SHORT REPORT

Biallelic TMEM260 variants cause truncus arteriosus, with or without renal defects

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Abstract
Only two families have been reported with biallelic TMEM260 variants segregating with structural heart defects and renal anomalies syndrome (SHDRA). With a combination of genome, exome sequencing and RNA studies, we identified eight individuals from five families with biallelic TMEM260 variants. Variants included one multi-exon deletion, four nonsense/frameshifts, two splicing changes and one missense change. Together with the published cases, analysis of clinical data revealed ventricular septal defects (12/12), mostly secondary to truncus arteriosus (10/12), elevated creatinine levels (6/12), horse-shoe kidneys (1/12) and renal cysts (1/12) in patients. Three pregnancies were terminated on detection of severe congenital anomalies. Six patients died between the ages of 6 weeks and 5 years. Using a range of stringencies, carrier frequency for SHDRA was estimated at 0.0007–0.007 across ancestries. In conclusion, this study confirms the genetic basis of SHDRA, expands its known mutational spectrum and clarifies its clinical features. We demonstrate that SHDRA is a...
TMEM260 is a 79.5 kDa protein with eight transmembrane spans (www.uniprot.org/uniprot/Q9NX78) located mainly in the nucleoplasm and within focal adhesion sites (www.proteinatlas.org).\(^1\) TMEM260 encodes at least four protein-coding transcripts. Of these, two (ENST00000261556.11 and ENST00000538838.5) are considered to be the main transcripts. They differ in the utilisation of an internal exon as well as the final three exons, which in the short isoform are non-coding.

Five individuals from two families with biallelic truncating TMEM260 variants and brain, cardiac, renal, and digit abnormalities were reported in 2017.\(^2\) The condition is now listed on OMIM as “structural heart defects and renal anomalies syndrome” (SHDRA; MIM #617478). Notably, the variants in both families mapped to the long isoform, raising the possibility of SHDRA being an isoform-specific disorder. Since the original publication, there have been no further reports in the literature. Knowledge about the variant and the clinical spectrum of this condition is therefore limited (Supplementary background). In this study, we describe eight affected individuals from five families, confirming that biallelic TMEM260 loss of function variants cause SHDRA and helping to define its clinical spectrum.

2 | MATERIALS AND METHODS

Whole genome sequencing in Families 1 and 2 was performed as part of the 100 000 genomes project (100KGP; https://doi.org/10.6084/m9.figshare.4530893.v6, Cambridge South REC: 14/EE/1112). Families 3–5 were identified via whole exome sequencing (WES) pipelines and international collaboration. RNA analysis was performed for Family 1 using PaxGene blood samples. Carrier frequency for SHDRA was calculated as described previously.\(^3\) More details are in Supplemental methods and Tables S1-S2.

3 | RESULTS

3.1 | Compound-heterozygous TMEM260 variants in foetuses with congenital heart anomalies

In F1-II-3 (Figure 1A), type I truncus arteriosus (TA) with pulmonary stenosis and ventricular septal defect (VSD) were detected on ante-natal anomaly scan at 20 weeks gestation (Table 1). The pregnancy was terminated at 21 weeks. Post-mortem examination confirmed the cardiac anomalies (Figure S1) and did not reveal any other abnormalities. In F1-II-4 a large peri-membranous outlet VSD, type I TA with small pulmonary trunk and small pulmonary artery branches were detected antenatally and the pregnancy was terminated at 24 weeks gestation. Post-mortem examination confirmed the cardiac anomalies and revealed a horseshoe kidney. The placenta was also abnormal with a two-vessel cord and omphalomesenteric duct remnant.

