Research Article

Identification of Novel Mutations in ABCA4 Gene: Clinical and Genetic Analysis of Indian Patients with Stargardt Disease

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Received 26 December 2014; Revised 13 February 2015; Accepted 9 March 2015

Academic Editor: Mitsuru Nakazawa

Stargardt disease (STGD) is the leading cause of juvenile macular degeneration associated with progressive central vision loss, photophobia, and colour vision abnormalities. In this study, we have described the clinical and genetic features of Stargardt patients from an Indian cohort. The next generation sequencing was carried out in five clinically confirmed unrelated patients and their family members using a gene panel comprising 184 retinal specific genes. Sequencing results were analyzed by read mapping and variant calling in genes of interest, followed by their verification and interpretation. Genetic analysis revealed ABCA4 mutations in all of the five unrelated patients. Among these, four patients were found with compound heterozygous mutations and another one had homozygous mutation. All the affected individuals showed signs and symptoms consistent with the disease phenotype. We report two novel ABCA4 mutations in Indian patients with STGD disease, which expands the existing spectrum of disease-causing variants and the understanding of phenotypic and genotypic correlations. Screening for causative mutations in patients with STGD using panel of targeted gene sequencing by NGS would be a cost effective tool, might be helpful in confirming the precise diagnosis, and contributes towards the genetic counselling of asymptomatic carriers and isolated patients.

1. Introduction

Stargardt disease (STGD: OMIM #248200/#600110) is an inherited genetic eye disease in which patients develop bilateral macular dystrophy leading to progressive loss of central vision in early childhood. It is the most common form of autosomal recessive juvenile macular dystrophy with a reported prevalence of 1:10000 [1, 2]. The disease is characterized by loss of central vision, fundus flavimaculatus, mottling or atrophy of the retinal pigment epithelium (RPE), bull’s eye maculopathy, flecks in the macula, beaten-bronze macular appearance, and cone-rod dysfunction [3]. STGD is associated with accumulation of lipofuscin content in RPE cells, failure in removal of toxic substances, and significant photoreceptor cell death [4–6]. So far, mutations have been reported in six candidate genes (ABCA4, ELOVL4, PROM1, PRPH2, and CRB1) in various forms of panretinal dystrophies with the possible phenotype of STGD1 [7–12]. Mutations in ABCA4 gene are implicated in recessive STGD type I [13, 14]. ABCA4 is a large gene which consists of 50 exons located in chromosome 1p13 [15]. It is a member of the subfamily A of the ATP-binding cassette (ABC) transporters that is expressed in the retinal outer segments of cone and rod photoreceptor cells. This gene is involved in the transport and clearance of all-transretinal aldehyde, a by-product of the retinoid cycle of vision, and other essential molecules across the disc membrane into the cytoplasm [16–18]. Defective ABCA4 leads to an accumulation of retinoids in the outer
segment or the retinal pigment epithelium and nonhomogenous slowing of retinoid cycle kinetics, as observed in earlier studies in STGD patients [13]. In the animal model, ABCA4-knockout mice show deposition of lipofuscin-like substance in the RPE. Additionally, increased levels of lipofuscin components such as fluorescent dienitroind, A2E, and dirental pyridinium are also observed [19, 20]. Interestingly, knockout of RDH8 (retinol dehydrogenase 8) which conducts all-transretinaldehyde metabolism together with ABCA4 showed the retinal degeneration in mice [21]. Moreover, aged ABCA4 knockout albino mice display a mild retinal degeneration [22]. It is well known that ABCA4 mutations are heterogeneous and cause various forms of retinal dystrophies, which include STGD, retinitis pigmentosa, age-related macular degeneration (AMD), and cone-rod dystrophy [12, 23–26]. In this study, we describe novel and reported ABCA4 gene variants and associated phenotypes in Indian patients with STGD.

