Craniofrontonasal Syndrome Caused by Introduction of a Novel uATG in the 5′ UTR of EFNB1

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Abstract
Craniofrontonasal syndrome (CFNS) is an X-linked disorder caused by EFNB1 mutations in which females are more severely affected than males. Severe male phenotypes are associated with mosaicism, supporting cellular interference for sex bias in this disease. Although many variants have been found in the coding region of EFNB1, only 2 pathogenic variants have been identified in the same nucleotide in 5′ UTR, disrupting the stop codon of an upstream open reading frame (uORF). uORFs are known to be part of a wide range of post-transcriptional regulation processes, and just recently, their association with human diseases has come to light. In the present study, we analyzed EFNB1 in a female patient with typical features of CFNS. We identified a variant, located at c.–411, creating a new upstream ATG (uATG) in the 5′ UTR of EFNB1, which is predicted to alter an existing uORF. Dual-luciferase reporter assays showed significant reduction in protein translation, but no difference in the mRNA levels. Our study demonstrates, for the first time, the regulatory impact of uATG formation on EFNB1 levels and suggests that this should be the target region in molecular diagnosis of CFNS cases without pathogenic variants in the coding and splice sites regions of EFNB1.

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Keywords
Genetic counseling · Molecular diagnosis and testing · Protein translation · Regulatory variant · Upstream ORF and ATG

Craniofrontonasal syndrome (CFNS; OMIM 304110) is a rare X-linked dominant disorder caused by loss-of-function mutations in EFNB1 [Twigg et al., 2004; Wieland et al., 2004]. Greater severity is observed in heterozygous female patients compared with hemizygous male individuals. Female patients can present with severe hypertelorism, a central nasal groove, craniofacial asymmetry, and coronal craniosynostosis associated with extracranial...
features such as sloping shoulders, mild cutaneous syndactyly, grooved nails, duplication of the first digit, and wiry hair. On the other hand, hemizygous males present only with hypertelorism and occasional cleft lip [Twigg et al., 2004, 2013; van den Elzen et al., 2014]. However, severe phenotypes have been observed in male patients with mosaic EFNB1 mutations [Twigg et al., 2013], a finding that supports the hypothesis that cellular interference underlies clinical severity [Wieacker and Wieland, 2005; Twigg et al., 2013; Niethamer et al., 2017].

More than a hundred EFNB1 mutations have been described in association with CFNS. Almost all pathogenic variants are in the coding region, with the majority (>95%) leading to premature termination codons [Twigg et al., 2004, 2006, 2013; Wieland et al., 2004, 2005, 2007; Shotelersuk et al., 2006; Torii et al., 2007; Wallis et al., 2008; Hogue et al., 2010; Makarov et al., 2010; Grasso et al., 2011; Apostolopoulou et al., 2012; Cui et al., 2012; Ramirez-Garcia et al., 2013; Seven et al., 2013]. Only 2 mutations affecting the same nucleotide have been reported outside the coding and splice site sequences of EFNB1: c.–95T>G and c.–95T>C. These variants are present in the 5′UTR and abolish the stop codon of a small upstream open reading frame (uORF), leading to an overlapped and out-of-frame ORF with the main downstream ORF (dORF, corresponding to EFNB1), with reduced translation of EFNB1 as a consequence [Twigg et al., 2013].

uORFs regulate gene expression through multiple functional mechanisms, e.g., in a peptide-dependent manner, by consumption of functional pre-initiation complex, or triggering nonsense-mediated decay [Rahmani et al., 2009; Barbosa et al., 2013; Hurt et al., 2013; Ebina et al., 2015]. These regulatory sequences are mostly linked to repression of translation of the main ORF, although under specific conditions, uORFs can confer induction of protein translation [Wethmar, 2014]. uORF-altering variants have been described in association with the etiology of human disorders [Barbosa et al., 2013]. Such variants can introduce new uORFs, as found in gonadal dysgenesis and van der Woude syndrome [Poulat et al., 1998; Kondo et al., 2002; Calvo et al., 2009], as well as disrupt uORFs in diseases such as Marie Unna hereditary hypotrichosis [Wen et al., 2009] and thrombocytopenia [Kondo et al., 1998].

In this study, we demonstrate the regulatory impact of a variant-created upstream ATG (uATG) on mRNA translation of EFNB1. Identification of such variants contributes to the expanding field of translation regulation and to the definition of critical noncoding sequences that can be targeted in genetic diagnosis.

