De Novo SOX6 Variants Cause a Neurodevelopmental Syndrome Associated with ADHD, Craniosynostosis, and Osteochondromas

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SOX6 belongs to a family of 20 *SRY*-related HMG-box-containing (SOX) genes that encode transcription factors controlling cell fate and differentiation in many developmental and adult processes. For *SOX6*, these processes include, but are not limited to, neurogenesis and skeletogenesis. Variants in half of the SOX genes have been shown to cause severe developmental and adult syndromes, referred to as SOXopathies. We here provide evidence that *SOX6* variants also cause a SOXopathy. Using clinical and genetic data, we identify 19 individuals harboring various types of *SOX6* alterations and exhibiting developmental delay and/or intellectual disability; the individuals are from 17 unrelated families. Additional, inconstant features include attention-deficit/hyperactivity disorder (ADHD), autism, mild facial dysmorphism, craniosynostosis, and multiple osteochondromas. All variants are heterozygous. Fourteen are *de novo*, one is inherited from a mosaic father, and four offspring from two families have a paternally inherited variant. Intragenic microdeletions, balanced structural rearrangements, frameshifts, and nonsense variants are predicted to inactivate the *SOX6* variant allele. Four missense variants occur in residues and protein regions highly conserved evolutionarily. These variants are not detected in the gnomAD control cohort, and the amino acid substitutions are predicted to be damaging. Two of these variants are located in the HMG domain and abolish SOX6 transcriptional activity *in vitro*. No clear genotype-phenotype correlations are found. Taken together, these findings concur that *SOX6* haploinsufficiency leads to a neurodevelopmental SOXopathy that often includes ADHD and abnormal skeletal and other features.

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Introduction

SRY-related high-mobility-group (HMG)-box-containing (SOX) genes code for transcription factors sharing at least 50% similarity with one another in a characteristic HMGtype DNA-binding domain.¹ The 20 SOX genes present in the human genome are distributed into eight groups (SOXA-SOXH) on the basis of the sequence conservation of their proteins within this domain. Further, SOX proteins belonging to the same group also feature significant conservation in sequences outside the HMG domain. SOX genes exhibit overlapping expression patterns and functions, especially among same-group members. Most SOX genes have been shown to participate pivotally in the control of cell fate and differentiation in one or multiple lineages, such that the SOX family as a whole has been implicated in almost every developmental, physiological, or pathological process. In line with these critical roles, to date, variants in half of the SOX genes have been associated with human developmental disorders, called SOXopathies.² The family founder, SRY (MIM: 480000), owes its name to its location in the sex-determining region of the Y chromosome. SRY is necessary for initiating male differentiation in the mammalian embryo,^{3–5} and variants inactivating the gene cause disorders of sex development, including 46,XY sex reversal 1 (MIM: 400044). Besides SRY and SOX3 (MIM: 313430), which are located on the Y and X chromosomes, respectively, all other SOX genes map to autosomal chromosomes, and diseasecausing variants in these genes are most often heterozygous, de novo, and inactivating. For instance, the presence of such variants within and around SOX9 (MIM: 608160) can cause campomelic dysplasia with or without XY sex reversal (CMPD1-SRA1 [MIM: 114290]); variants in SOX10 (MIM: 602229) cause peripheral demyelinating neuropathy, central demyelination, Waardenburg syndrome and Hirschsprung disease (PCWH [MIM: 609136]), and the Waardenburg syndrome types 2E (MIM: 611584) and 4C (MIM: 613266); and variants in SOX2 (MIM: 184429) cause anophthalmia, syndromic microphthalmia-3 (MCOPS3 [MIM: 206900]), optic nerve hypoplasia, and abnormalities of the central nervous system. Variants in several other SOX genes were recently shown to be responsible for intellectual disability and other features. Namely, the heterozygous loss of function of SOX5 (MIM: 604975) causes Lamb-Shaffer syndrome (LAMSHF [MIM: 616803]), which is associated with global developmental delay, intellectual disability, and mild dysmorphic features;^{6,7} SOX11 (MIM: 600898) haploinsufficiency causes Coffin-Siris-like syndrome-9 (MIM: 615866), which is characterized by intellectual disability, microcephaly, and dysmorphic features;^{8,9} and heterozygous de novo missense variants in SOX4 (MIM: 184430) cause Coffin-Siris-like syndrome-10 (MIM: 618506), which features intellectual disability and mild facial and digital skeletal abnormalities.¹⁰

Human *SOX6* (MIM: 607257) is located at 11p15 and is expressed in a variety of tissues.^{11,12} Its mouse ortholog

was shown to share essential redundant functions with its closest SOXD relative, Sox5, in the differentiation of chondrocytes^{13,14} but to be uniquely expressed and exert other important functions in skeletal myoblast differentiation,¹⁵ cardiomyocyte proliferation,¹⁶ erythroid cell maturation,¹⁷ and regulation of insulin secretion.¹⁸ Further, mouse Sox6 is also pivotal in the developing central nervous system, where it regulates the development of cortical interneurons, ^{19,20} dopaminergic neurons in the substantia nigra pars compacta,²¹ and oligodendrocytes.²² Genomewide association studies have linked human SOX6 variants to a variety of adult clinical conditions, such as variability in bone mineral density,²³ carotid plaque formation,²⁴ high blood pressure,²⁵ and obesity.²⁶ In oncology, SOX6 copy-number variants (CNVs) have been found in esophageal squamous cell carcinoma,²⁷ Ewing sarcoma,²⁸ and glioblastoma.²⁹ To date, however, no developmental disorder has been definitively associated with SOX6 variants.

Here, we report 19 individuals who carry *SOX6* variants, share milestone delays and intellectual disability, and exhibit inconstant abnormalities, including mild dysmorphism, craniosynostosis, and osteochondromas; the individuals are from 17 unrelated families. The variants are private in each affected individual (apart from relatives), and all except two missense variants are predicted to abrogate or impair *SOX6* expression or protein activity. We thus propose that *SOX6* haploinsufficiency underlies a neurodevelopmental SOXopathy associated with other variable features.

Material and Methods

Recruitment of Subjects

The study cohort consisted of 19 individuals from 17 unrelated families originating from Belgium, Canada, France, Germany, the Netherlands, Slovenia, the UK, and the US. All individuals had molecular karyotyping, whole-exome sequencing (WES), or whole-genome sequencing (WGS) performed as part of local neurodevelopmental studies on developmental delay and intellectual disability or congenital abnormalities. Informed consent for participating in the genetic studies was obtained via protocols approved by institutional review boards of local hospitals. The parents or legal guardians of subjects provided consent for publication of all photographs shown in this study. No consanguinity was noted in the families. All affected individuals with a CNV or single-nucleotide variant (SNV) have been registered in the DECI-PHER database (Table 1).

Genomic Variant Diagnosis

Genomic DNA was extracted from peripheral blood via standard protocols. Molecular karyotyping was conducted with various arrays according to the manufacturers' instructions (Table S1). Deletion breakpoints were mapped to the human genome assembly hg19 of the UCSC Genome Browser. Multiplex ligation-dependent probe amplification (MLPA) or qRT-PCR was used for confirming the genomic alteration in each individual and determining parental inheritance. WES was performed with target-enrichment designs according to the manufacturers' instructions. We filtered

