



SHORT REPORT

Mutations in *TMEM231* cause Joubert syndrome in French Canadians

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ABSTRACT

Background Joubert syndrome (JBTS) is a predominantly autosomal recessive disorder characterised by a distinctive midhindbrain malformation, oculomotor apraxia, breathing abnormalities and developmental delay. JBTS is genetically heterogeneous, involving genes required for formation and function of non-motile cilia. Here we investigate the genetic basis of JBTS in 12 French-Canadian (FC) individuals.

Methods and results Exome sequencing in all subjects showed that six of them carried rare compound heterozygous mutations in *CC2D2A* or *C5ORF42*, known JBTS genes. In addition, three individuals (two families) were compound heterozygous for the same rare mutations in *TMEM231* (c.12T>A[p.Tyr4*]; c.625G>A[p.Asp209Asn]). All three subjects showed a severe neurological phenotype and variable presence of polydactyly, retinopathy and renal cysts. These mutations were not detected among 385 FC controls. *TMEM231* has been previously shown to localise to the ciliary transition zone, and to interact with several JBTS gene products in a complex involved in the formation of the diffusion barrier between the cilia and plasma membrane. siRNA knockdown of *TMEM231* was also shown to affect barrier integrity, resulting in a reduction of cilia formation and ciliary localisation of signalling receptors.

Conclusions Our data suggest that mutations in *TMEM231* cause JBTS, reinforcing the relationship between this condition and the disruption of the barrier at the ciliary transition zone.

Joubert syndrome (JBTS [MIM213300]) is a predominantly autosomal recessive disorder characterised by oculomotor apraxia, abnormal breathing, ataxia and variable developmental delay or intellectual impairment (reviewed in Sattar *et al.*).¹ A cardinal sign of JBTS is the presence of a complex midhindbrain malformation consisting of hypoplasia of the cerebellar vermis, abnormally deepened interpeduncular fossa at the level of the upper pons, and elongated and thickened superior cerebellar peduncles. This malformation takes the appearance of a molar tooth on MRI. Extraneurological manifestations, including retinopathy, renal cysts and polydactyly, are present in a subset of affected individuals. JBTS is genetically heterogeneous, with 17 genes described to date,^{1–13} all of which appear

to play a role in the development and/or function of non-motile cilia.

There is a high prevalence of JBTS in the French-Canadian (FC) population of the Lower Saint-Lawrence region of Quebec. We recently performed exome sequencing in 15 individuals (11 families) with JBTS from that region and found that mutations in *C5ORF42* explain JBTS in nine individuals (seven families).¹² In addition, we identified pathogenic compound heterozygous mutations in *CC2D2A*, a previously known JBTS gene, in two affected individuals from two different families. The genetic basis of JBTS remained unexplained in four individuals (two families) from this initial study. Here, we follow-up on our previous investigation by performing exome sequencing in eight additional individuals with JBTS (six unrelated families) originating from other regions of Quebec.

The six probands had a molar tooth sign on imaging, and variable expression of the classical JBTS features. The two additional individuals are the uncle (II-4) and aunt (II-6) of subject IV-1 in family 385/447. These individuals were considered to have a variable expression of JBTS as they both had oculomotor apraxia and, additionally, II-4 had gait ataxia and a history of developmental delay. Brain MRI was normal in II-6 (see online supplementary figure S1, A-B) but was not done in II-4. Informed consent was obtained from each individual or legal guardian. This study was approved by our institutional ethics committee. Genomic DNA from each sample was captured with the Agilent SureSelect 50 Mb exome capture oligonucleotide library, and the captured DNA was sequenced with paired-end 100 bp reads on Illumina HiSeq 2000 resulting in an average of 12.7 gigabases (Gb) of raw sequence for each sample. Data were analysed as previously described.¹⁴ After removing putative PCR-generated duplicate reads using Picard (V1.48), we aligned reads to human genome assembly hg19 using a Burroughs-Wheeler algorithm (BWA V0.5.9). Median read depth of bases in consensus coding sequence (CCDS) exons was 99 (determined with Broad Institute Genome Analysis Toolkit V1.0.4418).¹⁵ On average, 87% (±2.0%) of bases in CCDS exons were covered by at least 20 reads. We called sequence variants using Samtools (V0.1.17) mpileup and varFilter, and required at least three variant reads as well as

≥20% variant reads for each called position, with Phred-like quality scores of at least 20 for single nucleotide variants (SNVs) and at least 50 for small insertions or deletions (indels). We used Annovar and custom scripts to annotate variants according to the type of mutation, occurrence in the Single Nucleotide Polymorphism database (dbSNP), Sorting Intolerant from Tolerant (SIFT) score, 1000 Genomes allele frequency, and National Heart, Lung and Blood Institute (NHLBI) exomes allele frequency.¹⁶ To identify potentially pathogenic variants we filtered out (1) synonymous variants or intronic variants other than those affecting the consensus splice sites; (2) variants seen in more than two of 416 exomes from patients with rare, monogenic diseases unrelated to JBTS that were sequenced at the McGill University and Genome Quebec Innovation Centre and (3) variants with a frequency greater than 0.5% in either the 1000 genomes or NHLBI exome datasets.