2. Methods

2.1. Study Subjects and Clinical Ascertainment. Five clinically confirmed unrelated patients with Stargardt disease and their available family members were enrolled for genetic analysis. Study subjects underwent a complete ophthalmic examination including measurement of visual acuity, intraocular pressure, slit lamp biomicroscopy, detailed fundus examination, fundus photography, fundus autofluorescence (FAF), spectral domain optical coherence tomography (SD-OCT), and full field electroretinography (ERG). Fifty normal controls without any history of eye diseases were included. This study adhered to the tenets of Declaration of Helsinki and was approved by the institutional ethical committee (IEC). Informed consent was obtained from all subjects for the study and publication of their data.

2.2. DNA Extraction. Five mL of peripheral blood samples was obtained from all subjects in EDTA coated vacutainers. Total genomic DNA was extracted using the salt precipitation method described previously [27]. Agarose gel electrophoresis was done to examine the integrity of genomic DNA.

2.3. Sample Preparation. Fifty nanograms of Qubit quantified DNA was used for library preparation using the Illumina Nextera protocol as per the manufacturer’s instructions. This protocol uses transposon-based shearing of genomic DNA and allows the DNA to be “tagmented,” that is, fragmented and tagged simultaneously in the same tube. Limited cycle PCR was carried out to incorporate adaptors and sample specific barcodes to prepare the sample library. The tagged and amplified sample library was checked for quality using BioAnalyzer (Agilent, USA) and quantified.

2.4. Target Enrichment and Exome Sequencing. Approximately 500 nanograms of library was used for enrichment, involving two successive hybridization steps with target specific biotinylated probes. These probes were targeted at the exons of 184 genes with previously known pathogenic variations associated with multiple eye disorders. Bound DNA was pulled down using streptavidin beads. The target library was amplified using limited cycles of PCR. 6–10 pM of the enriched library was loaded for sequencing on to a MiSeq sequencer (Illumina, USA) using a standard v2 kit.

2.5. Data Analysis

Alignment. The trimmed fastq files were generated using MiSeq reporter from Illumina. The reads were aligned against the whole genome hg19 build using Strand NGS v1.6 (Strand Life Sciences) after trimming at least 1 bp at the 3’ end and any others with quality below 10. Reads which had length less than 25 bp after trimming were not considered for alignment. A maximum of 5 matches of alignment score of at least 90% were computed. The reads were realigned using the local realignment tool in Strand NGS. Reads that failed vendor QC, reads with average quality less than 20, reads with ambiguous characters, and translocated and single-mate flip reads were all filtered out.

2.6. Variant Calling. The Strand NGS variant caller was used to detect variants at locations in the target regions covered by a minimum of 10 reads with at least 2 variants reads. Variants with a decibel score of at least 50 were reported. Results were further compared with the dbSNP138 to identify novel variations. The potential deleterious effect of variants was determined by using various in silico prediction algorithms (PolyPhen, SIFT, Mutation Taster. PhyloP, LRT GERP++RS).

2.7. Variant Interpretation. Variants were then imported into Strandomics v1.0 (Strand Life Sciences) for annotation, prioritization, and reporting based on ACMG guidelines. Six known genes (ABCA4, CNGB3, CRB1, ELOVL4, PROM1, and PRPH2) for STGD were included for interpretation.

2.8. Validation of ABCA4 Mutations. The potential ABCA4 variations were revalidated in all the affected individuals and in the available unaffected family members as well as 50 unrelated normal controls using polymerase chain reaction (PCR) and Sanger sequencing. ABCA4 coding exons (5, 17, 19, 35, 42, 46, and 47) and their flanking splice junctions were PCR amplified using the primers reported previously [28]. PCR amplicons were sequenced on a 3730xl DNA Sequencing Analyzer (Life Technologies). Sequencing results were analyzed in FinchTV software (Geospiza, Seattle, WA, USA) and compared with the reference databases (NM_000350, ENSEMBL, and ENST00000370225).