	Internal ID; DECIPHER ID	Sex	Genomic Variant (GenBank: NM_ 033326.3; hg19)	Protein	Inheritance	Age	OFC	Weight	Height	ID	Facial Dysmorphism	Osteochondromas	Behavior	Other Features
1	PIT-1; 406933	male	deletion of exons 1 and 2	-	affected father	7 years	0 SD	+1 SD	+1 SD	mild	large ears, large nose	multiple osteochondromas	ADHD, autism, aggressive, emotional lability, tantrums, staring spells	submucous cleft palate
2	PIT-2	male	deletion of exons 1 and 2	-	affected father	3 years	-0.5 SD	–0.5 SD	–0.5 SD	mild	-	multiple osteochondromas	ADHD, autism, sleep disruption, staring spells, and left-sided weakness	bilateral nasolacrimal duct obstructions, food allergies
3	PIT-3	female	deletion of exons 1 and 2	-	affected father	12 months	+0.5 SD	+0.5 SD	-2.5 SD	normal	-	-	-	diastasis recti, umbilical hernia cardiac rhabdomyoma
1	CHUN-1; 406934	male	deletion of exons 1–4	-	healthy father, mosaic 22%	10 years	+1 SD	+2 SD	+0.5 SD	mild	-	-	ADHD, anxiety	-
5	CHLM-1; 406929	female	deletion of exons 2 and 3	-	de novo	10 years	+1 SD	+3.5 SD	+4.5 SD	mild	-	-	-	arachnodactyly, precocious puberty
5	LJU-1; 406932	male	deletion of exons 2–13	_	affected father	7 years	-0.5 SD	0 SD	0 SD	mild to moderate	-	-	-	hypermetropia
7	IHG-1; 406930	male	deletion of exons 2–12	-	de novo	9 years, 4 months	+2 SD	+4 SD	+0.5 SD	mild	-	-	restlessness, short attention span, quick changes of mood, aggressive	_
8	CHUP-1; 406931	female	deletion of exons 5–7	-	de novo	12 years	-1 SD	-1 SD	-1 SD	moderate	oxycephaly (sagittal, metopic, and coronal craniosynostosis), micrognathia	-	attention deficit, anxiety	-
9	UK-1; 412103	male	c.242C>G	p.Ser81*	de novo	9 years, 7 months	-2 SD	-1 SD	0 SD	moderate	scaphocephaly (sagittal craniosynostosis), prominent occiput, hypertelorism	-	ADHD, lack of sense of danger, destructive, easily upset	short fifth fingers with clinodactyly, flat feet with valgus heels
10	UK-2; 412119	female	c.277C>T	p.Arg93*	unknown	6 years	-1 SD	-0.8 SD	+1 SD	mild	scaphocephaly (sagittal and left coronal craniosynostosis)	-	attention deficit, sleep disturbance, aggressive episodes	-
11	CHUN-2; 412120	male	c.293C>G	p.Ser98*	de novo	6 years	-1 SD	-1 SD	-1 SD	mild to moderate	high forehead	-	ADHD	ogival palate

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	Internal ID; DECIPHER ID	Sex	Genomic Variant (GenBank: NM_ 033326.3; hg19)		Inheritance	Age	OFC	Weight	Height	ID	Facial Dysmorphism	Osteochondromas	Behavior	Other Features
12	GDX-3; ^a 412130	male	c.483G>C	p.Trp161 Cys	de novo	13 years	+1 SD	+1 SD	+1.5 SD	severe	-	-	ADHD	obesity
13	GDX-1; 412121	female	c.718C>T	p.Gln240*	de novo	10 years	0 SD	+2 SD	+1 SD	mild	bitemporal narrowing, small mouth	-	ADHD, anxiety	bilateral inverted nipples, sensorineural hearing loss, vestibular dysfunction
14	GDX-2; 412122	female	c.878delC	p.Pro293 Leufs*3	de novo	13 years	0 SD	+1.5 SD	-1 SD	mild	short palpebral fissures, hooded eyelids, hypertelorism, wide nasal bridge, low-set ears	-	ADHD	high arched palate; long, tapering fingers; hypotonia
15	PS-1; 412123	female	c.1728del	p.Glu577 Argfs*29	de novo	27 years	0 SD	+4 SD	-1.5 SD	moderate	synophrys	-	-	short extremities, hirsutism, dental abnormalities, Hashimoto's thyroiditis
16	CHOP-1; 412124	female	c.1814T>C	p.Met605 Thr	de novo	6 years	0 SD	+1 SD	+2 SD	mild	-	multiple osteochondromas	ADHD	dysgraphia, developmental delay
17	LEUV-1; 412126	female	c.1915T>A	p.Trp639 Arg	de novo	3 years	-0.9 SD	+0.1 SD	+1.2 SD	mild to moderate		-	ADHD, autism	tremor hands
18	LEID-1; ^b 412127	male	c.2237C>T	p.Ser746 Leu	de novo	27 years	-0.2 SD	0 SD	-0.5 SD	severe	long face, triangular; full eyebrows	_	autism, abnormal movements, automutilation, hyperventilation	-
19	CHUG-1	male	breakpoint in intron 13 of <i>SOX6</i>	-	de novo	2.5 years	+0.7 SD	+0.6 SD	+0.8 SD	mild to moderate		-	ADHD, oppositional disorder, self- endangerment, sleep disorders	gestural dyspraxia, graphomotor difficulties, troubles in neurosensory integration

Abbreviations are as follows: ID, intellectual disability; OFC, occipitofrontal circumference. ^aNote that GDX-3 also harbored a hemizygous likely pathogenic *MECP2* variant that is consistent with his clinical presentation. ^bIn addition to trio WES, other targeted analyses were performed (e.g., *MECP2* and *TCF2* were analyzed by Sanger sequencing, and Smith-Magenis syndrome was analyzed by fluorescence *in situ* hybridization), but no other candidate variant was detected.

sequence variants in a stepwise manner to exclude synonymous variants, non-exonic SNVs, indels and variants with a minor allele frequency > 1% in gnomAD (version v2.1.1), the 1000 Genomes Project, and internal exome databases. WGS was performed for affected individual CHUG-1 at the Centre Hospitalier Universitaire de Grenoble and for affected individuals UK-1 and UK-2 by the Genomics England 100,000 Genomes Project. SNVs were confirmed by Sanger sequencing. The *SOX6* reference sequence GenBank: NM_033326.3 was used in naming these variants.

In Silico Assessment of SOX6 Variant Pathogenicity

SOX6 synonymous and missense variants present in control individuals were downloaded from gnomAD (version v2.1.1)³⁰ and analyzed with paired t tests. SOX sequences were downloaded from NCBI (Tables S2 and S3) and aligned with the ClustalW tool embedded in MacVector16 software (MacVector). The effects of missense variants on protein structure were predicted with HOPE,³¹ SWISS-MODEL,³² and PEP-FOLD3.³³ The best-scoring models were selected.

Functional Assessment of SOX6 Variants In Vitro

Expression plasmids for SOX6 missense variants were generated by PCR mutagenesis using appropriate primers (Table S4) and an expression plasmid for mouse wild-type (WT) SOX6 tagged with an N-terminal 3FLAG epitope.¹² Plasmid integrity was verified by Sanger sequencing. For reporter assays, HEK293 cells (CRL-1573; ATCC) were transfected in triplicate cultures with 3.5 µL ViaFect Transfection Reagent (Promega) containing 100 ng pNL1.1.TK[Nluc/TK] (NanoLuc Renilla luciferase control reporter plasmid), 400 ng mouse Acan [4xA1]-p89Luc (SOX6/9-dependent firefly luciferase reporter),³⁴ 75 ng 3FLAG-SOX9 plasmid,³⁵ and the indicated amounts of plasmids encoding no protein, 3FLAG-SOX6, and a 3FLAG-SOX6 variant for a total of 1,000 ng DNA. Cells were collected after 24 h in Tropix Lysis buffer (Applied Biosystems) supplemented with a protease inhibitor cocktail (Thermo Fisher Scientific), and extracts were subjected to the Nano-Glo Dual-Luciferase Reporter Assay (Thermo Fisher Scientific). Reporter activities were measured with a GloMax Explorer Multimode Microplate Reader (Promega). They are presented in the figures as means \pm the standard deviation of firefly luciferase values measured for biological triplicates and normalized for transfection efficiency with NanoLuc values.

We tested SOX6 intracellular localization by transfecting HEK293 or COS-1 cells (CRL-1650; ATCC) with ViaFect (3.5 μ L) and empty or SOX6 expression plasmid (1,000 ng per 10 cm² dish). Extracts were prepared the next day with NE-PER Nuclear and Cytoplasmic Extraction Reagents (Thermo Fisher Scientific). Immunoblots were carried out with FLAG M2-peroxidase-conjugated antibody (A8592, Sigma-Aldrich), P84 antibody (GTX70220-01, GeneTex), and β -actin antibody (sc-47778 [c4], Santa Cruz Biotechnology) as previously described.¹⁰

The abilities of SOX6 variants to homodimerize and to bind DNA were tested upon transfection of COS-1 cells with ViaFect ($3.5 \ \mu$ L) and empty or SOX6 expression plasmid (1,000 ng per 10 cm² dish). Whole-cell extracts were prepared after 40 h in 14 mM HEPES buffer (pH 7.9) containing 1.5 mM MgCl₂, 6.0 mM KCl, 0.44 M NaCl, 0.08 mM EDTA, 2.3 mM DTT, 10% glycerol, and a protease inhibitor cocktail. Protein homodimerization was assessed by immunoblot after a 10 min incubation of cell extracts with varying amounts of glutaraldehyde, up to 0.01%. SOX6's binding to DNA was tested in an electrophoretic mobility

shift assay (EMSA) with a DIG Gel Shift Kit, 2^{nd} Generation (Sigma-Aldrich). The DNA probe contained a SOXD (SOX5 or SOX6) binding site that is located in the *Acan* enhancer used in the reporter assay described above (Agc1 K).³⁴ Mixtures contained 0.5 µL cell extract, 10 fmol DIG-labeled probe, and 25 ng poly [dG.dC] in a 20 mM HEPES buffer (pH 7.9) containing 40 mM KCl, 0.5 mM EDTA, 0.5 mM DTT, and 10% glycerol. After 30 min of incubation at 30°C, they were electrophoresed in 4% native polyacrylamide gels in TGE buffer (50 mM Tris, 0.4 M glycine, and 4.5 mM EDTA [pH 8.0]). We transferred DNA to a Zeta-Probe GT nylon membrane (Bio-Rad Laboratories) by electroblotting, and we detected signals according to the manufacturer's instructions by using a Chemi-Doc Imaging System (Bio-Rad Laboratories).