Trio WGS was performed as part of the 100KGP on F1-II-4 and both parents. Although the initial analysis focussing on several panels from PanelApp was negative, a scan for Mendelian inconsistencies highlighted an apparently homozygous NM_017799.4:c.344G > A:p.(Arg115Lys) TMEM260 variant in F1-II-4 (Figure 1B). As expected, the father (F1-I-1) was heterozygous for the c.344G > A TMEM260 variant, but the mother (F1-I-2) was apparently homozygous for the wild-type allele. Review of read alignments revealed a maternally inherited 4891 bp deletion (Figure 1B) encompassing exons 2–3. Sanger sequencing and digital droplet PCR confirmed that F1-II-3 was also hemizygous for c.344G > A. Neither of the unaffected brothers (F1-II-1 and F1-II-2) had inherited both TMEM260 variants.

Although the c.344G > A variant was initially annotated as p.Arg115Lys, it involves the last base of exon 3 and results in a drop in the predicted splicing efficiency (MaxEntScan: 9.65 → 2.69). We, therefore, performed RT-PCR on peripheral blood sample from F1-I-1, which showed the presence of two bands, with only the larger band seen in controls (Figure 1C). Sanger sequencing confirmed exon 3 skipping (Figure 1D, Figure S2), resulting in a frameshift of exon 4 (p.Val65AlafsTer32). Similarly, the expected exon 1–4 junction was detected by RT-PCR in the maternal sample (Figure S3), resulting in p.(Glu55PhefsTer20). Collectively, the genetic studies, RNA analysis and similarity of the foetuses’ phenotypes with features reported previously,\(^2\) strongly suggest these TMEM260 variants are disease-causing.

3.2 | Identification of additional SHDRA patients

To expand the cohort of patients with SHDRA we interrogated the 100KGP database further and identified a homozygous c.1410C > G: p.Tyr470Ter TMEM260 variant in Family 2 (Figure 1E). F2-II-2 exhibited a common arterial trunk, tricuspid atresia, VSD, partial anomalous pulmonary venous connection, bilateral hearing loss, global developmental delay, protein losing enteropathy and deteriorating renal function from the age of 15 months (Figure S4). Multi-organ severe condition associated with substantial mortality in early childhood and characterised by congenital cardiac malformations with a variable renal phenotype.