Case Report

The female patient (referred to as F5162-1), first evaluated 10 days after birth, presented with typical features of CFNS: unilateral coronal craniosynostosis (left), hypertelorism, asymmetric craniofacial contour, widows peak, arched eyebrows, hypoplastic supraorbital ridges, telecanthus, downsloping palpebral fissures, short nose with depressed nasal bridge, bifid nasal tip, deep philtrum, tented upper lip, widened metopic suture, and a frontal bone defect (Fig. 1a, b). A fronto-orbital advancement surgery was performed at the age of 5 months. No clinical manifestations were observed in her parents, and there is no additional family history.

Materials and Methods

DNA Samples

Genomic DNA from the patient and her clinically unaffected parents was extracted from peripheral blood using Gentra Systems Autopure LS (AutoGen), according to the manufacturer’s instructions.

DNA Sequencing

Standard genetic screening of the coding regions of EFNB1 was carried out with the patient’s DNA using Sanger sequencing. Primers were designed with Primer3 [Untergasser et al., 2012] (primer sequences available upon request). Sequencing reactions were performed using the BigDye Terminator v3.1 Cycle Sequencing Kit, and the amplicons were sequenced using an ABI 3730 DNA Analyzer (Thermo Fisher Scientific). Data were analyzed using Sequencher v.5.1 software.

A more comprehensive analysis of the patient’s DNA was performed by targeted next-generation sequencing. Specific probes were used to capture 5′ and 3′ UTR sequences and all coding regions of EFNB1 (RefSeq NM_004429) as well as splice acceptor and donor sites (Nextera – custom target enrichment, Illumina), using MiSeq (Illumina) according to the manufacturer’s protocols. Data were aligned to the human reference sequence (hg19) with the Burrows-Wheeler Aligner (BWA) [Li and Durbin, 2009], and PCR duplicates were removed with the Picard toolkit. Variants were called by the Unified Genotyper tool from Genome Analysis Toolkit [McKenna et al., 2010] and annotated with ANNOVAR [Wang et al., 2010]. Sanger sequencing was used to analyze the candidate variant in parental DNA (primer sequences available upon request).

Variant Analysis

Variants present in the 1000 Genomes database, dbSNP150, Genome Aggregation database (gnomAD), or in the Online Archive of Brazilian Mutations (ABrOAM) [Naslavsky et al., 2017] were excluded from downstream analysis as unlikely to be pathogenic. Evolutionary sequence conservation was assessed using the Vertebrate Multiz Alignment & Conservation track (100 vertebrate species) and Genomic Evolutionary Rate Profiling (GERP) (35 species of mammals), both through the UCSC Genome Browser. For multiple vertebrate sequence alignment, 20 EFNB1 orthologues were extracted from the Ensembl genome browser along with the human gene. All sequences comprised the first coding exon plus the 5′UTR (when it was found annotated) or ~700 bp of
A Variant-Created uATG in \textit{EFNB1}

the upstream sequence from the main ATG start codon. The se-
quen
ces were aligned using MAFFT \cite{Katoh2017} through
the AliView v.1.18.1 program \cite{Larsson2014}; consensus se-
quen
cese for the alignment was obtained using Jalview v.2.10.2b2 \cite{Water-
house2009}. Open Reading Frame Finder (ORFfinder, NCBI) was utilized
to search for uORFs in the sequences. NetStart 1.0 (http://www.cbs.dtu.
dk/services/NetStart/) was taken to pre-
dict and score translation initiation sites.

\textbf{Microsatellite Analysis}

Microsatellite genotyping was conducted using fluorescently
labeled PCR primer pairs from the Linkage Mapping Set v2.5, ac-
cording to the manufacturer’s protocol. PCR fragments were read
in an ABI 3730 DNA Analyzer (Thermo Fisher Scientific) and an-
alyzed with GeneMapper v5.0.

\textbf{Plasmids}

Site-directed mutagenesis was carried out to introduce spe-
cific nucleotide substitutions within the 5′ UTR of \textit{EFNB1}, using
the QuikChange II Site-Directed Mutagenesis Kit (Agilent Tech-
nologies), following the manufacturer’s instructions. The wild
type (WT) 5′UTR of \textit{EFNB1} was inserted into a previously
modified psiCHECK-2 vector \cite{Calvo2009; referred to
as WT construct} in which the ATG of \textit{Renilla} was modified
to TTG, so that \textit{Renilla} expression was driven from the primary
\textit{EFNB1} ATG codon. We used this vector as a template to gener-
ate a construct containing the c.–411C>G variant (referred
to as F5162 construct) and a second construct containing
c.–227G>T and c.–225G>A in addition to c.–411C>G (referred
to as F5162\textsuperscript{SHORT}; further explanation below). Sanger sequencing
was used to confirm the incorporated modifications (primer se-
quen
ceses available upon request).