Evaluation of SOX6 Expression in the Human Brain

We used RNA sequencing (RNA-seq) data available in the BrainSpan Atlas of the Developing Human Brain³⁶ to assess the expression of *SOX6* in various anatomical sites and at various developmental and adult ages.

Results

De Novo SOX6 Variants Associate with a Neurodevelopmental Syndrome

We identified 19 individuals sharing developmental milestone delays and/or intellectual disability and carrying various types of SOX6 variants (Table 1). Ten individuals were males, and nine were females. At the most recent examination, their ages ranged from 12 months to 27 years. Pregnancies were uneventful, and postnatal growth parameters were generally in the normal range. Most individuals had intellectual disability (18/19) varying from mild (8/19) to mild-moderate (3/19), moderate (3/19), and severe (2/19). Abnormal behavior was observed in many affected individuals (14/19) and included attention-deficit/hyperactivity disorder (ADHD; 10/19) or autism spectrum disorder (4/19). Additional behavioral concerns included anxiety, aggressiveness, and automutilation. Dysmorphic features and skeletal anomalies were noted in many affected individuals. They included a mild facial dysmorphism, but this feature was nonspecific and thus not consistent with a clinically recognizable syndrome (Figure 1A). Of note, sagittal craniosynostosis with scaphocephaly was observed in two unrelated affected individuals, and oxycephaly with synostosis of the sagittal, metopic, and coronal sutures was observed in a third unrelated individual. Moreover, multiple diffuse osteochondromas were present in two brothers (7-year-old PIT-1 and 3-year-old PIT-2) and in one girl from an unrelated family (6-year-old CHOP-1) (Table S5). Variants in EXT1 (MIM: 608177) and EXT2 (MIM: 608210), which are deleted in a large percentage of individuals with exostoses (MIM: 133700 and MIM: 133701, respectively) and are not chromosomally linked to SOX6, were not detected in any of the individuals with osteochondromas. Interestingly, several of the osteochondromas that were initially detected in affected individual CHOP-1 at 4 years of age

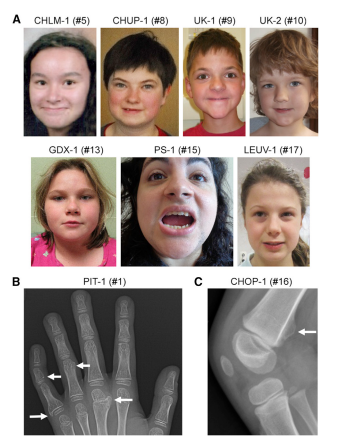


Figure 1. Clinical Findings in Subjects with SOX6 Variants (A) Photos of seven subjects showing mild, nonspecific facial dysmorphism.

(B) X-ray showing multiple osteochondromas (marked by arrows) in the right hand of affected individual PIT-1.

(C) X-ray showing an osteochondroma (marked by arrow) at the right distal femur of affected individual CHOP-1.

were no longer detected on X-rays or were no longer palpable 2 years later (Table S5 and Figure 1B). The girl had no complaints of pain, and she ran and played without difficulty. Taken together, these data suggest that *SOX6* variants cause a unique form of human neurodevelopmental disorder often associated with mild dysmorphism and occasionally associated with craniosynostosis or osteochondromas.

SOX6 Translocation, CNVs, and SNVs Associate with Disease

The 19 affected individuals in our study displayed a spectrum of *SOX6* variants (Table 1). All individuals were heterozygous for the *SOX6* variant. Whereas 14 of them carried a *de novo* variant, four (including the PIT-1, PIT-2, and PIT-3 siblings) inherited the variant from their affected father, and one inherited it from an unaffected father who showed 22% mosaicism for the variant. The father of the PIT-1, PIT-2, and PIT-3 siblings presented with mild intellectual disability, but no further clinical evaluation was available, especially regarding the presence of osteochondromas.

Eight affected individuals (1–8), including the PIT-1– PIT-3 siblings, harbored a CNV, and one individual (19, CHUG-1) carried a balanced structural variant (Figure 2A). *SOX6* gives rise to four main transcripts that differ from one another in alternative promoter usage and alternative splicing of an exon encoding an internal, dispensable protein sequence. The most abundant transcript (GenBank: NM_033326.3) encodes a 5' untranslated exon 1 and 15 coding exons. The reciprocal translocation detected in CHUG-1 occurred between the chromosomal regions 11p15.2 and 2p12 (Figure 2B). The 11p15.2 breakpoint was located in intron 13 of *SOX6*, thus separating exons 14–16, which encode the HMG domain and C terminus of the protein, from the more upstream exons.

The CNVs detected in all eight affected individuals (DECIPHER: 406929-406934) were different from one another (except for the those detected in the three siblings; Figure 2A and Table S6). All were partial deletions of SOX6 and did not involve any other gene bodies. The PIT-1, PIT-2, and PIT-3 siblings' deletion involved exons 1 and 2 of all main SOX6 transcripts. CHUN-1's deletion included exons 1-4 of all main SOX6 transcripts. CHLM-1's deletion encompassed exons 2 and 3 of GenBank: NM_033326.3 and exons 1-3 of the other transcripts. IHG-1's and LJU-1's deletions comprised exons 2-12 and 2-13 of GenBank: NM_033326.3, respectively, and exons 1-11 and 1-12 of the other transcripts, respectively. CHUP-1's deletion removed exons 5-7, including most of the primary coiled-coil (CC1) domain sequence (discussed later in the manuscript). Splicing from exon 4 to exon 8 of GenBank: NM_033326.3 would not affect the coding sequence frame but would result in a SOX6 lacking a functional CC1 domain, and splicing of exons 4-9 would result in a frameshift with a stop codon after 27 residues. Thus, the balanced translocation and all deletions most likely inactivated the affected SOX6 allele. Because no other abnormalities were detected in the microarray tests, the resulting diseases were postulated to reflect SOX6 haploinsufficiency.

The other affected individuals in the study carried an SNV in SOX6. All SNVs were distinct and corresponded to four nonsense, two frameshift, and four missense mutations (Table 1). All, except one in an individual whose parents were unavailable, arose de novo and were absent in gnomAD control populations. The SOX6 encoded by GenBank: NM_033326.3 comprises 808 amino acids (Figure 3A). Its known functional domains are the HMG domain and two coiled-coil homodimerization domains, of which CC1 is longer and more critical than the secondary coiled-coil (CC2) domain.¹² The four nonsense variants caused SOX6 truncation upstream of CC1 (p.Ser81* [UK-1], p.Arg93* [UK-2], and p.Ser111* [CHUN-2]) or within CC1 (p.Gln240* [GDX-1]). In addition to encoding a short protein lacking all known functional domains, the mutant mRNAs were likely to undergo nonsense-mediated decay. The two frameshift variants (p.Pro293Lfs*3 [GDX-2] and p.Glu577Argfs*29 [PS-1]) caused SOX6 truncation

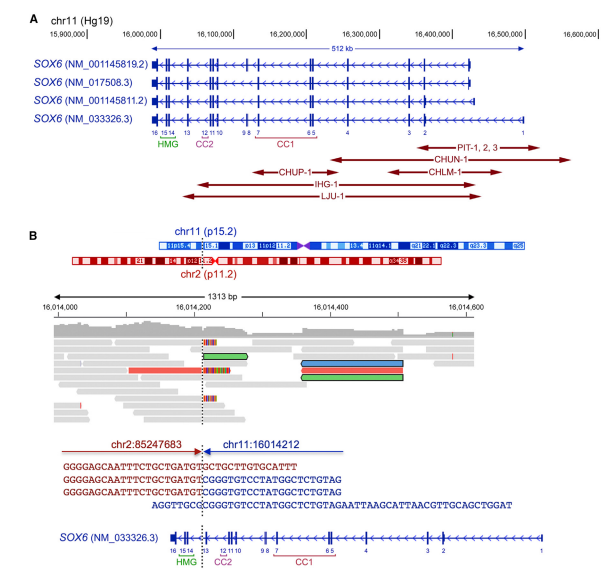


Figure 2. Structural Variants Detected in Affected Individuals 1-8 and 19

(A) Location of CNV variants. The main *SOX6* transcript isoforms are schematized; taller vertical lines correspond to coding sequences, smaller vertical lines correspond to 5' and 3' untranslated sequences, and arrowheads in introns point to the transcriptional direction. NCBI accession numbers are indicated. The GenBank: NM_033326.3 coding exons are labeled 1–16. Coordinates of chromosomal region 11p15 are shown above the schematics. Double-arrowed lines depict the microdeletions identified in subjects 1–8. CC1, primary coiled-coil domain; CC2, secondary coiled-coil domain; HMG, HMG domain.

