A Novel Splice-Site Mutation in the ELN Gene Suggests an Alternative Mechanism for Vascular Elastinopathies

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Abstract: The ELN gene encodes elastin, a fundamental protein of the extracellular matrix that confers elasticity to different tissues including blood vessels. The formation of elastin fibers is a complex process involving monomer coacervation and subsequent crosslinking. Mutations in exons 1–29 of the ELN gene have been linked to supravalvular aortic stenosis (SVAS) whereas mutations in exons 30–33 are associated with autosomal dominant cutis laxa (ADCL). This striking segregation has led to the hypothesis that distinct molecular mechanisms underlie both diseases. SVAS is believed to arise through haploinsufficiency while ADCL is hypothesized to be caused by a dominant negative effect. Here, we describe a patient with SVAS harboring a novel splice-site mutation in the last exon of ELN. The location of this mutation is not consistent with current knowledge of SVAS, since all mutations reported in the C-terminus have been found in ADCL patients, and a thorough evaluation did not reveal significant skin involvement in this case. RT-PCR analysis of skin tissue showed that C-terminal mutations in the region can lead to the production of aberrant transcripts through intron retention and activation of cryptic splice sites and suggest that disruption of the very last exon can lead to functional haploinsufficiency potentially related to SVAS.

Keywords: SVAS, ELN, ADCL, coacervation, splicing

Introduction

Elastin is a key protein of the extracellular matrix present in connective tissue. This protein is one of two components of elastic fibers, which confer elasticity to organs and tissues including the heart, skin, lungs, and blood vessels.1 The formation of elastic fibers, or elastogenesis, is a stepwise process whereby elastin monomers bind to receptors on the membrane of fibroblasts and are subjected to coacervation and subsequent cross-linking.2–4 Elastogenesis is important during embryogenesis and shortly after birth. Mutations in the elastin gene are associated with diverse forms of elastinopathies, including supravalvular aortic stenosis (SVAS), autosomal dominant cutis laxa (ADCL) and pulmonic stenosis.5,11

Supravalvular aortic stenosis (SVAS) (OMIM 185500) is a congenital narrowing of the lumen of the ascending aorta with an estimated incidence of 1 in 20,000 live births.6,12 This condition can present in both syndromic and nonsyndromic forms. The syndromic form is usually associated with Williams-Beuren syndrome (WBS) (OMIM 194050), which shows an autosomal dominant inheritance pattern and is caused by a deletion in 7q11.23. This syndrome is characterized by intellectual
disability, distinctive facial features, hypercalcemia, connective tissue abnormalities, growth abnormalities and a gregarious personality.\textsuperscript{13} Mutations in the elastin gene (\textit{ELN}) are associated with non-syndromic SVAS with compromised coronary and pulmonary artery branches, but normal descending aorta, cerebral and renal arteries.\textsuperscript{12} \textit{ELN} arteriopathy is inherited in an autosomal dominant pattern, with incomplete penetrance and variable expressivity.\textsuperscript{6}

Autosomal Dominant Cutis Laxa (ADCL) is an inherited connective tissue disease characterized by an excess of skin tissue with loss of elasticity and inguinal hernia, SVAS, pulmonary stenosis and emphysema in some cases.\textsuperscript{14} Histological findings in this condition show fiber fragmentation with reduced deposition, whereas in SVAS only reduced total deposition of elastin fibers without additional abnormalities can be observed.\textsuperscript{1}

Even though both conditions are caused by mutations in the same gene, SVAS and ADCL are mutually exclusive in most patients, suggesting distinct underlying molecular mechanisms. Animal and in-vitro studies suggest that non-syndromic SVAS mutations lead to insufficient levels of elastin, resulting from lower mRNA levels because of degradation via the NMD pathway.\textsuperscript{10} Similarly, in-vitro studies show that ADCL mutations lead to the expression of dominant negative proteins which interfere with its function.\textsuperscript{12} Remarkably, mutations associated with one or the other show distinct clusters within the \textit{ELN} gene. Analysis of ClinVar and HMGD databases showed that all frameshift (n=42) and intronic (n=14) mutations associated with SVAS are located in exons 1–29 and all frameshift (n=17) mutations associated with ADCL are found in exons 30–33 (Figure 1A).\textsuperscript{1}

In this report, we present a patient with a mutation in the intron-exon boundary of the last exon of the \textit{ELN} gene

![Diagram](https://www.dovepress.com/)

\textbf{Figure 1} Mutations on \textit{ELN} gene associated to SVAS. (A) Schematic representation of the \textit{ELN} gene and the mutations associated with SVAS and Cutis Laxa showing the location of the reported mutations (B) Family pedigree. Reported case and different phenotypes in the family.