3. Results

Genetic analysis was performed on a group of five unrelated patients with clinical findings of STGD (Figure 1). In this study, we performed NGS analysis in a panel of 184 genes encompassing multiple eye disorders including Stargardt disease. However the data analysis was performed for six known Stargardt related genes (ABCA4, ELOVL4, CNGB3, PROM1, PRPH2, and CRB1). This genetic analysis revealed a total of
five mutations in the \textit{ABCA4} gene. One of these was a previously reported homozygous mutation (p.Arg2149X) with two asymptomatic heterozygous carriers in a five-generation consanguineous family. Two novel heterozygous mutations, p.Phe191Valfs42 and p.Tyr872X, were identified in two different patients along with previously reported heterozygous mutations (p.Gly172Ser, p.Gly1961Glu, resp.), resulting in a compound heterozygous state (p.Phe191Valfs42/p.Gly172Ser and p.Tyr872X/p.Gly1961Glu). In addition, previously reported compound heterozygous mutations (p.Arg1640Trp/p.Gly1961Glu) and a previously reported homozygous mutation (p.Thr971Asn) were found in two isolated cases. There are no pathogenic mutations that were identified in other candidate genes associated with STGD. The clinical features of the patients are described in Table 1. We also found nonpathogenic reported polymorphisms (SNPs) in the Stargardt related genes (\textit{ABCA4}, \textit{ELOVL4}, \textit{CNGB3}, \textit{PROM1}, \textit{PRPH2}, and \textit{CRB1}) which are listed in Table 2.

\subsection*{3.1. Family: SG-01}

Patient (II:1) was a 27-year-old male from nonconsanguineous parents with progressive decrease of vision in both eyes (Figure 1(a)). His best corrected visual acuity was 6/60 and 6/36 in the right eye (RE) and left eye (LE), respectively. Anterior segment examination and the intraocular pressure (IOP) in both eyes were normal. Fundus showed an appearance of bull's eye maculopathy in both eyes. There were no visible flecks. Fundus autofluorescence (FAF) showed an area of hypoautofluorescence corresponding to the atrophic macula surrounded by a ring of hyperautofluorescence (Figure 2(a)). The ring of hyperautofluorescence corresponds to the lipofuscin-laden RPE cells while the central hypofluorescent area corresponds to an area where there is complete RPE atrophy. Full field electroretinography of both eyes was normal suggestive of group 1 STGD. SD-OCT of both eyes showed a central area of photoreceptor loss corresponding to the atrophy seen on the fundus and FAF (Figure 2(a)). Genetic analysis in patient (II:1) revealed two previously reported missense mutations, c.4918C>T (p.Arg1640Trp) and c.5882G>A (p.Gly1961Glu), in compound heterozygous state in the \textit{ABCA4} gene. These compound heterozygous missense mutations were not present in 50 unrelated normal controls. The parents (I:1 and I:2) were clinically normal and did not consent to genetic analysis.

\subsection*{3.2. Family: SG-02}

Patient (II:1) in the second family was a 31-year-old male with complaints of blurred vision for distance since the age of 20 years. He was the first sibling of nonconsanguineous parents (Figure 1(b)). His BCVA was 6/60, N24 in the RE and 4/60, N36 in the LE, respectively. Fundus examination of both eyes showed a central atrophic lesion with flecks restricted to the macula (Figure 2(b)).
Table 1: Clinical features of STGD patients with ABCA4 mutations.

| Family ID | Age/sex | BCVA | Refraction | Fundus | FAF | SD-OCT | Full field ERG |
|-----------|---------|------|------------|--------|-----|--------|---------------|
|           |         |      | RE | LE | RE | LE | | |
| SG-01 salty 27/M 6/60, N8 6/36, N8 | −2.5 DS | −2.75 DS | Macular atrophy, no flecks | Hypo-AF surrounded by ring of hyper-AF | IS/OS loss | Normal |
| SG-02 salty 31/M 6/60, N34 4/60, N36 | 0 | 0 | Macular atrophy, macular flecks | Hypo-AF surrounded by hyper- and hypo-AF flecks | IS/OS loss | Normal |
| SG-03 salty 22/M 6/60, N18 6/45, N18 | −1.25 DS/−1.00 DC 90° | −1.50 DS/−0.50 DC 20° | Macular atrophy, extensive flecks | Hypo-AF surrounded by hyper- and hypo-AF flecks | IS/OS loss | Rod-cone dysfunction |
| SG-04 salty 26/M 6/38, N12 6/38, N12 | −1.00 DS/−0.5 DC 70° | −1.50 DS | Macular atrophy, no flecks | NA | IS/OS loss | Normal |
| SG-05 salty 16/F 6/60, N12 6/38, N12 | +0.5 DS/−0.75 DC 5° | +0.5 DS/−0.75 DC 170° | Macular atrophy, temporal pallor of optic discs | Hypo-AF surrounded by hyper-AF flecks | IS/OS loss | Cone-rod dysfunction |