For comparative analysis in the luciferase assays, a previously
described construct taken to investigate the c.–95T>G variant
\cite{Twigg2013} was also used (referred to as 1330).

\textbf{Luciferase Assay}

Human embryonic kidney 293T (HEK293T) cells were cul-
tured in Dulbecco’s modified Eagle’s medium (DMEM) supple-
cmented with 10% FBS, 100 U/mL penicillin, and 100 μg/mL strep-
tomycin (all provided by Life Technologies). Cells were seeded at
6 \times 10^3 cells/well in 96-well optical-bottom plates (Nunc\textsuperscript{TM}, Ther-
mo Scientific) 24 h prior to transfection. Transient transfections
were performed using 100 ng of the vector constructs, in triplicate,
with TurboFectin 8.0 (OriGene), according to the manufacturer’s
instructions. Forty-eight hours after DNA transfection, \textit{Renilla} ac-
tivity was measured with the Dual-Glo Luciferase Assay System
(Promega), in a GloMax-Multi Detection System (Promega). All
constructs also contained the Firefly luciferase gene, used as a
transfection control. \textit{Renilla} luminescence results were normal-
ized by Firefly luciferase. Statistical analysis was performed by one-
way ANOVA with a post-hoc Bonferroni multiple comparison test
using the GraphPad Prism 6.0. Data were presented relative to the
WT construct as mean ± SEM.
Real-Time Quantitative PCR

HEK293T cells were seeded at 10^5 cells per well in 6-well culture plates 24 h prior to transfection. Cells were transfected with 1 μg of vector construct, in duplicate, with TurboFectin 8.0. Forty-eight hours after transfection, cells were washed with PBS, and total RNA was obtained using the Nucleospin RNA II extraction kit (Macherey-Nagel), following the manufacturer’s recommendations. cDNA synthesis was performed with the SuperScript IV First-Strand Synthesis System (Life Technologies) and oligo-dT primers using 1.5 μg of total RNA from each cell sample as starting material. RT-qPCR reactions were performed with 2X Fast SYBR Green PCR Master Mix (Applied Biosystems) and 200 nM of each primer. Primer pairs for Renilla luciferase 5′-TCCATTGCTGAG-AGTGTCTGTG-3′ (forward) and 5′-CAAGCACCTTTTCTCGCCC-3′ (reverse) and Firefly luciferase 5′-TCTGGCGACATTGCTCCTACTCTG-3′ (forward) and 5′-CGGCGTCGAAAATGTTAGGG-3′ (reverse) were designed with the Primer-BLAST tool (NCBI) [Ye et al., 2012], and their amplification efficiencies (E) were determined by serial cDNA dilutions; primers were supplied by Exxtend. Fluorescence was detected using the QuantStudio 5 System (Applied Biosystems) under standard temperature protocols. RT-qPCR reactions were performed with 2X Fast SYBR Green PCR Master Mix (Applied Biosystems) and 200 nM of each primer. Primer pairs for Renilla luciferase 5′-TCCATTGCTGAGAGTGTCTGTG-3′ (forward) and 5′-CAAGCACCTTTTCTCGCCC-3′ (reverse) and Firefly luciferase 5′-TCTGGCGACATTGCTCCTACTCTG-3′ (forward) and 5′-CGGCGTCGAAAATGTTAGGG-3′ (reverse) were designed with the Primer-BLAST tool (NCBI) [Ye et al., 2012], and their amplification efficiencies (E) were determined by serial cDNA dilutions; primers were supplied by Exxtend. Fluorescence was detected using the QuantStudio 5 System (Applied Biosystems) under standard temperature protocols. RT-qPCR reactions were performed with 2X Fast SYBR Green PCR Master Mix (Applied Biosystems) and 200 nM of each primer. Primer pairs for Renilla luciferase 5′-TCCATTGCTGAGAGTGTCTGTG-3′ (forward) and 5′-CAAGCACCTTTTCTCGCCC-3′ (reverse) and Firefly luciferase 5′-TCTGGCGACATTGCTCCTACTCTG-3′ (forward) and 5′-CGGCGTCGAAAATGTTAGGG-3′ (reverse) were designed with the Primer-BLAST tool (NCBI) [Ye et al., 2012], and their amplification efficiencies (E) were determined by serial cDNA dilutions; primers were supplied by Exxtend. Fluorescence was detected using the QuantStudio 5 System (Applied Biosystems) under standard temperature protocols.