\textbf{Abbreviation:} SVAS, supravalvular aortic stenosis.
that leads to the production of aberrant transcripts. However, unlike all reported frameshift mutations in the C-terminus of ELN, this mutation causes SVAS and not ADCL. This finding suggests that the distinction between haploinsufficiency and dominant negative mechanisms in the pathophysiology of SVAS and ADCL is not as clear-cut as previously thought. Alternatively, this mutation suggests that there could be an additional mechanism for SVAS beyond insufficient levels of ELN.

Patient Report

The proband was born at 40w2d of gestation after an uneventful pregnancy with birth weight of 2940 g and birth length of 49 cm. When he was delivered, his mother was 23 years old and his father was 28 years old. At 4 years of age, the patient presented two syncopal episodes. In the study of these episodes, echocardiography revealed aortic valvular stenosis with a valvular gradient of 60mmHg, asymmetric thickened tri-leaflet valve and hypertrophy of the left ventricle. Balloon valvuloplasty was performed successfully. At the age of 7, a new echocardiogram and contrast cardiac computed tomography were performed and showed SVAS of 33% of the lumen with gradient of 45mmHg without coronary arteries compromise. Later, cardiac catheterization was performed, revealing evidence of the SVAS, but renal and pulmonary circulation was normal. Thus, the aorta was repaired with an autologous pericardial patch. At the age of nine, a skin biopsy was performed, which showed no significant alterations (Figure 2). Physical examination was unremarkable; the proband has neither dysmorphic characteristics nor hypermobility. The parents had another male son which died at the age of 11 months with supravalvular aortic stenosis. Additionally, the proband’s paternal half-sister died at age 3 with asymptomatic pulmonary stenosis and epilepsy (Figure 1B).

Materials and Methods

Whole Exome Sequencing

Total DNA from the patient was extracted from a blood sample with a QIAamp DNA mini kit (QIAGEN, Germantown, USA) according to the manufacturer’s instructions. DNA concentration was measured using a Qubit DNA assay Kit in a Qubit 2.0 Fluorometer (Life Technologies, CA, USA). A total amount of 1.0 µg genomic DNA was used for Whole Exome Sequencing (WES) as input material for library preparation. Sequencing libraries were generated using Agilent Sure Select Human All Exon Kit v6 (Agilent Technologies, CA, USA), after generation of blunt-ended 180–280 bp fragments by hydrodynamic shearing and enzyme treatment. Then, adapter oligonucleotides were ligated after adenylation at the 3’ DNA fragments with ligated adapter molecules on both ends were selectively enriched in a PCR reaction. After PCR reaction, libraries were hybridize in liquid phase with biotin-labeled probes, and captured with streptavidin-linked magnetic beads. Products were purified using the AMPure XP system (Beckman Coulter, Beverly, USA) and quantified on the Agilent Bioanalyzer 2100 system. 150 bp paired ends were sequenced on the HiSeq Illumina platform. The raw reads were aligned to the hg19 reference genome with Burrows-Wheeler Aligner (BWA) software. The variants were called with GATK. >80% of bases have a sequencing quality score >Q30 and minimum coverage of 100X. Following genomic variant detection, variants were annotated using ANNOVAR to show affected genomic regions, protein coding changes, and allele frequency reported by Genome Aggregation Database (gnomAD). Library preparation and sequencing were carried out by Novogene Inc. (Beijing-China).

The VarSeq v2.2 (Golden Helix) software was used to filter variants in the ELN gene. These variants were classified into missense, nonsense, splice site and frameshift. For our analysis, novel variants, or variants with MAF≤0.005 were selected.

Skin Biopsy and Histology

A skin sample was collected from the scapular region by punch biopsy. Half of the sample was refrigerated in trizol and the rest was fixed with formaldehyde for hematoxylin and eosin staining and histologic analysis.
RNA Extraction, RT-PCR and Cloning

To determine the effect of the splicing mutation, we performed RNA extraction using Trizol Reagent and the PureLink RNA Mini Kit (Thermo Scientific). We generated cDNA using the SuperScript™ III First-Strand Synthesis System following the manufacturer’s recommendations.

We carried out two PCR approaches to detect all possible aberrant transcripts: a first one to generate a fragment containing exon 32 and the 3’UTR (primers F3 and R3), and a second one to generate a fragment spanning exon 30 and intron 31 (primers F1 and intR1) (Figures 3A and 4A). PCR primers were generated using Primer Blast (https://www.ncbi.nlm.nih.gov/tools/primer-blast/). The target regions of ELN were amplified by PCR and primer sequences are available upon request.