BCVA: best corrected visual acuity, RE: right eye, LE: left eye, IOP: intraocular pressure, PR: photoreceptors, AF: autofluorescent, SD-OCT: spectral domain optical coherence tomography, FAF: fundus autofluorescence, and ERG: electroretinography.

3.3. Family: SG-03. The patient (II:1) in the third family was a 22-year-old male who presented with difficulties in night vision and blurred vision for distance since the age of 10. He was the only child of nonconsanguineous parents (Figure 1(c)). His BCVA was 6/60, N18 in the RE and 6/45, N18 in the LE, respectively. Slit lamp examination of the anterior segment revealed no abnormalities in both eyes with normal IOP. Fundus examination of both eyes showed an atrophic macula with retinal flecks (Figure 2(c)). FAF showed a central area of hypoautofluorescence corresponding to the atrophic macula and areas of multiple hyperautofluorescence corresponding to the flecks. SD-OCT of both eyes showed foveal thinning with extensive loss of photoreceptors (Figure 2(c)). The full field ERG of both eyes showed rod-cone dysfunction suggestive of group 3 STGD. Genetic analysis of patient (II:1) showed a previously reported homozygous mutation c.2912C>A (p.Thr971Asn) in ABCA4 gene that was not present in 50 unrelated healthy controls analyzed. His parents were normal on ophthalmic examination. We were unable to examine the segregation of this mutation in the parents.

3.4. Family: SG-04. The patient (II:1) in the fourth family was a 26-year-old male with loss of central vision. He was the only child of nonconsanguineous parents (Figure 1(d)). His BCVA was 6/38, N12 in both eyes. Fundus of both eyes showed atrophic macular lesions (Figure 2(d)). Anterior segment examination and the IOP in both eyes were normal. There were no flecks; FAF showed central hypoautofluorescence corresponding to the atrophic macula and areas of multiple hyperautofluorescence corresponding to the flecks. SD-OCT of both eyes showed foveal thinning with extensive loss of photoreceptors (Figure 2(c)). The full field ERG of both eyes showed rod-cone dysfunction suggestive of group 3 STGD. Genetic analysis of patient (II:1) revealed a previously reported novel mutation c.2616C>G (p.Tyr872X) and a previously reported mutation c.5882G>A (p.Gly1961Glu) occurring in a compound heterozygous state in the ABCA4 gene (Figure 3(b)). His unaffected mother (I:1) harbored only one heterozygous variant (p.Gly1961Glu) in...
Table 2: List of nonpathogenic variations identified in patients with STGD by NGS analysis.