KEYWORDS

exome sequencing, genome sequencing, kidney, phenotypic variability, renal failure, SHDRA, structural heart defects and renal anomalies syndrome, TMEM260, truncus arteriosus
| Family number | Family 1 | Family 2 | Family 3 | Family 4 | Family 5 | Ta-Shma et al. 2017* | Totals |
|---------------|---------|---------|---------|---------|---------|-------------------|-------|
| Ethnicity     | White British | Pakistani | Ashkenazi Jewish | Sudanese | Chinese | Ashkenazi Jewish, Arabic |       |
| Parental consanguinity | No | Yes | No | No (no ROH >10 Mb detected) | No | 2/2 Families | 3/7 Families |
| Genomic coordinates (GRCh38) | chr14:56585912G > A, chr14:56625393C > G | chr14:56621697C > T, chr14:56633308GT > G | chr14:56633141GTAT C > G | chr14:56658579A > G, chr14:56634918G > C | chr14:56621697C > T, chr14:56633141GTAT C > G |       |
| cDNA coordinates | c.44A > G, A30del | c.1410C > G | c.193-2A > G, c.1744G > C | c.1393C > T, c.1698_1701delCTAT | c.1393C > T, c.1698_1701delCTAT |       |
| Protein coordinates | p.Tyr470Ter | p.Tyr567Terfs*27 | p.Tyr567Terfs*27 | p.Tyr567Terfs*27 | p.Tyr567Terfs*27 |       |
| CADD (splice predictions) | 34 | 41 (SpliceAI = 0.06), 32 | NA | 34 | 41 (SpliceAI = 0.06), NA | 28.3–41 |
| Isoforms involved | Short and long | Long | Long | Long | Short and long, long | Long |       |
| gnomAD AF (v.2.1.1) | 2/249708: NA | 2/250342 | 19/273758, 2/251170 | Absent | Absent | Absent | Allide counts of 0–19 |
| Individual ID | F1-II-3 | F1-II-4 | F2-II-2 | F3-II-2 | F3-II-3 | F4-II-2 | F5-II-1 | F5-II-3 | NA |       |
| Gender | Female | Male | Male | Male | Male | Female | Female | 2 M, 2F | 7 M, 5 F |       |
| Methods | Sanger sequencing | Trio genome sequencing (HiSeqX) as part of 100KGP | Trio genome sequencing (HiSeqX) as part of 100KGP | Exome + Sanger | Exome + Sanger | Exome + Sanger | Nextera exome + Sanger | Nextera exome + Sanger | Exome sequencing (Agilent Sureselect) | Sanger, exome and genome sequencing |
| Deceased | TOP at 21 weeks | TOP at 24 weeks | Died at 5 years | No (currently 5 years old) | Died at 3 months | No (currently 3 years old) | Died at 4 months | TOP at 22 weeks | 3/4 are deceased (6 weeks, 2 months, 1 year) | 21 week TOP to 5 years |       |
| Cardiac defects | Septal defects(s) | VSD | VSD (subarterial) | VSD | VSD | VSD | VSD and ASD | VSD |       |
| Truncus arteriosus | + (type I) | + (type I) | + (type I, l.s/p complete repair) | + (type I, complete repair at 1 week of age) | + (type I, complete repair at 1 week of age) | + (type I, complete repair at 3.5 months of age) | + (type I) | 2/4 | 10/12 |
| Pulmonary atresia | – | – | – | – | – | – | – | 1/4 | 4/12 |
| Right aortic arch | – | – | – | – | – | – | – | 1/4 | 1/12 |
| Interrupted aortic arch | – | – | – | – | – | – | – | 1/4 | 1/12 |
| (Continues) |       |       |       |       |       |       |       |       |       |       |
| TABLE 1 (Continued) |
|---------------------|
| **Cardiac defects** |
| Tricuspid valve atresia | – | – | + | – | – | – | – | – | 1/4 | 2/12 |
| Partial anomalous pulmonary venous return | – | – | + | (Partially anomalous pulmonary venous connection(s)–left pulmonary vein–innominate) | – | – | – | – | – | 1/4 | 2/12 |
| **Neurodevelopmental defects** |
| Agenesis of corpus callosum | – | – | – | – | – | – | – | – | 2/3 |
| Microcephaly | – | – | – | – | – | – | – | – | 1/4 |
| Developmental delay | N/A | N/A | + | – | N/A | – | N/A | N/A |
| **Renal defects** |
| Renal failure | N/A | N/A | – | – | – | – | – | – | N/A |
| Elevated creatinine levels (normal: 0.2–0.9 mg/dl) | N/A | N/A | Elevated from 15 months of age (Figure S4A,B) | – | Elevated (Figure S5) | Elevated from 10 days of age (Figure S6) | N/A | 3/3 elevated (1.3–1.9 mg/dl) | 6/12 |
| Anuria/Oliguria | N/A | N/A | – | + | – | – | – | – | N/A | 2/4 | 3/12 |
| Urine protein | N/A | N/A | – | – | – | – | – | N/A | NA |
| Renal cysts | – | – | – | – | – | – | – | – | N/A | 1/4 | 1/12 |
| Horseshoe kidney | – | + | – | – | – | – | – | – | NA | 1/12 |
| **Limb defects** |
| Polydactyly | – | – | – | – | – | – | – | – | 1/4 |
| Overriding toes | – | – | – | – | – | – | – | – | 1/4 |
| **Other** |
| Other features | Severe and prolonged chylothorax following surgery | None |
| Horseshoe kidney | Vertical midline linear skin defect over chest wall, single umbilical artery, left pre-auricular sinus nevocili, enterocolitis with perforation of transverse colon at day 10 |
| Edema (3/4), cyanosis (2/4), facial dysmorphism (1/4), low set ears (1/4), webbed neck (1/4), hypotonia (1/4), bilateral preauricular skin tags (1/4), facial port wine nevus (1/4), undescended right testis (1/4) |
| **Prenatal findings/polyhydramnios** |
| Two-vessel cord with omphalomesenteric duct remnant, non-transformed maternal deciduous vessels associated with extra-pleural membranes (of uncertain significance) | Single umbilical artery |

Abbreviations: ASD, atrial septal defect; NA, not available; N/A, not applicable; ROH, region of homozygosity; VSD, ventricular septal defect.