Detection of a uAUG Codon-Creating EFNB1 Variant

As there was a clinical diagnosis of CFNS, the patient was referred to our center for analysis of EFNB1. Sanger sequencing of the coding regions did not identify potential causative variants in EFNB1, and next-generation sequencing also confirmed no potential pathogenic variants in the coding regions or in splice sites. However, we identified a variant located within the 5′UTR of EFNB1, NM_004429.4:c.–411C>G. This variant (chrX:68049209C>G) was not inherited from the patient’s parents (Fig. 1c), and the analysis of microsatellite marker patterns was consistent with the correct biological relationships (correct segregation of 15/15 markers). The identified variant was novel and absent from 1000 Genomes, dbSNP150, ABraOM, and gnomAD.

The WT 5′UTR sequence of EFNB1 contains 2 uORFs on the positive strand and downstream of the c.–411C>G variant: the first one is the functional 4 codon uORF (uORF1), studied by Twigg et al. [2013]; the second is a 10 codon uORF (uORF2) that was predicted by ORFfinder in silico analysis. Substitution of C by G at position c.–411 was predicted to introduce a uAUG start codon, which is in-frame with the uUGA stop codon of uORF2, establishing the creation of a new 76 codon uORF (hereafter referred to as uORF3) overlapping with uORF2 and sharing the same uUGA stop codon. All uORFs1–3 are out-of-frame with the AUG of the main dORF (Fig. 2a).

Additionally, we investigated whether reported variants in gnomAD would alter, create, and/or delete some uORFs in the 5′UTR of EFNB1. Forty-seven point variants (allele frequency <0.09) were found. None of them were predicted to modify uORFs in the sequence, according to predefined parameters of the ORFfinder.

As the sequence around uAUGs has been associated with the likelihood of translation initiation, we evaluated the AUG context in uORFs1–3 and the main dORF. In the Kozak consensus, the most critical bases are a purine (A or G) at position −3 and a G at position +4 (the A of AUG, is designated as +1) [Kozak, 1987, 1989, 2002]. The uORF3 sequence has a match with the Kozak sequence at purine −3. The uORF2 sequence context lacks a purine at position −3 and the G base at position +4, while uORF1 and the main dORF do have a match at position −3 and +4 of the Kozak sequence. The AUGs of uORF1–3 and the main dORF were scored as probable translation sites (NetStart score > 0.5) (online suppl. Fig. 1; for suppl. material see www.karger.com/doi/10.1159/000490635).

The c.–411C site was found to be evolutionary conserved (GERP score = 2.48) and no uATG was identified in any species at this position. Analysis of EFNB1 orthologues in 20 vertebrates revealed a high degree of identity, most strongly among mammalian species. No uORFs were observed near the human c.–411C, with the exception of 24 and 16 codon uORFs found in zebra finch and zebrafish, respectively. The region comprising uORF2 shows a high degree of sequence homology in several species of mammals, while zebra finch and zebrafish also have a uORF just a few bases downstream the start codon of human uORF2. The multispecies alignment comprising uORF1–3 and the main ORF show more than 1 uORF in any species (online suppl. Fig. 1). Collectively, these observations suggest that the introduction of a uATG, by the c.–411C>G variant, could potentially interfere with EFNB1 translation.

c.–411C>G Reduces Translation of the Main EFNB1 ORF

We performed dual luciferase reporter assays to functionally evaluate the regulatory importance of the 5′UTR and to test the effect of c.–411C>G on protein expression from the main dORF, which could explain the observed CFNS phenotype. In the presence of c.–411C>G, transla-
of-frame with and overlaps the main dORF. Asterisks indicate nucleotide changes, and blue bars represent uORFs. The AUG sequence context is shown for each uORF/ORF up to base –3 and +4. b Graph depicting the relative Renilla luciferase activity ([experimental sample ratio – negative control ratio] / [positive control ratio – negative control ratio]) for each construct. c RT-qPCR analysis of relative Renilla luciferase mRNA levels in transfected HEK293T cells. All values are relative to wild type as mean ± SEM of 3 experiments which were performed in triplicate. One-way ANOVA with a post-hoc Bonferroni multiple comparison test was done. * p < 0.05, ** p < 0.01, *** p < 0.001.

To determine if the c.–411C>G variant in our patient has similar regulatory effects to c.–95T>G, previously described in CFNS by Twigg et al. [2013], we compared luciferase activity using the c.–95G construct. Both c.–411C>G (F5162) and c.–95T>G (1330) variants significantly reduced the main dORF translation on average 63% compared to WT.

