing and extension temperatures were 60 °C and 72 °C respectively for this reaction. Later, this product was used as a template to perform nested PCR with primer combination of F2 and R4. In this nested PCR reaction mix, the annealing temperature was 60 °C and 3% DMSO was added. Nested PCR with primer combination of F2 and R4 resulted in better amplification but nonspecific products were present (Fig. 3[c]).

The band of the desired size of 1120 bp was isolated using Gel extraction after electrophoresis. This 1120 bp DNA was sequenced using the Sanger method from both primers. PCR products of primer pair F1+R1, F2+R4, and pair F5+R5 overlap each other and cover the whole coding region of the FOXC2 gene. So, these three products were sequenced to get the complete sequence of the gene (Fig. 4).

Sequence analysis of FOXC2 gene and locating deletion mutation

Primer pairs of PCR products that provided good quality sequence are F1+R1, F5+R5 and F2+R4. When blast search against the NCBI nucleotide database were performed, no mutation was found in the sequence derived from pair F1+R1 and F5+R5. Product of primer combination F2+R4 was sequenced with both F2 and R4 primer to get the sequence of the whole segment. Sequence from R4 was reverse-complemented and after trimming they were assembled in full 1120 bp sequence. When the assembled sequence was blast searched, an 8 bp deletion (ACGCCGCC) was identified (Fig. 5[a]). The chromatogram also indicates that the patient is homozygous for this deletion (Fig. 5[b]). This
deletion is identified to be between base no 6423 and 6431 of RefSeqGene on chromosome 16, (which refers to mRNA position between 914 and 921). This mutation had been reported before for the same disease. ClinVar accession code for this deletion is RCV000007679.3. Although most other mutations of this gene are unique among different families, as of October, 2021 only this 8 bp deletion has been reported four times in four independent studies for families from different geographical region (Bell et al., 2001; Erickson et al., 2001; Bahuau et al., 2002; Brice et al., 2002). These changes resulted in the production of a premature stop codon that terminated the predicted protein earlier than the wild-type and produced novel C-termini. Codons after 304 are disrupted for this mutation and new 154 amino acids are added.

In contrast to most other mutations, which appear to be unique, the eight-base-pair deletion reported in this study was observed in four additional unrelated pedigrees with lymphedema and distichiasis, suggesting that it is recurrent. One study reported this deletion in a French family showing autosomal-dominant segregation of upper- and lower-eyelid distichiasis in seven affected relatives over three generations, in addition to below-knee lymphedema of pubertal onset in three (Bahuau et al., 2002). Two children had cleft palate as well as distichiasis, but without any association with the Pierre–Robin sequence. Divergent strabismus and early-onset myopia were other ophthalmologic anomalies. Another study reported this deletion in a British family with the father and all three of his children being affected with both distichiasis and bellow-knee lymphedema (Bell et al., 2001). Another study reported this mutation in fourteen-year-old boy with distichiasis, cleft palate webbed neck and very early onset (6 week) lymphedema (Erickson et al., 2001). The patient of this study stated he developed lymphedema during his puberty. He had distichiasis in the lower lids of his both eyes. He did not have myopia or cleft palate like some other LD patients with this deletion. Being reported in this study makes this 8 bp deletion the only FOXC2 mutation that has been found in five totally unrelated LD affected families. The reason behind the recurrence of this 8-base-pair-deletion is not fully clear. It indicates that the region is prone to DNA polymerase slippage during replication process. In fact, high GC percentage has been reported to make DNA more prone to deletion through polymerase slippage and recombination which partly explains why most of the FOXC2 mutations are frameshift deletions. Elucidating the reason behind this particular deletion being relatively more frequent might shed light on to important features that make a sequence prone to deletion and has implications in mutational hotspot identification.
Fig. 2. Clinical presentation of a 32-year-old Bangladeshi male with lymphedema distichiasis, (a) the lymphedema of the legs and feet, (b) Lymphoscintigraphy image of both legs, (c, d) Distichiasis in lower eye lids of the left and right eye respectively.

Fig. 3. (a) PCR amplification with all FOXC2 primer pairs. Among these, good quality sequence was determined from pair 1 and pair 5; (b) Gel electrophoresis of PCR amplification with DMSO, band was visible only for primer pair: F1+R5. The band of 1.5 kb size (indicated by the white box) was extracted from gel to use as template for nested PCR; (c) Nested PCR with product of F1+R5 as template, PCR product of primer F2+R4 was loaded in the gel and band of 1120 bp is visible with some nonspecific smears.
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Fig. 4. PCR amplification maps of the FOX2 gene. F: forward primer, R: reverse primer, numbers after F/R means pair number, green bars indicate sequenced region, pink arrows indicate primers used during cycle sequencing.

Fig. 5. (a) Blast result showing 8-base pair deletion, (b) Chromatogram of patient’s DNA, yellow arrow indicates site of mutation. (c) Genomic locations [1] and the relative position of primer pairs [2], locations of FOXC2 coding region [3], sequence gap [4] and identified mutation [5]; genomic locations are given according to GRCh38.
Conclusion
This is the first reported case of LD in Bangladesh. Although Hereditary lymphedemas are rare, analysis and investigation of such disease bear significance. The patient diagnosed in this study draws special attention due to the relatively low expressivity of the disease even though he is homozygous for the deletion in FOXC2. The deletion disrupts codons after 304. It means that the C-terminal activation domain and inhibitory domain are disrupted but the amino acid sequence for the N-terminal activation domain, nuclear localizing segment, and forkhead DNA binding domain remains identical to the native protein. As the homozygous deletion did not result in serious health complications for the patient, the mutated protein probably retains most of its native functions. Although the proper function of the protein might depend on the interactions among its domains, the function of the N terminal activation domain and forkhead DNA binding domain was not fully disrupted otherwise no native function would have been retained. A study found eight proline-directed Ser/Thr phosphorylation sites in FOXC2 clustered in a relatively short region of the protein encompassing amino acids 219 to 367. Phosphorylation at these sites was found to be important for FOXC2-induced vascular remodeling in vivo. It indicates that this eight-base-pair deletion probably hampers the proper regulation of the protein, but its major functional domains maintain their functions.
Unfortunately, the sequence of a 40 bp segment could not be clearly read from the chromatograms due to the limitation of Sanger sequencing to produce peaks of good resolution at the ends of the DNA molecule (Fig. 5[c]). Extensive literature search has ensured that there is no report of mutation in this region and no LD patients were found to simultaneously carry two different mutations of FOXC2 gene until now.
Declarations
Ethics approval and consent to participate: Provided by patient.

Consent for publication: Attached
Availability of data and materials: All data are included in this manuscript.
Competing interests
The authors declare no competing interests
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Authors' contributions: The clinical assessments were conducted by Md. Lutfar Rahman, Md. Mostafizur Rahman, Mohammad Ali and A.F.M Helal Uddin. The molecular study was planned and supervised by Dr. Mustak Ibn Ayub and bench work was conducted by Nahid Parvez.
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References
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