*Deletion removes exons 2 and 3.

*Annotated as missense but results in exon skipping and c.193_344del/p.(Val65Alafs*32) in a splicing defect.

*Only molecularly confirmed cases from Ta-Shma et al. 2017 (PMID: 28318500) are included.
failure following cardiac surgery led to death at age 5 (Table S1, Supplementary Case Histories).

Through international collaboration, we uncovered three further families ascertained via WES (Figure 1E,F). This included a novel c.1644del:p.Pro549LeufsTer46 allele in trans with the previously described2 c.1393C > T:p.Gln465Ter in a sib-pair (Family 3). The elder brother presented with type I TA, mild truncal valve insufficiency, a large VSD, with normal renal function (max. Creatinine of 39 μmoL/L) and normal neurodevelopment at the age of 5 years. The younger brother was born with Type I TA, mild to moderate truncal valve insufficiency and a large VSD (Video S1). A few days after cardiac surgery he developed severe chylothorax and progressive renal failure (Figure S5) and died at approaching 4 months from multi-organ failure. In Family 4 we identified previously described2 homozygous c.1698_1701del:p.Tyr567ThrfsTer27 TMEM260 variant in 2 year old boy with Type I TA and VSD, and normal renal function (maximum creatinine level of 37 μmoL/L) and normal neurodevelopment. In Family 5 we identified novel compound-heterozygous TMEM260 variants: c.193-2A > G and c.1744G > C:p.Glu582Gln in a girl who died at the age of 4 months. The missense variant is predicted damaging by SIFT/PolyPhen2. The girl was born with TA type I, hypoplastic right ventricle, small main and branch pulmonary arteries, VSD and atrial septal defect. During her neonatal period, she developed necrotizing enterocolitis with perforation of transverse colon, renal impairment (Figure S6) and died at 4 months. Type I TA and VSD were detected antenatally in the pro-band’s sibling (Figure S7) and the pregnancy was terminated at 22 weeks. The same compound-heterozygous TMEM260 variants were also identified in the foetus.

3.3 | SHDRA carrier frequency is 0.7–7 per 1000 individuals

Next, we estimated the carrier frequency of SHDRA using a range of stringency thresholds (Figure S8A). The most stringent criteria included only variants that would be predicted to result in loss-of-
function (without low-confidence flags) and known ClinVar pathogenic or likely pathogenic variants. The least stringent criteria included all variants with a CADD score over 30, a spliceAI score over 0.8 as well as loss-of-function alleles and ClinVar pathogenic and likely pathogenic alleles. This analysis showed that per-ancestry gene carrier rate (GCR) for TMEM260 ranges between 0.001 and 0.007 for least stringent parameters to 0.0007–0.005 for the most stringent. Only 16/94 predicted deleterious variants using the lowest stringency threshold are missense variants (Figure S8B). The GCR was found to be higher in individuals with African/African–American ancestry and lowest in Finnish ancestry. The higher GCR in the African/African–American population is due to a possible founder variant (p.Lys696ThrfsTer7, rs568247949) which has “Likely pathogenic” status in ClinVar with a single submission (SCV00092576.2).