(B) Location of the breakpoint of the 46,XY,t(2;11)(p11.2;p15.2)-balanced reciprocal translocation identified in affected individual 19. From top to bottom are schematics of chromosomes 2 and 11, in which a vertical dotted line indicates the breakpoint involving 11p15.2 and 2p11.2; reads (BAM file) aligned by the Integrative Genomics Viewer (IGV); the sequence of the breakpoint, in which red and blue lines show the normal sequences of chromosomes 2 and 11, respectively, and red sequences followed by blue sequences show the sequence junction; and the same representation of *SOX6* as in (A).

before CC2 and the HMG domain, respectively, and thus also most likely generated non-functional proteins. The missense variants affected different protein regions. p.Met605Thr (CHOP-1) and p.Trp639Arg (LEUV-1) occurred in the HMG domain, whereas p.Trp161Cys (GDX-3) and p.Ser746Leu (LEID-1) occurred in regions of unknown functions. p.Trp161Cys was located in the N terminus of SOX6, close to the CC1 domain, and p.Ser746Leu (LEID-1) was located in the C terminus of the protein. None of the missense variants were located at or near exon boundaries or in the alternatively spliced exon. Thus, although we predicted the nonsense and frameshift SNVs to be pathogenic, we needed to undertake further analyses to determine whether the missense SNVs could be pathogenic too.

SOX6 Missense Variants Affect Residues Highly Conserved Evolutionarily and Mutated in Several SOXopathies

MetaDome analysis of mutation tolerance in the *SOX6* coding sequence in control human populations showed, not surprisingly, that the HMG domain is the most

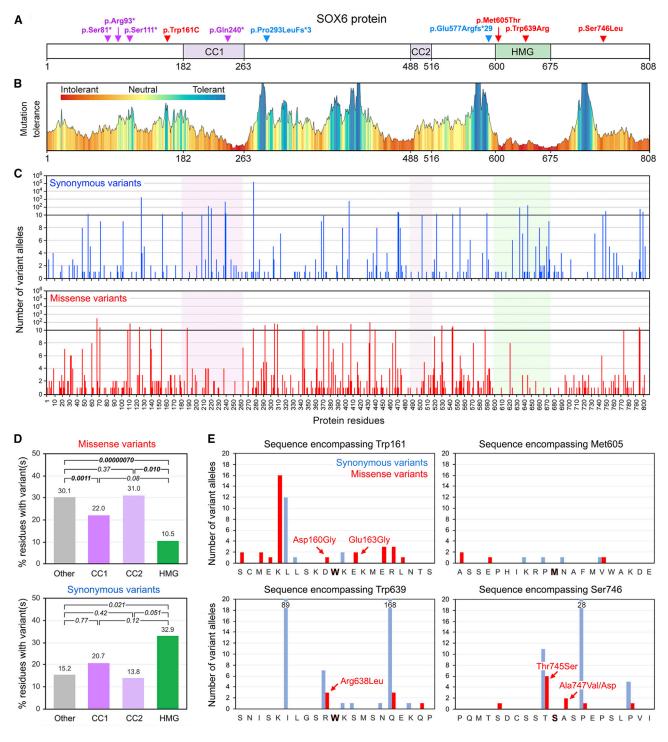


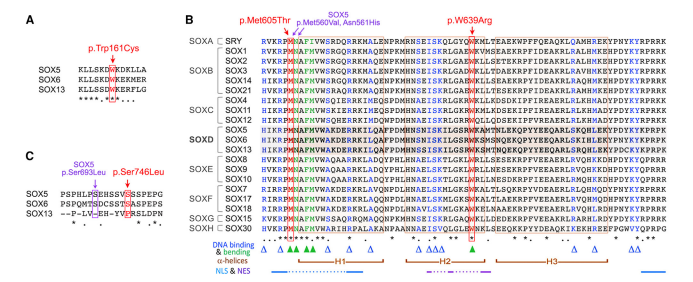
Figure 3. Analysis of SOX6 SNVs in Affected Individuals 9–18 and in gnomAD Individuals

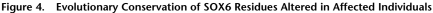
(A) Location of study subjects' SNVs in the SOX6 isoform encoded by the GenBank: NM_033326.3 transcript. The protein and domain residue boundaries are indicated underneath the schematic. CC1, primary coiled-coil domain; CC2, secondary coiled-coil domain; HMG, HMG domain. Red represents missense variants, purple represents nonsense variants, and blue represents frameshift variants. (B) Plot of the mutation tolerance of SOX6 residues downloaded from MetaDome.

(C) Counts and distribution of SOX6 synonymous and missense variants found in gnomAD individuals.

(D) Percentages of residues carrying at least one missense or synonymous variant in the functional and other domains of SOX6 in gnomAD individuals. We performed paired t tests to calculate the statistical significance of differences between domains. p values are indicated.

(E) Numbers of synonymous and missense variants detected in gnomAD individuals in the sequences encompassing the missense variants identified in four study subjects. The nature of the missense variants closest to the residues altered in the four affected individuals is indicated.





(A) Alignment of human SOXD sequences encompassing Trp161. Asterisks represent fully conserved residues, and dots represent semiconserved residues.

(B) Alignment of the HMG domain sequences of all human SOX proteins: residues altered in SOX6 in affected individuals in our study are shown in red, and residues altered in SOX5 in affected individuals with LAMSHF syndrome are shown in purple. Triangles represent residues mediating DNA binding (blue) and bending (green). Brackets represent H1, H2, and H3 α helices. Lines linked with dots represent key amino acids in nuclear localization signal sequences (NLSs) and nuclear export signal sequences (NESs). (C) Alignment of human SOXD sequences encompassing Ser746 with indication of the position of SOX5 and SOX6 variants identified in

affected individuals in our study.

intolerant region, but it also showed other intolerant regions, including CC1 (Figure 3B). On the basis of this finding, we analyzed the frequency and nature of SOX6 variants in control individuals by using gnomAD, a database containing genomic and exonic sequences from more than 140,000 unrelated people of various genetic backgrounds.³⁵ Constraint metrics indicated that SOX6 is under tight conservation constraint because 42.6 loss-offunction (nonsense) variants were expected, but only four were observed, resulting in a probability of loss-offunction intolerance of 1.00. Interestingly, whereas 169.7 synonymous variants were expected and 167 were observed (constraint Z score = 0.16), 454.7 missense variants were expected, but only 339 were observed (constraint Z score = 1.93). These findings thus suggested that SOX6 missense variants might often be pathogenic.

Examination of the SNV distribution revealed that missense and synonymous variants occurred throughout the *SOX6* coding sequence (Figure 3C). However, although the three protein domains showed no statistically significant difference (t test, $p \le 0.01$) in their percentages of residues carrying at least one synonymous variant, the HMG ($p = 7 \times 10^{-7}$) and CC1 ($p = 1.1 \times 10^{-3}$) domains had significantly fewer residues with missense variants than the CC2 domain and all other protein regions combined (Figure 3D). This result was consistent with the critical functions of the HMG and CC1 domains. Importantly, no missense variant was recorded in gnomAD individuals in the codons carrying a missense variant in our subjects (Figure 3E). Several gnomAD missense variants were located close to Trp161 and Ser746, and a few were close

to Met605 and Trp639. Thus, although *SOX6* missense variants are frequent in control individuals, those found in the affected individuals in our study have not been detected, leaving open the possibility that they are pathogenic.