We first examined the eight exome datasets to look for rare variants in the 17 known JBTS genes (*INPP5E*[MIM613037], *TMEM216*[MIM613277], *AHI1*[MIM608894], *NPHP1*[MIM607100], *CEP290*[MIM610142], *TMEM67*[MIM609884], *RPGRIP1L*[MIM610937], *ARL13B*[MIM608922], *CC2D2A*[MIM612013], *CXORF5*[MIM300170], *KIF7*[MIM611254], *TCTN1*[MIM609863], *TCTN2*[MIM613885], *TMEM237*[MIM614424], *CEP41*[MIM610523], *TMEM138*[MIM614459], *C5ORF42*[MIM614571],^{1–13} as well as in the JBTS candidate gene *TTC21B*(MIM612014).¹⁷ Five individuals from three families (II-1 from family 379, II-4, II-6 and IV-1 from family 385/447, and II-1 from family 492 online supplementary figure S2) were each found to carry two rare heterozygous variants in *CC2D2A*(NM_001080522.2). One in-frame amino acid deletion (c.3450_3452del[p.Val1151del]) and four different missense variants (c.3376G>A[p.Glu1126Lys], c.4559A>G[p.Asn1520Ser], c.4667A>T[p.Asp1556Val], c.4702T>C[p.Tyr1568His]) were identified, two of which, c.3376G>A(p.Glu1126Lys) and c.4667A>T(p.Asp1556Val), were identified previously in FC individuals with JBTS.¹² The novel mutations c.4559A>G(p.Asn1520Ser) and c.4702T>C(p.Tyr1568His) are predicted to be damaging (by SIFT, Polyphen-2 and Mutation Taster) and neither variant has been reported in the Exome Variant Server (EVS; NHLBI GO Exome Sequencing Project), dbSNP135 or 1000 Genome datasets. These five *CC2D2A* mutations cluster in either the C2 domain (amino acids 1062–1174) or the C-terminal part of the protein, as do most missenses that cause JBTS.¹⁸ Segregation analysis revealed that all the affected individuals, but none of their unaffected relatives, were compound heterozygous for the mutations (see online supplementary figure S1). We conclude that these mutations are pathogenic and responsible for JBTS in these five individuals.

We also identified a frameshift mutation (c.8257_8258insA [p.Lys2753fs]) and a splice-site mutation (c.7400+1G>A) in *C5ORF42*(NM_023073.3) in individual II-2 from family 551. Sanger sequencing showed that the proband is compound heterozygous for these mutations. The splice site (c.7400+1G>A) mutation has been previously identified in patients with JBTS and shown to result in skipping of exon 35 and the creation of a premature stop codon while c.8257_8258insA(p.Lys2753fs), which is novel, is predicted to truncate *C5ORF42* in the middle of its sequence, close to where other truncating mutations have been previously identified in JBTS patients.¹² Both *C5ORF42* mutations are thus considered pathogenic in this individual. Table 1 summarises the genotypes and phenotypes of these patients with mutations in *CC2D2A* and *C5ORF42*, as well as those of FC patients previously described with mutations in these genes. Individuals in our cohort with mutations

in *CC2D2A* do not have any extraneural manifestations, and appear to have a milder phenotype, with all affected individuals walking independently before the age of 4 years, and intelligence ranging from normal to mild intellectual impairment. Individuals with mutations in *C5ORF42* have a more variable phenotype. They have borderline to moderate cognitive impairment and variable age at walking, ranging between 30 months and 8 years. Some patients also showed limb abnormalities, including one individual with combined pre- and postaxial polydactyly, an unusual finding in JBTS, which is typically associated with postaxial polydactyly.