| Gene   | Patient ID: SG-01,E1 | SNPID | Variation identified | SNP ID | Variation identified | SNP ID | Variation identified | SNP ID | Variation identified | SNP ID |
|--------|----------------------|-------|----------------------|--------|----------------------|--------|----------------------|--------|----------------------|--------|
| ABCA4  | c.5844A>G            | rs2275029 | c.5844A>G            | rs2417863 | c.5844A>G            | rs4147863 | c.5844A>G            | rs3789452 | c.5844A>G            | rs2297634 |
|        | c.302+26A>G          | rs2297634 | c.302+26A>G          | rs2297634 | c.302+26A>G          | rs2297634 | c.302+26A>G          | rs2297634 | c.302+26A>G          | rs2297634 |
|        | c.6729+21C>T         | rs1800699 | c.6729+21C>T         | rs1800699 | c.6729+21C>T         | rs1800699 | c.6729+21C>T         | rs1800699 | c.6729+21C>T         | rs1800699 |
|        | c.6006-16G>G         | rs4147863 | c.6006-16G>G         | rs4147863 | c.6006-16G>G         | rs4147863 | c.6006-16G>G         | rs4147863 | c.6006-16G>G         | rs4147863 |
|        | c.302+26A>G          | rs2297634 | c.302+26A>G          | rs2297634 | c.302+26A>G          | rs2297634 | c.302+26A>G          | rs2297634 | c.302+26A>G          | rs2297634 |
|        | c.6729+21C>T         | rs1800699 | c.6729+21C>T         | rs1800699 | c.6729+21C>T         | rs1800699 | c.6729+21C>T         | rs1800699 | c.6729+21C>T         | rs1800699 |
|        | c.6006-16G>G         | rs4147863 | c.6006-16G>G         | rs4147863 | c.6006-16G>G         | rs4147863 | c.6006-16G>G         | rs4147863 | c.6006-16G>G         | rs4147863 |
| CRB1   | c.964+475dupT         | rs7756947 | c.964+475dupT         | rs7756947 | c.964+475dupT         | rs7756947 | c.964+475dupT         | rs7756947 | c.964+475dupT         | rs7756947 |
| CNGB3  | c.2264A>G            | rs3735972 | c.2264A>G            | rs3735972 | c.2264A>G            | rs3735972 | c.2264A>G            | rs3735972 | c.2264A>G            | rs3735972 |
| PROM1  | c.2347-4dupC         | rs34269395 | c.2347-4dupC         | rs34269395 | c.2347-4dupC         | rs34269395 | c.2347-4dupC         | rs34269395 | c.2347-4dupC         | rs34269395 |
| PRHP2  | c.13C>T              | rs361524 | c.13C>T              | rs361524 | c.13C>T              | rs361524 | c.13C>T              | rs361524 | c.13C>T              | rs361524 |
| ELOVL4 | c.895A>G             | rs3812153 | c.895A>G             | rs3812153 | c.895A>G             | rs3812153 | c.895A>G             | rs3812153 | c.895A>G             | rs3812153 |

*Nomenclature based on cDNA position according to human genome variation society (HGVS).
ABCA4. His father (I:2) was clinically normal and was not included for genetic testing. These compound heterozygous mutations were not present in the 50 normal controls.

3.5. Family: SG-05. We identified a 12-year-old girl (V:1) from a five-generation Indian family with the characteristic features of STGD (Figure 1(e)). She presented with inability to see the letters on the classroom board since the age of 8 years. She was the first child of a consanguineous marriage. Her BCVA was 6/60, N12 and 6/38, N12 in the RE and LE, respectively, on the Snellen's chart. Slit lamp examination showed normal anterior segment and IOP in both eyes.
Figure 3: Sequencing analysis of two patients with novel *ABCA4* mutations. (a) and (b) show the comparison of sequence chromatograms of two patients (SG-02 II:1, SG-04 II:1) and the normal controls. Arrows indicate the positions of nucleotides change.

were normal. Fundus examination of both eyes showed mild temporal pallor of the optic nerves with an atrophic macula (Figure 2(e)). FAF of both eyes showed an area of hypoautofluorescence corresponding to the atrophic macula with multiple hyperautofluorescent areas corresponding to flecks. SD-OCT of both eyes showed foveal thinning with extensive photoreceptor loss. Full field ERG in both eyes showed a cone-rod dysfunction suggestive of group 2 STGD (Figure 2(e)). The parents and sibling were clinically normal. Genetic analysis of the patient (V:1) showed a previously reported homozygous change c.6445C>T (p.Arg2149X) in the *ABCA4* gene. Her unaffected parents (IV:1 and IV:2) were both heterozygous for the same mutation (p.Arg2149X) showing an asymptomatic carrier status. This mutation was not present in 50 unrelated controls.