4 | DISCUSSION

We present eight individuals, from five independent families, with biallelic TMEM260 variants (Figure 1). In combination with clinical data published previously,2 our results suggest congenital cardiac malformations to be the most consistent phenotype of SHDRA. All 12 patients are reported to have VSD and 10/12 had TA (Table 1). In most of these patients, VSD is likely to be secondary to TA. Notably, TA is one of the rarest congenital cardiac anomalies with few known genetic associations in NKX2-5,4 NKX2-6,5 GATA6 as well as TBX1.7 Interestingly, TMEM260, is predicted to be one of 1442 target genes for GATA6 predicted using known transcription factor binding site motifs from the TRANSFAC database.8 The JASPAR database of transcription factor binding sites predicts a GATA6 binding site within intron 5 of TMEM260 although the functionality of this motif is unknown.9

Our results show that the renal phenotype of SHDRA is highly variable. Horseshoe kidney and cysts were noted in one patient each. The renal failure seen in three individuals could be pre-renal injury and acute tubular necrosis secondary to cardiac failure and systemic illness. However, the decline in glomerular filtration prior to the onset of cardiac failure in F2-II-2 suggests the possibility of underlying renal dysfunction. Further studies should address whether the variable renal involvement is secondary to cardiac complications or a primary component of the condition. The intra-familial variability in renal phenotype indicates that this may not be solely due to the precise TMEM260 variant(s) that are involved. A more likely hypothesis is that there is a congenital predisposition to renal failure, leaving the individual vulnerable to a rapid deterioration that can be precipitated by clinical (e.g., cardiac/intestinal) insults.

The combination of congenital heart disease, especially conotruncal defects with renal abnormalities is unusual. Conotruncal abnormalities are seen in 22q11.21 deletion syndrome, in which renal abnormalities, such as hypoplasia or agenesis of the kidney, multicystic dysplasia and vesicoureteral reflux, are thought to occur in over 30% of patients.10 Another dominant disorder with some phenotypic overlap, including TA and hypoplastic kidneys, is Townes-Brocks syndrome (MIM #107480) due to heterozygous mutations in SALL1.11,12 The association of cardiac, cerebral and renal malformations is also reminiscent of ciliopathies, although generally the cardiac features linked to these group of disorders do not include TA.

Antenatal detection of severe congenital malformations led to termination of pregnancy in three cases described here. Out of nine live born pregnancies, six patients died within the age ranges of 6 weeks to 5 years. One of the two individuals whom survived to 5-years old (F2-II-2) had developmental delay and hearing loss. However, due to insufficient numbers it is difficult to confidently associate these features with SHDRA. We note that two other individuals in the present study who survived beyond their first year, were cognitively normal. Although facial dysmorphism was reported in 1/4 of the original cohort, that feature was not replicated here.

This study substantially expands the known mutation spectrum of SHDRA. Including the patients presented here, a total of eight different TMEM260 variants in 12 individuals from seven families have now been identified (Figure 1F). Of these, two variants are stop-gains, two are frameshifts, one is a multi-exon deletion, two disrupt splicing and one is missense. All variants are supported by in silico tools, including CADD scores, which are 28.3–41 (Table 1). The distribution of the variants confirms that variants affecting only the longer isoform are sufficient to cause SHDRA.

We show that the carrier frequency for SHDRA could be up to 1 in 140 in certain populations (Figure S8). This analysis also identified a potential founder variant in the African/African–American population that requires further functional studies to validate its “Likely Pathogenic” status in ClinVar. The c.1698_1701del seen in Family 4 and an Arabic family described previously7 may also represent a founder mutation.

In conclusion, our description of five families with biallelic TMEM260 variants confirms the genetic basis of SHDRA and helps delineate the mutational/phenotypic spectrum of the condition. The strong association with TA has important implications for genetic counselling, prenatal diagnostics as well as postnatal targeted genetic testing.

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CONFLICT OF INTEREST

No conflicting interests are declared.

PEER REVIEW

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DATA AVAILABILITY STATEMENT

Researchers can apply to access 100KGP data at www.genomicsengland.co.uk/join-a-gecip-domain. Other data can be made available upon request.
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SUPPORTING INFORMATION

Additional supporting information may be found in the online version of the article at the publisher’s website.

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APPENDIX A

A.1. | The Genomics England Research Consortium (27th May 2021)

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