For cloning, 1µL of PCR products were cloned into pCR™4-TOPO™ Vector (Invitrogen) using TOPO™ TA Cloning™, as the manufacturer’s instructions. Ligated plasmids were inserted in One Shot TOP10 E. coli (Invitrogen) through thermic shock during 30 sec. The bacteria were incubated during 1h in SOC medium. Transformed cells were plated on LB agar plates containing 100 µg/mL ampicillin and were grown at 37°C overnight. Plasmid DNA was extracted from colonies with a Qiagen Plasmid Mini kit (Qiagen).

Sanger Confirmation

We confirmed the mutation identified by WES and the cloned RT-PCR products using sanger sequencing. The genomic DNA encompassing the variant ELN-c.2132–14_2161del was amplified by PCR using primers designed with the same tools mentioned previously (Primers available upon request). The pCR™4-TOPO™ Vector plasmids containing the PCR products of RT-PCR were sequenced with T7 and T3 primers. The obtained sequences were compared to reference sequence ENST00000252034.12 (NM_000501.3).

A detailed protocol is available at: dx.doi.org/10.17504/protocols.io.bmcnk2ve

Results

Germline sequencing of genomic DNA from the patient revealed a new splice site mutation in the ELN gene, which corresponds to a deletion of 44 bp across intron 32/exon 33 boundary (ELN-c.2132–14_2161del). This mutation has not been previously described in public databases (gnomAD) nor in the literature.

The functional effect of the mutation was determined by amplifying putative aberrant transcripts resulting from intron retention and activation of cryptic splice sites using intronic and 3’UTR primers (Figures 3A and 4A). Amplicons resulting from RT-PCR with the 3’UTR primer (F3 and R3) revealed a band smaller than expected for the wild type allele (WT), and another that was consistent with a normal allele (Figure 3B). Cloning and sequencing analysis of these PCR products identified an aberrant transcript that skipped all the coding region of exon 33 and used a new cryptic splice site in the 3’UTR (Figure 3C). This alteration, combined with a common insertion (rs34208922) in the 3’UTR, led to a predicted C-terminus containing 49 additional residues and lacking the conserved KxxxRKKR motif (r=[., 2132_2175del]; p. G712Pex*44). (Protein #1, Figure S1).

Amplicons resulting from RT-PCR using the intronic primer (F1 and IntR1) showed two abnormal bands of 281bp and 227bp (Figure 4B). Sequencing of the 281 bp product revealed the presence of 92 bp of intrinsic sequence adjacent to exon 32 indicating intron 32 retention. In this case, the disruption of the splice acceptor site of intron 32 inhibits splicing and results in retention of part of intron 32 in the aberrant transcript (Figure 4C). The aberrant protein has 17 additional amino acids at the C-terminus and disrupts the KxxxRKKR motif (protein #2) (Figure S1). The 227 bp product shows an identical intron 32 retention, combined with the alternative splicing of exon 31, also observed in WT transcripts. The protein generated (protein #3) is likely to have the same functional implications as described for protein #2 (Figure S2).

Histological analysis of the skin biopsy revealed normal elastic fibers in configuration and number with adequate cellularity and without inflammatory infiltrates (Figure 2).

Discussion

SVAS is a rare disease caused by mutations in the ELN gene. The molecular mechanism proposed for the pathogenesis of SVAS is ELN haploinsufficiency, explained by premature termination codon mutations (PTCs), extensively described in affected patients.8,11,12,15 PTCs lead to insufficient expression of ELN because mutant mRNA is presumably degraded by the nonsense mediated decay (NMD) pathway.10 A reduction in ELN expression has also
been found in skin fibroblasts and aortic smooth muscle cells of SVAS affected patients, thus supporting ELN haploinsufficiency as the mechanism involved in the pathogenesis of the vasculopathy.\(^{11}\)

The location of protein truncating mutations within a gene can have a significant influence on the levels of mutant transcripts by affecting the efficiency of nonsense mediated decay. For instance, mutations that occur near
the 3’ end of the dystrophin gene related to Duchenne Muscular Dystrophy (DMD) result in mild phenotypes, suggesting that these truncating proteins are capable of a partial rescue of the DMD phenotype. PTCs near the 5’ end, however, are associated with severe forms of disease and fail to rescue the phenotype because of the NMD pathway. Furthermore, PTCs near the 3’ end of gene are invisible to the NMD machinery.16 Similarly, mutations near the C-terminus of ELN can be less affected by NMD and more likely to lead to the production of dominant negative proteins as observed in ADCL.

This model has been used to explain why all frameshift and nonsense mutations in exons 1–29 lead to SVAS and similar mutations in exons 30–33 lead to ADCL. However, this model does not explain why peptides with dominant negative effects overwhelmingly affect the structure of the extracellular matrix in the skin but not the vasculature and why missense mutations can cause SVAS. Only a few missense mutations have been conclusively associated with this condition, although most seem to affect protein expression by disrupting the initial methionine or affecting splicing. However, the p.Ala707Asp mutation, which seems to affect the putative microassembly domain in exon 29 is particularly interesting, since it can potentially reduce protein expression by affecting post-transcriptional regulation.1 This suggests that most, if not all, ELN variants associated with SVAS result in reduced expression or functional haploinsufficiency.