Since residues important for protein function or regulation are generally conserved evolutionarily, we sought to determine whether Trp161, Met605, Trp639, Ser746, and their neighboring residues are conserved in SOX6 orthologs and other SOX proteins. The alignment of full-length SOX6 sequences from multiple vertebrate species revealed a high degree of evolutionary conservation both within and outside the known functional domains (Figure S1). This conservation included the residues altered in the affected individuals in our study and their neighboring residues. The alignment of SOX5, SOX6, and SOX13 sequences showed that Trp161 and its neighbors are also highly conserved among SOXD proteins (Figure 4A). The alignment of all SOX HMG domains showed full conservation at the positions corresponding to Met605 and Trp639 in SOX6 (Figure 4B). Furthermore, the Met605 neighbors are fully conserved too, and the Trp639 neighbors are conserved in SOXD and several other SOX proteins. Finally, Ser746 is conserved between SOX5 and SOX6, even though its neighbors are only weakly conserved (Figure 4C).

We next asked whether variants in other SOX genes at residues matching those found in *SOX6* in our subjects have been reported to cause disease. Although no SOX5 variant of the residue matching Trp161 or its neighbors has been associated with LAMSHF disease yet, a SOX5

Table 2.Pathological Missense Variants in Various SOX ProteinsMatch the Two SOX6 HMG Domain Variants Identified in AffectedIndividuals

Protein	Variant	Phenotype	Reference
sox va	ariants Matc	hing the SOX6 p.Met605Th	ır Variant
SOX6	p.Met605Thr	ID with mild skeletal defects	this study
SRY	p.Met64Ile	XY sex reversal	Berta et al. ³⁷
SRY	p.Met64Arg	gonadal dysgenesis and XY sex reversal	Scherer et al. ³⁸
SOX5	p.Met560Val	Lamb-Shaffer syndrome	Zawerton et al. ⁷
SOX10	p.Met108Thr	Kallmann syndrome	Pingault et al. ⁴⁹
SOX Va	ariants Matc	hing the SOX6 p.Trp639Ar	g Variant
SOX6	p.Trp639Arg	ID with mild skeletal defects	this study
SRY	p.Trp98Arg	XY sex reversal with primary amenorrhea and low testosterone	Philibert et al. ⁵⁰
			Bastian et al. ⁵¹
SRY	p.Trp98Cys	gonadal dysgenesis and XY sex reversal	Dastiali et al.
	p.Trp98Cys p.Trp79Ser		Chassaing et al. ⁵²
SRY SOX2 SOX9		XY sex reversal bilateral anophthalmia	
SOX2 SOX9	p.Trp79Ser	XY sex reversal bilateral anophthalmia with GH deficiency campomelic dysplasia	Chassaing et al. ⁵²

p.Met560Val variant matching the SOX6 p.Met605Thr variant and a SOX5 p.Asn561His variant have caused LAMSHF syndrome⁷ (Figure 4B and Table 2). In the SRY residue, two variants matching SOX6 p.Met605Thr (p.Met64Ile and p.Met64Arg) have caused gonadal dysgenesis, and a SOX10 p.Mer108Thr variant, identical to the SOX6 p.Met605Thr variant, has caused Kallmann disease (MIM: 147950).^{37,38} Variants in the residue equivalent to SOX6 Trp639 were found to cause disease in SRY, SOX2, SOX9, SOX10, and SOX17; several of these variants are identical to the SOX6 variant (Trp \rightarrow Arg substitution). Finally, an affected individual with a SOX5 p.Ser693Leu variant, located close to the SOX6 Ser746 residue, has LAMSHF syndrome⁷ (Figure 4C). Together, sequence conservation and SOXopathy data strongly support the notion that the four missense variants found in the affected individuals in our study might be pathogenic.

Affected Individuals' Missense Variants Might Affect the Local Structure of SOX6

We used several software programs to get insights into the possible structural impact of the *SOX6* missense variants detected in the affected individuals in our study (Figure S2). HOPE indicated that the p.Trp161Cys variant could alter SOX6 interactions with other molecules because cysteine is smaller than tryptophan. SWISS-MODEL, a template-based structure-prediction software, and PEP-FOLD3, a *de novo* software predicting protein

structures directly from amino acid sequences, predicted that residues 153-170, which are centered around Trp161, could form an α helix juxtaposed to CC1, and PEP-FOLD3 proposed that the replacement of tryptophan by cysteine would disrupt this structure. Regarding p.Met605Thr, HOPE indicated that the substitution of the sulfur-containing nonpolar methionine by a hydroxyl-containing, polar, and smaller threonine could affect the ability of the HMG domain to bind and bend DNA. For p.Trp639R, PEP-FOLD3 did not predict a change in the helical structure of the HMG domain, but HOPE indicated that the substitution of the bulky aromatic tryptophan with a positively charged arginine would most likely affect the interaction of the HMG domain with DNA. In regard to p.Ser746Leu, SWISS-MODEL could not predict a specific structure for the region encompassing Ser746, but PEP-FOLD3 predicted that this region forms a small α helix regardless of whether the 746 position is occupied by a serine or a leucine. Thus, a change in protein structure is unlikely for this variant. However, although both serine and leucine have a short side chain, this chain is polar and contains a hydroxyl group in serine, whereas it is aliphatic and nonpolar in leucine. It is thus possible that the p.Ser746Leu variant affects the ability of SOX6 to interact with other proteins. In conclusion, each SOX6 variant could destabilize the local structure or biochemical properties of SOX6 and thereby impair protein function or stability.

Affected Individuals' Missense Variants Affect the Nuclear Localization and Transcriptional Activity of SOX6 In Vitro

To test whether the missense variants detected in our subjects could affect SOX6 function, we transiently transfected expression plasmids for the WT and variant SOX6 proteins in COS-1 and HEK293 cells. Immunoblots of cytoplasmic and nuclear extracts showed that all variants were efficiently expressed and that the p.Met605Thr and p.Trp639Arg proteins were not translocated or retained into the nucleus as efficiently as WT SOX6 and the other two variants (Figure 5A). Because Trp161 could interact with CC1, we treated cell extracts with glutaraldehyde, an assay previously used to detect SOX6 homodimers¹² and observed that the p.Trp161Cys variant protein could homodimerize as efficiently as WT SOX6 and the p.Met605Thr variant (Figure 5B). Next, we tested cell extracts in EMSA with an Acan enhancer sequence previously shown to bind SOX5 and SOX6.³⁴ The proteins with variants outside the HMG domain behaved like WT SOX6, whereas the proteins harboring the p.Met605Thr and p.Trp639Arg variants failed to bind to the DNA probe (Figure 5C).

Finally, we tested the variants for transcriptional activity by co-transfecting cells with a reporter plasmid containing an *Acan* enhancer synergistically activated by SOX9 and SOX proteins (SOX5 or SOX6)³⁴ and with SOX6 and SOX9 expression plasmids. As expected, SOX6 alone,

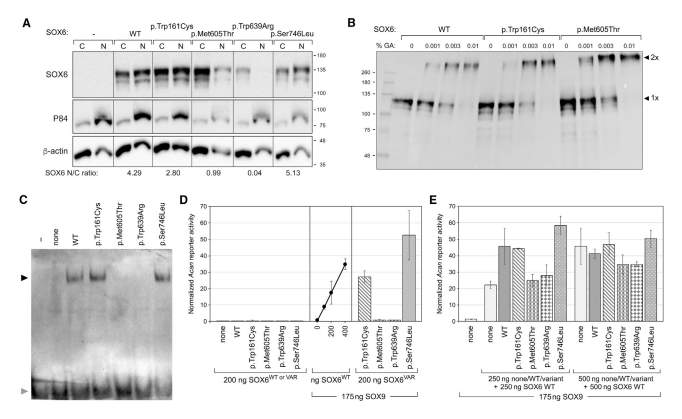


Figure 5. Functional Tests of SOX6 Missense Variants

(A) Stability and intracellular distribution of SOX6 variants. COS-1 cells were transfected with plasmids encoding 3FLAG-tagged SOX6 WT and variant proteins, as indicated. Cytoplasmic (C) and nuclear (N) extracts were tested via immunoblotting with a FLAG antibody. As expected, the P84 protein was enriched in nuclear extracts, and β -actin was enriched in cytoplasmic extracts. The distribution of SOX6 relative to β -actin in the nuclear versus cytoplasmic compartment is indicated underneath the blots. The migration of protein markers (Mr in k values) is indicated.

(B) Ability of SOX6 variants to homodimerize. Lysates of COS-1 cells, transfected as described in (A), were incubated without (0) or with 0.001%, 0.03%, or 0.01% glutaraldehyde. SOX6 was visualized via immunoblotting with a FLAG antibody. The migration of protein standards (Mr in k values) and SOX6 monomers $(1\times)$ and homodimers $(2\times)$ is indicated.