4. Discussion

Mutational screening for Stargardt patients has not been described in detail in Indian patients earlier. An appreciation of the genetic basis is important for understanding the biology of the disease. In this study, we report five mutations in the *ABCA4* gene, two of which have not been previously reported, and we determine the clinical phenotype associated with these genotypes in five unrelated patients with STGD. Mutations in *ABCA4* gene are the most common causes of STGD in inherited as well as sporadic cases. Several pathogenic mutations in *ABCA4* have been reported in various ethnic backgrounds of STGD and documented in the HGMD (http://www.biobase-international.com/product/hgmd) and dbSNP databases (http://www.ncbi.nlm.nih.gov/SNP/). Apart from STGD, *ABCA4* mutations have also been reported in other ocular disorders including cone-rod dystrophy [29] and autosomal recessive retinitis pigmentosa [25].

The patient (SG-01 II:1) carries two pathogenic missense mutations p.Gly1961Glu and p.Arg1640Trp, both well-known disease-causing mutations in STGD. p.Gly1961Glu has been reported most frequently in Caucasian and Spanish patients with STGD [30, 31], in 11 probands, who exhibited compound heterozygosity with other missense variations in *ABCA4* [17]. p.Arg1640Trp was identified in a compound heterozygous Stargardt patient who harbored this variant along with p.Gly863Ala and p.Trp1408Arg. *In vitro* functional analysis showed a synergistic effect of the variants (p.Arg1640Trp and p.Trp1408Arg) in causing a severe reduction in *ABCA4* protein expression [32]. The p.Gly1961Glu variation has been associated with a milder STGD phenotype; this is consistent with our findings that the patient had a normal full field ERG with no evidence of either cone or rod dysfunction. The fundus appearance in this patient was similar to the phenotype described previously in individuals carrying the variant p.Arg1640Trp [23].

Another compound heterozygous mutation (p.Phe191Valfs42/p.Gly172Ser) was found in the patient (SG-02 II:1) with typical signs of atrophic macula and paramacular flecks in both eyes. The missense amino acid substitution p.Gly172Ser was previously identified in STGD probands from South African and Italian cohorts [33, 34]. The novel essential splice site mutation c.570+1G>A (p.Phe191Valfs42) is predicted to cause missplicing of the gene, resulting in a frameshift and subsequent truncation in the protein. A neighboring mutation c.570G>C (p.Gln190His) has previously been reported in compound heterozygous state along with p.Gly1961Glu in a patient with STGD [14].

We found a homozygous missense mutation (p.Thr971Asn) in a patient (SG-03 II:1) with a severe phenotype of STGD (atrophic macula, extensive flecks, and rod-cone dysfunction on ERG). We could not check for p.Thr971Asn in the parents, who were possibly asymptomatic.
carriers. This missense change (p.Thr971Asn) has been previously observed in a STGD patient but not in any of the 96 healthy control individuals investigated in that study [35]. The mutation alters a residue within the Walker A ATP binding motif (residues 963–970), adjacent to the NBD1 domain, which is located in a highly conserved region. It has been reported that the NBD1 domain is responsible for basal ATPase activity [36]. Moreover, it has been shown that the variant p.Thr971Asn leads to reduced ATP-binding capacity and abolishes retinal-stimulated ATP hydrolysis [37].

Patient (SG-04, II:1) had a compound heterozygous change (p.Tyr872X, p.Gly1961Glu) and his unaffected mother harbored only one heterozygous variant (p.Gly1961Glu) in the ABCA4 gene. It is well known that p.Gly1961Glu is found in compound heterozygous state along with another pathogenic variant in STGD [17]. The truncating mutation p.Tyr872X has not been previously reported. It is found in the topological domain of the ABCA4 chain and leads to premature truncation after 872 amino acids, which is likely to be nonfunctional and would be degraded or not be translated. Interestingly, nearby nonsense mutations in ABCA4, for example, p.Ser878X, have been identified in a Chinese patient with STGD [38].