In this study, we identified a deletion of 44 base pairs in the ELN gene situated across the intron-exon boundary of intron 32 and exon 33 in a patient with SVAS and no clinical or histological skin abnormalities. The patient had a significant family history of severe SVAS, suggesting an autosomal dominant pattern of inheritance with incomplete
penetrance. cDNA analysis indicated the formation of several aberrant transcripts through the activation of cryptic splice sites in the 3’UTR (protein #1) and intron retention (proteins # 2 and 3). Since all of the transcripts generate an aberrant stop codon in the final exon, they are unlikely to trigger NMD. Since NMD is unlikely to play a role in this case of SVAS, the data suggests that an alternative mechanism can lead to SVAS.

The aberrant transcripts produced in this case are unusual because they preserve the amino acid composition of almost all of the elastin protein, with the exception of the 13 C-terminal residues. However, this region contains the highly conserved KxxxRKRK motif, known to mediate the interaction between tropoelastin and the GAGs on the cell membrane. We propose this disruption can lead to defects in the coacervation and/or crosslinking of tropoelastin monomers, impeding their assembly into elastin fibers. This can lead to a functional haploinsufficiency, whereby the mutant tropoelastin is not incorporated into elastin fibers but remains in its soluble form. Functional studies of the elastin fiber should be conducted to determine if these hypotheses are indeed the case.

Alternatively, Schmitz et al recently showed that intron retaining mRNAs have significantly longer 3’UTR sequences that are enriched for miRNA binding sites. miRNA sequences of 18–21 nucleotides are involved in mRNA degradation and/or inhibition of translation. In this case, the retention of intron 32 in transcripts #2 and #3 permits the formation of transcripts with longer 3’UTR sequences, with a greater potential of undergoing miRNA-mediated degradation or translational inhibition. This could lead to a reduced quantity of ELN protein leading to haploinsufficiency via an NMD-independent mechanism.

Although a few other ELN mutations seem to also defy the haploinsufficiency/dominant negative model for SVAS/ADCL, careful analysis can explain why they are behaving the way they do. For example, a splicing mutation associated with ADCL was found in exon 25. However, the aberrant transcript is predicted to produce an inframe deletion of exon 25, likely avoiding the activation of NMD and producing an aberrant protein. This is consistent with the dominant negative model for ADCL. Similarly, a frameshift mutation affecting only the last four residues of ELN (preserving the KxxxRKRK motif) was found in a family with ADCL, suggesting that an unusual feature of the transcripts produced in this case is causing the unexpected phenotype.

**Conclusion**

This study suggests that the functional consequence of extreme C-terminal ELN mutations can have more diverse effects than previously thought and must be analyzed carefully when counselling patients. Furthermore, the mechanisms proposed here can also help explain why a handful of missense mutations can cause SVAS, without likely affecting protein expression. These mutations can affect critical residues necessary for the incorporation of tropoelastin monomers into elastin fibers and can be used to map critical protein–protein interactions. Functional haploinsufficiency can play a role in several other mutations in ELN and other genes which rely on extensive protein–protein interactions for their function.

**Abbreviations**

ADCL, autosomal dominant cutis laxa; cDNA, complementary DNA; DMD, Duchenne Muscular Dystrophy; DNA, deoxyribonucleic acid; EJC, exon junction complex; ELN, elastin; GAG, glycosaminoglycan; HGMD, human gene mutation database; mRNA, messenger ribonucleic acid; NMD, nonsense mediated decay; OMIM, Online Mendelian Inheritance in Man; PTC, premature terminal codon; RT-PCR, reverse transcription-polymerase chain reaction; SVAS, supravalvular aortic stenosis; UTR, untranslated region; WBS, Williams-Beuren syndrome; WT, wild type.

**Ethical Compliance**

Informed consent for genetic testing, skin biopsy and publication of clinical data and images was signed by the proband and guardian and the study protocol was approved by the institutional review board of the Fundacion CardioInfantil as a part of project “Programa para la Innovación en Cardiopatías Congénitas Humanas infrecuentes para Colombia” PINOCCHIO. (Number 21-2016).

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**Author Contributions**

All authors made a significant contribution to the work reported, whether that is in the conception, study design, execution, acquisition of data, analysis and interpretation, or in all these areas; took part in drafting, revising or critically reviewing the article; gave final approval of the version to be published; have agreed on the journal to
which the article has been submitted; and agree to be accountable for all aspects of the work.

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**Disclosure**

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