(C) Ability of SOX6 variants to bind DNA. An *Acan* enhancer probe was used in EMSA with no protein extract (-) or with extracts from COS-1 cells transfected with expression plasmids for no protein (none), SOX6 WT, or variant proteins, as indicated. SOX6-probe complexes (indicated with a black arrowhead) and free probes (indicated with a gray arrowhead) were resolved by electrophoresis.

(D) Ability of SOX6 variants to synergize with SOX9 in transactivation. HEK293 cells were transfected with an *Acan* reporter, a control reporter, and plasmids encoding no SOX (none), SOX9, SOX6 WT, or SOX6 variant proteins, as indicated. The amounts of SOX plasmids are indicated too. *Acan* reporter activities were normalized for transfection efficiency. They are presented as the mean \pm standard deviation obtained for triplicates in an experiment representative of five independent ones. The arrowhead in the middle panel points to the amount of WT SOX6 plasmid (200 ng) that was also used for variant plasmids in the right panel.

(E) Interference of SOX6 variant proteins with WT SOX6 in transactivation. HEK293 cells were transfected as described in (D) with the indicated types and amounts of plasmids encoding no protein (none) or SOX proteins. *Acan* reporter activities were normalized for transfection efficiency. They are presented as the mean \pm standard deviation obtained for triplicates in an experiment representative of four independent ones.

whether WT or mutant, and SOX9 alone hardly activated the reporter, whereas co-expression of WT SOX6 and SOX9 resulted in a SOX6 dose-dependent activation of the reporter (Figure 5D). In line with the EMSA results, the SOX6 variants located within the HMG domain were transcriptionally inactive. In contrast, the other variants were as potent as, or even more potent than, WT SOX6. Because all affected individuals in our study are heterozygous for their SOX6 variant, we next asked whether the variant proteins could have a dominant-negative influence on WT SOX6. When a dose of WT SOX6 within the linear dose-response range (250 ng plasmid) was doubled, the reported activity doubled as well. When this dose was supplemented with the same dose of p.Trp161Cys or p.Ser746-Leu variant, a similar or slightly higher reporter activity was measured, whereas the same dose of p.Met605Thr or p.Trp639Arg variant failed to increase the reporter activity. These findings were consistent with the activities of the variant proteins when tested alone (Figure 5D) and thus suggested that none of the variant proteins negatively interfered with WT SOX6. When the SOX6 proteins were tested under competitive conditions, that is, at doses in the plateau of maximum reporter activity (500 ng plasmid), none of the variant proteins prevented WT SOX6 from reaching this plateau, again suggesting a lack of a dominant-negative effect of the variants.

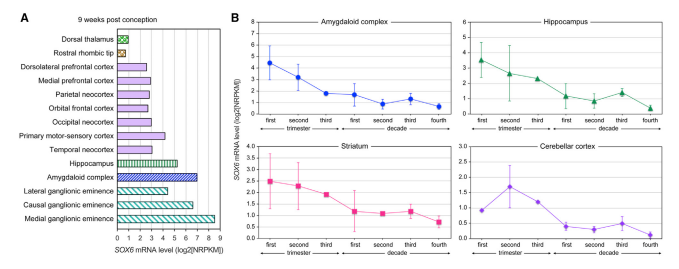


Figure 6. Analysis of SOX6 Expression in the Human Brain

(A) SOX6 RNA expression measured by RNA-seq in various regions of the developing brain of a representative human fetus 9 weeks after conception.

(B) *SOX6* RNA expression measured by RNA-seq in the amygdaloid complex, hippocampus, striatum, and cerebellar cortex of multiple individuals whose ages are within the range spanning the three trimesters of gestation *in utero* and the first four decades of life. Data are presented as averages with standard deviation for each age category (n = 1-10 per group). The four brain structures were selected because their *SOX6* RNA expression is higher than that of other structures.

Taken together, these functional assays indicated that the two missense variants in the HMG domain abolished the ability of SOX6 to function as a transcription factor. The two missense variants located outside this domain exhibited SOX6 WT activities in our assays, but we cannot rule out the possibility that they could impair critical activities of SOX6 *in vivo* and might thus be genuinely pathogenic.

SOX6 Is Highly Expressed in the Developing Human Brain

Whole-transcriptome profiling data deposited in the Brain-Span Atlas of the Developing Human Brain provided information on where and when SOX6 is expressed in the brain of human fetuses, children, and adults. At 9 weeks after conception, fetuses were found to express SOX6 in many prospective brain structures (Figure 6A). SOX6 RNA expression was highest in the ganglionic eminence, the amygdaloid complex, and the hippocampus, all of which have central roles in brain development. Expression lower than that measured in the aforementioned regions were measured in sub-regions of the cortex and neocortex. The expression of SOX6 declined in all brain structures in the final stages of gestation and in the neonatal period, such that it was already as low in infants as in adults (Figure 6B). These data thus provide support to the notion that SOX6 has important roles in several regions of the developing human brain.

Discussion

We reported here on 19 individuals, from 17 families, who exhibited genetic variants predicted in most cases to cause *SOX6* haploinsufficiency. All subjects shared core clinical features of a neurodevelopmental syndrome that included developmental delay and intellectual disability. Inconstant features, such as ADHD (11 affected individuals), mild facial dysmorphism (9 affected individuals), craniosynostosis (3 affected individuals), and osteochondromas (3 affected individuals), were also shared by several subjects. This disease thus expands the currently known list of SOXopathies, i.e., developmental disorders due to pathogenic variants in SOX genes.

The spectrum of SOX6 variants carried by the affected individuals in our study ranged from a balanced chromosomal translocation to partial or complete gene deletions and to nonsense, missense, and other SNVs. Although no individual affected by a SOX6 SNV was reported prior to our study, two individuals were described as having a SOX6 CNV. The first one was a 4-year-old girl with global developmental delay, spinal cord syrinx, and recurrent episodes of parkinsonian symptoms, including gait instability, tremors, and dysarthria.³⁹ Because molecular karyotyping showed a *de novo* 2.36 Mb deletion including SOX6 and several other genes, no conclusion could be drawn regarding the implication of the SOX6 deletion in the clinical phenotype. The second individual was a 15-year-old boy with significantly delayed speech development and ADHD.⁴⁰ This boy also had generalized dystonic and frequent athetoid movements of the arms, trunk, and neck. His gait was severely impaired secondarily to frequent dystonic postures. A whole-genome SNP array revealed a de novo 84 kb deletion encompassing SOX6 exons 14-16, which encode the HMG domain and C terminus of SOX6. In combination with the affected individuals reported in our study, these additional subjects link SOX6 variants to a complex neurodevelopmental syndrome.

The linking of SOX6 variants with a neurodevelopmental syndrome fits with the evidence that SOX6 is dynamically expressed in the human developing brain and that its mouse ortholog is also expressed in this structure and plays key roles in creating neuronal diversity.^{19,20,41} Sox6 is specifically expressed in neuronal progenitors in the developing dorsal telencephalon and induces differentiation of the cells.¹⁹ This function might explain the neurodevelopmental disorders observed in the affected individuals in our study. In contrast to the individual with an 84 kb deletion, none of the affected individuals in our study had movement disorders. SOX6 is also strongly expressed in the medial ganglionic eminence, where it is necessary for the normal positioning and maturation of cortical interneurons. As a consequence, specific removal of Sox6 from these cells results in a severe epileptic encephalopathy in mice.42 None of the affected individuals reported here or previously, however, presented with epilepsy. A possible explanation is that the human individuals were heterozygous for the SOX6 alteration, whereas the phenotype was seen in mice only when Sox6 was inactivated homozygously.

Besides neurogenesis, chondrogenesis is another developmental process critically dependent upon Sox6. The gene is co-expressed with Sox5 in chondrocytes, and the two genes act largely in redundancy to promote chondrocyte differentiation.¹²⁻¹⁴ SOX5 and SOX6 form homodimers and heterodimers through their coiled-coil domains and cooperatively bind with SOX9 on enhancers driving hundreds of cartilage-specific genes. They thereby potentiate the ability of SOX9, which is a master chondrogenic factor, to transactivate these genes.⁴³ Sox5-null and Sox6-null mice are born with discrete skeletal malformations, and fetuses null for both Sox5 and Sox6 die in utero with severely underdeveloped cartilage structures and cartilage-derived bones. These chondrogenic roles of SOX5 and SOX6 could explain the mild facial dysmorphism and the shortness of fingers and extremities noted in some of the affected individuals in our study, as well as in some individuals with LAMSHF syndrome, who are heterozygous for *SOX5* pathogenic variants.⁷ Dysmorphic features have not been described in Sox5- and Sox6-heterozygous-null mice but cannot be excluded because detailed morphometric measurements have not been performed.