Another homozygous mutation (p.Arg2149X) was identified in the young girl (SG-05, V:1) from a consanguineous family with a severe phenotype of STGD. Interestingly, a compound heterozygous combination of p.Arg2149X and another variant IVS45+1G>C has been found in patients who displayed an extensive atrophic appearance of the RPE and abnormal responses from rods and cones [39]. This mutation (p.Arg2149X) leads to a predicted truncated protein in ABC transporter 2 domain in ABCA4 and has been previously described in STGD patients from Japanese and Caucasian origins [40, 41]. There are very few reports describing the variants in CNGB3, PROM1, PRPH2, ELOVL4, and CRB1 that are associated with rare causes of Stargardt or Stargardt-like disease phenotypes [8, 10–12]. Our results suggest that ABCA4 pathogenic mutations might be a major cause in Indian patients with STGD; it has been reported in other ethnic groups as well [31, 33, 34, 40, 42, 43]. In this study we found the most frequent ABCA4 mutation p.G1961E in two unrelated STGD patients; this has been reported in patients from African American and European ancestry with the frequencies of 2% and 11%, respectively [44, 45]. Recently, the different stages of an “optical gap” have been described on the SD-OCT in patients with p.G1961E mutations [46]. Stage 1 was characterized by mild disruption of the ellipsoid zone over the fovea, while stage 2 showed a progressive loss resulting in an apparent “optical gap” on the OCT. Stage 3 showed a collapse of this zone associated with RPE atrophy. In our study cohort, both the patients SG-01, II:1 and SG-04, II:1 who carried a p.G1961E did not demonstrate any optical gap phenotype on the SD-OCT (Figures 2(a) and 2(d)) and exhibited complete loss of photoreceptors and RPE atrophy in the foveal region.

Through our study, we showed the usefulness of next generation sequencing in making the precise molecular diagnosis of STGD and helping in clinical situation. However, a single candidate gene could not be selected for genetic analysis, since STGD is associated with variable clinical features and can be caused by six other known genes. The gold standard for identifying nucleotide changes is direct sequencing, but screening of genes like ABCA4 (large and consists of 50 exons) or other STGD related genes is expensive but time consuming. The panel of targeted gene sequencing by NGS might be a cost effective tool, which enables the rapid sequencing of a large gene like ABCA4 and multiple other genes simultaneously in large cohorts of patients with STGD. This could be a convenient tool for rapid determination of genotype-phenotype correlation in STGD or Stargardt-like diseases and this method could be routinely implemented in future gene therapy approaches in retinal dystrophies. Though we have suggested that performing clinical exome would give us an advantage of validating the clinical diagnosis, in our study cohort we identified nonambiguous STGD phenotype and had ABCA4 gene mutation out of half a dozen other genes. However, this may not be the case when others use the same methodology in their clinics, where there could be clinical and genetic heterogeneity and in such circumstances clinical exome could come in handy for a clinician [47] in further management of the case for focused genetic counseling and also in helping parents to make appropriate reproductive choices.

**Conflict of Interests**

The authors declare no conflict of interests regarding the publication of this paper.

**Authors’ Contribution**

Nallathambi Jeyabal and Arkasubhra Ghosh conceptualized and designed the study, enrolled patients and collected data, and drafted the paper. Rajani Battu enrolled patients and collected clinical data. Anshuman Verma, Ramesh Haritharan, Shuba Krishna, Ravi Kiran, Jemima Jacob, Aparna Ganapathy, and Vedam L. Ramprasad carried out NGS experiment, conducted data analysis, and revised the paper. Govindasamy Kumaramanickavel critically revised the paper. All the authors approved the final version of the paper.

**Acknowledgments**

The authors are grateful to the patients who participated in this study. The authors would like to acknowledge the funding by the Narayana Nethralaya Foundation for Arkasubhra
Ghosh and Nallathambi Jeyabalan and Strand Life Sciences to Rajani Battu for partial support.

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