As an RNA sample was not available from the patient to evaluate if the reduced protein levels could be a consequence of impaired EFNB1 transcription, we transfected HEK293T cells with each construct (F5162, F5162SHORT, 1330, and WT) and analyzed the Renilla luciferase RNA expression. No significant difference was observed between any of the constructs relative to WT (Fig. 2c), suggesting that the effect of the c.–411C>G variant is most likely at the level of translation.
Discussion

There are an increasing number of examples of diseases in which the underlying mechanisms are caused by variants at 5'UTR [Calvo et al., 2009; Barbosa et al., 2013]. However, pathogenicity interpretation of such variants is usually difficult, and detailed functional analyses are required to provide evidence of causality. Here, we add an additional, novel, and de novo c.–411C>G variant in EFNB1, to the still scarce list of pathogenic 5'UTR variants causative of CFNS.

The c.–411C>G change is a de novo variant located in a highly evolutionary conserved region, providing evidence that it may be functional. In silico analysis predicted the creation of a new uAUG, leading to a new uORF (uORF3) or representing a 5' elongation of a presumed uORF2. Favoring this hypothesis, we observed that this new uAUG of uORF3 has a good Kozak consensus sequence that scored as a probable translation site. The potential functional effect of this variant was also supported by the observation that none of the 47 single nucleotide variants, annotated at the 5'UTR in gnomAD, seemed to disrupt, create, or alter uORFs. Therefore, all the in silico analysis supported a functional effect for the c.–411C>G variant, identified in a region usually not sequenced in CFNS patients.

Addressing this functional hypothesis, through in vitro assays of sequences containing uORF3, our findings suggest that this variant leads to translation dysregulation. Both the F5162 and F5162SHORT assays provided evidence that the c.–411C>G variant decreases protein translation of the main dORF. The F5162SHORT analysis showed a slightly greater reduction in protein expression of the main dORF (compared to F5162 construct), in agreement with the literature for an additional number of uORFs [Iacono et al., 2005; Calvo et al., 2009; Chew et al., 2016; Johnstone et al., 2016]. Furthermore, the c.–411C>G uORF3-creating variant and c.–95T>G uORF1-disrupting variant [Twigg et al., 2013] showed highly similar effects on the protein levels of the main dORF, despite their different positions and sizes. Taken together, these results strongly suggest that the c.–411C>G variant has a functional effect on translation. Although one of the mechanisms thought to be triggered by uORFs is the nonsense-mediated decay [Barbosa et al., 2013], this is not supported by our in vitro mRNA analysis.

To date, at least 24 diseases have been described to be caused by uORF-altering variants in the 5'UTR of different genes. It is interesting to note that the c.–411C>G variant is one of the most distantly located uORF variants compared with other pathogenic uORF-altering mutations in different disorders [Barbosa et al., 2013; Hornig et al., 2016]. Therefore, in CFNS patients negative for pathogenic variants in coding and splice sites of EFNB1, a careful analysis of the entire 5'UTR should be carried out as causative variants in the entire region could result in decreased protein levels and disease.

About 49% of human transcripts contain uORFs that are most likely to be physiologically relevant, with their functional importance supported by conservation in vertebrate evolution. The mean length of uORFs across vertebrates was found to be <60 nucleotides [Johnstone et al., 2016]. The novel 228 nucleotide uORF3 described in this work is much longer than the majority of uORFs present within the EFNB1 5'UTR of the evaluated species, except for 1 predicted uORF found in the mammal hyrax (>680 bp; data not shown in the alignment), in agreement with depletion of long ORFs in vertebrate 5'UTRs [Johnstone et al., 2016].

This study contributes an additional CFNS case with a regulatory variant in the EFNB1 5'UTR sequence and supports screening of this noncoding region in CFNS patients where other variants or gene copy number changes are absent. The 5'UTRs of disease-related genes are not routinely screened, raising the possibility that similar pathogenic variants contribute more widely to disease. In light of this, it is interesting to note that uORFs can be targeted by antisense oligonucleotides to reestablish normal levels of the main protein [Liang et al., 2016]. In this context, a CRISPR-Cas9 approach may be useful to generate different mutations in a disease-relevant cell type, such as iPSC-derived neuroepithelial cells, to provide further insights into pathophysiology of the disease along with investigation of antisense oligonucleotide strategies. In summary, this report adds to the understanding of the functional effects of uAUG/uORFs on protein expression and pathophysiological mechanisms in CFNS.

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Statement of Ethics

Ethics approval for this study was provided by the Ethics Committee of the Instituto de Biociências at Universidade de São Paulo, São Paulo, SP, Brazil (Protocol 024/2004). Patients donated biological samples after providing signed informed consent.

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Disclosure Statement

The authors declare no conflicts of interest.