Three unrelated affected individuals reported here presented with craniosynostosis. For two of them (UK-1 and UK-2), the sequences of genes known to be involved in craniosynostosis were analyzed in depth, but no variant likely to be pathogenic was identified. Of note, a 5-year-old boy with mild developmental speech delay and craniosynostosis was previously described.⁴⁴ This boy exhibited brachycephaly, proptosis, midfacial hypoplasia, and low-set ears. He carried a *de novo* balanced translocation, 46,XY,t(9;11)(q33;p15), whose breakpoint on chromosome 11 disrupted *SOX6. Sox5* and *Sox6* single- and double-mutant mice were not described to have defects in the intramembranous bones that form the skull vault, but such defects and the expression of *Sox5* and *Sox6* in skull progenitor cells and osteoblasts might have been overlooked in view of the strong expression of both genes in chondrocytes and the severity of cartilage and endo-chondral bone underdevelopment in mutant mice.¹³

Among the affected individuals in our study, two brothers and one unrelated girl presented with diffuse osteochondromas. The former each had a SOX6 deletion, and the latter had a missense variant in the HMG domain. Thus, these individuals did not have SOX6 variant types that differed from those of other affected individuals in study. Osteochondromas are benign tumors our developing as ectopic bone and cartilage, most often in the perichondrium adjacent to cartilage growth plates. Because SOX6 promotes essential steps of the chondrocyte differentiation pathway, the finding that SOX6 haploinsufficiency is associated with such tumors is intriguing; this calls for researchers to undertake future investigations to determine the importance of SOX6 in preventing ectopic cartilage and bone formation. At present, a possible explanation might come from evidence that SOX6 is a tumor suppressor in numerous types of cancer, including osteosarcoma, the most common type of malignant bone tumor (MIM: 259500).^{25,27,45-47}

Most of the affected individuals in our study had SOX6 variants that surely inactivated one SOX6 allele and thus caused a disease that revealed SOX6 haploinsufficiency. These variants were microdeletions and frameshift and nonsense variants. That missense variants in the HMG domain inactivated the protein made from the SOX6 carrier allele was supported both by our in vitro functional assays and by evidence that many variants in the HMG domain, including those detected in affected individuals in our study, have been shown in other SOX genes to cause SOXopathies. One should, however, not deduce that any SOX variant in the HMG domain is pathogenic. gnomAD indeed contains multiple control individuals with variants in the HMG domain, and we showed in the case of SOX4 that these variants were retaining functions in vitro.¹⁰ The identification of additional individuals with variants causing disease along with expansion of gnomAD should help future researchers establish algorithms to accurately make pathogenicity diagnoses for newly affected individuals.

Two of the affected individuals in our study carried a *SOX6* missense variant outside the HMG domain. Although these two individuals appeared to have more severe intellectual disability than all others and *in silico* tools based on amino acid structure and protein sequence predicted pathogenicity, *in vitro* functional assays did not consolidate this argument. A possible explanation is that these assays, which assessed the ability of SOX6 to synergize with SOX9 to activate a chondrocyte-specific enhancer, were not suitable for assessing the impact of the variants on SOX6 function or regulation in neurogenesis and other processes *in vivo*. A few individuals with LAMSHF disease whose variants were located outside the HMG domain (of SOX5),

were not present in gnomAD individuals, and were fully active in the functional assay in vitro used in the present study were also reported.⁷ The pathogenicity of such variants is further supported by the extremely high degree of conservation observed between SOX5 and SOX6 throughout the proteins and among vertebrate orthologs. More studies are thus needed for determining how such variants might critically affect SOX5 and SOX6 function or regulation and thereby cause diseases. Of note, one of the two patients also carried a hemizygous MECP2 (MIM: 300005) variant that was most likely pathogenic. Variants in this gene have been associated with X-linked autism susceptibility (MIM: 300496), mental retardation (MIM: 300260), and Rett syndrome (MIM: 312750).⁴⁸ It is highly likely that this variant has a major contribution to the clinical features of this patient.

In conclusion, the findings from this study concur that *SOX6* haploinsufficiency leads to a specific form of neurodevelopmental SOXopathy characterized by mild to severe intellectual disability and inconstantly associated with skeletal anomalies, such as mild facial dysmorphism, craniosynostosis, and osteochondromas.

Supplemental Data

Supplemental Data can be found online at https://doi.org/10. 1016/j.ajhg.2020.04.015.

Acknowledgments

We thank all individuals and their families who contributed to this study. A.O.M.W. was supported by the National Institute for Health Research (NIHR) Oxford Biomedical Research Centre Programme. This research was made possible in part through access to the data and findings generated by the 100,000 Genomes Project. The 100,000 Genomes Project is managed by Genomics England Limited (a wholly owned company of the Department of Health and Social Care) and funded by the National Institute for Health Research and the National Health Service (NHS) England. The Wellcome Trust, Cancer Research UK, and the Medical Research Council have also funded research infrastructure. The 100,000 Genomes Project uses data provided by affected individuals and collected by the NHS as part of their care and support. This work was also funded by the Children's Hospital of Philadelphia.

Declaration of Interests

A.B., C.F., L.B.H., and P.R. are employees of GeneDx. All other authors declare no competing interests.

Received: March 13, 2020 Accepted: April 24, 2020 Published: May 21, 2020

Web Resources

1000 Genomes Project, http://www.1000genomes.org/ ClinVar, https://www.ncbi.nlm.nih.gov/clinvar/ DECIPHER, https://decipher.sanger.ac.uk/

gnomAD Browser, http://gnomad.broadinstitute.org/ GenBank, https://www.ncbi.nlm.nih.gov/genbank/

Genomics England 100,000 Genomes Project, https://www.genomicsengland.co.uk/

HOPE, http://www.cmbi.ru.nl/hope/

NCBI, https://www.ncbi.nlm.nih.gov/

OMIM, http://www.omim.org/

PEP-FOLD-3, http://bioserv.rpbs.univ-paris-diderot.fr/services/ PEP-FOLD3/

SWISS-MODEL, https://swissmodel.expasy.org/

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The American Journal of Human Genetics, Volume 106

Supplemental Data

De Novo SOX6 Variants Cause a Neurodevelopmental

Syndrome Associated with ADHD, Craniosynostosis,

and Osteochondromas

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SUPPLEMENTAL MATERIALS

Supplemental Tables

Subject Numbers	Internal IDs	Referring Centers	Methods of diagnostic			
#1	PIT-1	Pittsburgh	Agilent 180K Human Genome CGH + SNP ISCA			
#2	PIT-2 Pittsburgh		Agilent 180K Human Genome CGH + SNP ISCA			
#3	PIT-3 Pittsburgh		Agilent 180K Human Genome CGH + SNP ISCA			
#4	4 CHUN-1 Nantes		Agilent 60K ISCA v.2			
#5	CHLM-1	Le Mans	Agilent 60K ISCA v.2			
#6	LJU-1	Ljubljana	Agilent 180K			
#7	IHG-1	Essen	Affy CytoScan HD			
#8	CHUP-1	Poitiers	Agilent 105K			
#9	UK-1	Nottingham	WGS (Genomics England 100,000 Genomes Project)			
#10	UK-2	Nottingham	WGS (Genomics England 100,000 Genomes Project)			
#11	CHUN-2	Nantes	WES; Agilent SureSelect Human All Exome 50 Mb Kit			
#12	GDX-3	Sickkids	WES; Agilent SureSelect Clinical Research Exome Kit			
#13	GDX-1	UCLA	WES; Agilent SureSelect Clinical Research Exome Kit			
#14	GDX-2	Duke	WES; Agilent SureSelect Clinical Research Exome Kit			
#15	PS-1	Pitié	WES; Medexome Nimblegen 47 Mb Kit			
#16	CHOP-1	Philadelphia	WES; Agilent SureSelectXT Clinical Exome version 1			
#17	LEUV-1	Leuven	WES; Medexome Nimblegen, custom EZ choice XL V4			
#18	LEID-1	Leiden	WES; Agilent SureSelect V5			
#19	CHUG-1	Grenoble	WGS			

 Table S1. Referring Centers and Methods of Diagnostic for the 19 Affected Individuals.

SRY	CAA37790.1
SOX1	NP_005977.2
SOX2	NP_003097.1
SOX3	NP_005625.2
SOX4	AAH72668.1
SOX5	AAH60773.1
SOX6	AAK26115.1
SOX7	CAC84226.1
SOX8	NP_055402.2
SOX9	CAA86598.1
SOX10	CAG38808.1
SOX11	BAA88122.1
SOX12	AAH67361.1
SOX13	AAD50120.1
SOX14	AAC95380.1
SOX15	AAH72003.1
SOX17	BAB83867.1
SOX18	BAA94874.1
SOX21	AAC95381.1
SOX30	BAA37146.1

Table S2. Accession Numbers for the Human SOX Sequences Aligned in Figure 4.

Vertebrate species	Scientific name	Reference sequence
Human	Homo sapiens	NM_033326.3
Mouse	Mus musculus	AAC52263.1
Dog	Canis lupus familiaris	XP_022263597.1
Cow	Bos taurus	NP_001178347.1
Armadillo	Dasypus novemcinctus	XP_023438566.1
Platypus	Ornithorhynchus anatinus	XP_028916408.1
Chicken	Gallus gallus	NP_001305380.1
Western clawed frog	Xenopus tropicalis	XP_017945061.1
Zebrafish	Danio rerio	NP_001116481.1

Table S3. Accession numbers of SOX6 ortholog sequences aligned in Figure S1.

Variant	Forward primer	Reverse primer
WT	CAAAAGAT TGG AAAGGAAAAAATGGA	TTTCCTT <u>CCA</u> ATCTTTTGAAAGTAG
p.Trp161Cys	CAAAAGAT TGC AAGGAAAAAATGGA	TTTCCTT <u>GCA</u> ATCTTTTGAAAGTAG
WT	CGACCA ATG AATGCGTTCATGGTTTG	CGCATT <u>CAT</u> TGGTCGCTTGATGTGG
p.Met605Thr	CGACCA ACG AATGCGTTCATGGTTTG	CGCATT <u>CGT</u> TGGTCGCTTGATGTGG
WT	AGGATCTCGC TGG AAATCCATGTCCAACC	ATGGATTT <u>CCA</u> GCGAGATCCTAAGATTTTGC
p.Trp639Arg	AGGATCTCGC CGG AAATCCATGTCCAACC	ATGGATTT <u>CCG</u> GCGAGATCCTAAGATTTTGC
WT	GCACC <u>TCT</u> GCCAGCCCCGAGCCCA	GGCTGGC AGA GGTGCTAGAGCAGT
p.Ser746Leu	GCACC <u>TTG</u> GCCAGCCCCGAGCCCA	GGCTGGC <u>CAA</u> GGTGCTAGAGCAGT

Table S4. Primers Used to Generate Expression Plasmids for SOX6 Variants. Wild-type (WT) sequences are shown as references. Codons affected by variants are in bold font and underlined.

Affected individual	Body part	Region of the body part exhibiting endochondroma(s)
PIT-1	right arm	5 th proximal phalanx distal 4 th middle phalanx distal metaphysis of the 3 rd metacarpal bone trapezoid bone
	left arm	proximal humerus distal 4 th metacarpal bone metaphysis proximal end of 1 st metacarpal bone
	left leg	proximal end of 1 st metatarsal bone
	spine	right 12 th vertebral body (transverse process)
PIT-2	right leg	distal femur metaphysis
	left leg	proximal fibula proximal phalanx of 4 th toe
CHOP-1	right arm	proximal humerus*
	left arm	proximal humerus
	right leg	distal femur**
	left leg	distal tibia
	thorax	right rib*

Table S5. List of Endochondromas Found in Affected Individuals. *Osteochondromas no longerpalpable, but still visible on X-rays two years later. **Osteochondromas no longer palpable and no longervisible on X-rays two years later.

Subject	Internal	Genomic variant	Telomeric	Centromeric	Size of the
number	identifier	(NM_033326.3)	boundary	boundary	deletion
#1	PIT-1	del ex. 1 to 2	16,358,927	16,497,834	138,907 bp
#2	PIT-2	del ex. 1 to 2	16,358,927	16,497,834	138,907 bp
#3	PIT-3	del ex. 1 to 2	16,358,927	16,497,834	138,907 bp
#4	CHUN-1	del ex. 1 to 4	16,238,166	16,554,120	315,954 bp
#5	CHLM-1	del ex. 2 to 3	16,309,520	16,419,313	109,793 bp
#6	LJU-1	del ex. 2 to 13	16,023,626	16,446,051	422,425 bp
#7	IHG-1	del ex. 2 to 12	16,049,440	16,399,572	350,132 bp
#8	CHUP-1	del ex. 5 to 7	16,128,647	16,239,865	111,218 bp

Table S6. Location, Boundaries and Size of the SNVs Affecting *SOX6* in Eight Study Subjects (hg19).

Supplemental Figures

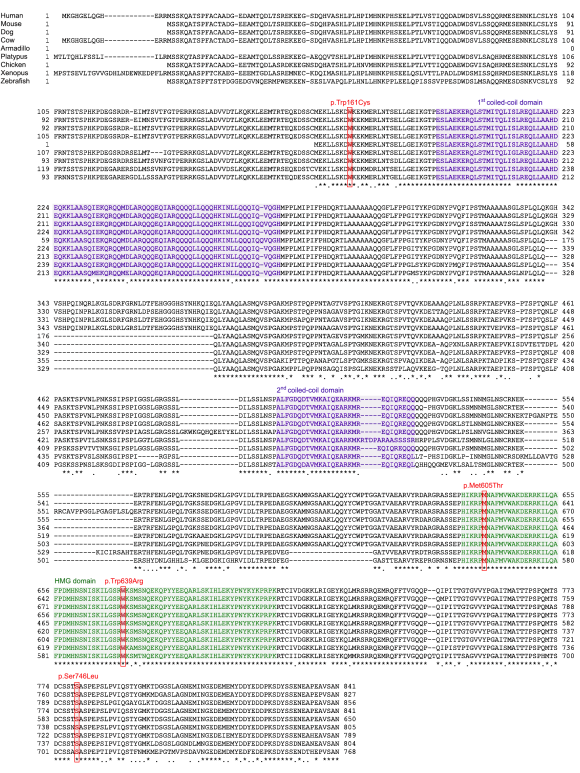


Figure S1. ClustalW Alignment of SOX6 Vertebrate Orthologs. The known functional domains of SOX6 are colored and labeled. Fully conserved residues are marked with asterisks and semi-conserved residues with dots underneath the sequences. Residues altered in study subjects are indicated in red.

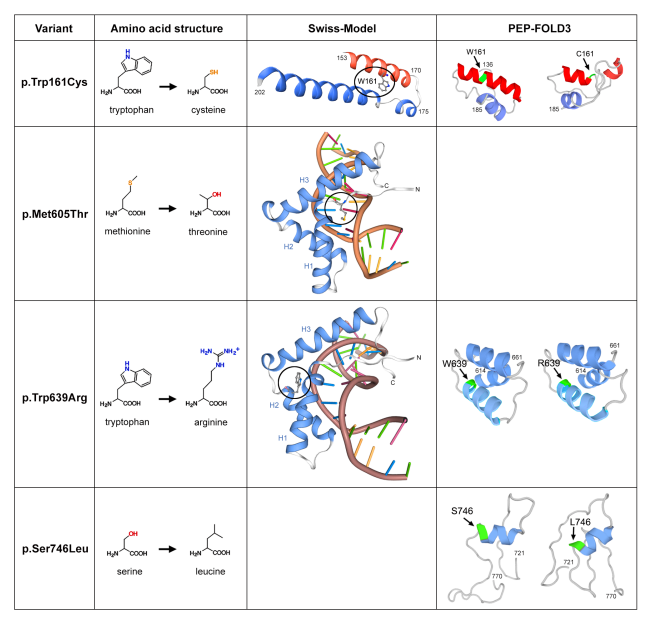


Figure S2. Prediction of Structural Changes Imposed by Missense Variants. The first column identifies the missense variants detected in four study subjects. The second column shows schematics of wild-type and missense variant residues. The third column shows structural models proposed by Swiss-Model for the SOX6 regions containing three missense variants. For p.Met605Thr and p.Trp639Arg, the models correspond to the SOX5 HMG domain (with the α -helices H1, H2, and H3 in blue, and flanking sequences in grey) wrapped around the DNA helix (backbone in brown and nucleotide pairs colored). Altered amino acids are circled in black and their side chains represented. The last column shows structural models predicted by PEP-FOLD3 for the wild-type and mutant SOX6 regions that encompass missense variants. The wild-type and variant residues are indicated, as are the boundaries of the sequences used for modeling.

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