We then combined the exome data of the two remaining individuals with unexplained JBTS and the exome data of four individuals with unexplained JBTS from our previous study,¹² making a total of six individuals from four different families. We analysed the data by looking for protein-coding genes that contained homozygous or multiple heterozygous variants in each affected individual. For multiplex families, we only considered genes with the same variants in the affected siblings (see online supplementary tables S1 and S2). Only one gene, *TMEM231*, harboured multiple rare mutations in more than one family. Three JBTS individuals from 2 families (II-1 and II-2 from family 387, and II-1 from family 483) harboured the same two variants in *TMEM231*(NM_001077418.1): c.12T>A (p.Tyr4*) and c.625G>A(p.Asp209Asn). Sanger sequencing showed that all affected individuals were compound heterozygous for these variants (figure 1A). The c.12T>A(p.Tyr4*) mutation targets exon 1 of the canonical isoform of *TMEM231* (NM_001077418.1; ENST00000258173), as well as the two other predicted protein-coding isoforms reported in the Ensemble Genome Browser. In ENST00000565067, it leads to the same nonsense change (p.Tyr4*), while in the longer isoform ENST00000398114, it abolishes the translation initiation methionine (c.2T>A[p.Met1?]), which would likely prevent translation of this isoform due to the absence of any other in-frame methionine in exons 1 and 2. The c.625G>A(p.Asp209Asn) causes the same amino acid change in the different *TMEM231* predicted isoforms (figure 1C). It affects a highly conserved amino acid (figure 1D), and is predicted to be damaging by Polyphen-2 and Mutation Taster but not by SIFT. Both p.Tyr4* and p.Asp209N are extremely rare. Among the 416 in-house exomes, the c.12T>A(p.Tyr4*) was not found, and the c.625G>A(p.Asp209Asn) variant was seen in the heterozygous state in one FC individual. No additional *TMEM231* coding/splicing variants were present in this individual's exome. To determine the carrier rate of c.625G>A and c.12T>A, we genotyped 385 healthy FC controls by Sanger sequencing, but did not find any carriers of either of these mutations, indicating that they are very rare. Only p.Asp209Asn is reported in the heterozygous state in the 1000 genomes and EVS, but at a very low frequency (0.01%), while p.Tyr4* is not reported in any of these public single nucleotide polymorphism (SNP) databases. Furthermore, no truncating mutations in *TMEM231* were seen in 416 control exomes of patients with other rare diseases, and only one other truncating variant (stopgain SNV) is reported in EVS, at a frequency of 0.04%. For the three individuals with compound heterozygous *TMEM231* mutations, we examined all SNV genotypes in regions surrounding the two mutations. This revealed a region of shared genotypes (two shared haplotypes) extending over at least 1.7 Mb, suggesting the existence of founder effects (see online supplementary table S3).

The three individuals with mutations in *TMEM231* are among the most severely affected in our French–Canadian JBTS

Table 1 Genotypes and phenotypes of French Canadian individuals with JBTS

Genotypes	Srouf <i>et al</i> ¹²												This study								
	406/301			394		474	480	489	479	468	473	484	385/447			379	492	551	387		483
	IV-1	IV-2	IV-3	II-1	II-2	II-1	II-1	II-1	II-1	II-1	II-2	II-1	II-4	II-6	IV-1	II-1	II-1	II-2	II-1	II-2	II-1
<i>C5ORF42</i>																					
c.4006C>T(p.Arg1336Trp)	+	–	–	+	+	+	+	–	+	+	–	–	–	–	–	–	–	–	–	–	–
c.7400+1G>A	+	+	+	+	+	+	+	–	–	–	–	–	–	–	–	–	–	+	–	–	–
c.6407del(p.Pro2136Hisfs*31)	–	–	–	–	–	+	–	–	–	–	–	–	–	–	–	–	–	–	–	–	–
c.7477C>T(p.Arg2493*)	–	–	–	–	–	–	–	+	–	–	–	–	–	–	–	–	–	–	–	–	–
c.4804C>T(p.Arg1602*)	–	–	–	–	–	–	–	–	+	–	–	–	–	–	–	–	–	–	–	–	–
c.7957+288G>A; c.4690G>A(p.Ala1564Thr)	–	+	+	–	–	–	–	+	–	+	–	–	–	–	–	–	–	–	–	–	–
c.8257_8258insA(p.K2753fs)	–	–	–	–	–	–	–	–	–	–	–	–	–	–	–	–	–	+	–	–	–
<i>CC2D2A</i>																					
c.4667A>T(p.Asp1556Val)	–	–	–	–	–	–	–	–	–	–	+	+	–	–	+	–	+	–	–	–	–
c.3376G>A(p.Glu1126Lys)	–	–	–	–	–	–	–	–	–	–	+	+	+	+	+	+	–	–	–	–	–
c.4559A>G(p.Asn1520Ser)	–	–	–	–	–	–	–	–	–	–	–	–	+	+	–	–	–	–	–	–	–
c.4702T>C(p.Tyr1568His)	–	–	–	–	–	–	–	–	–	–	–	–	–	–	–	+	–	–	–	–	–
c.3450_3452del(p.Val1151del)	–	–	–	–	–	–	–	–	–	–	–	–	–	–	–	–	+	–	–	–	–
<i>TMEM231</i>																					
c.12T>A(p.Tyr4*)	–	–	–	–	–	–	–	–	–	–	–	–	–	–	–	–	–	–	+	+	+
c.625G>A(p.Asp209Asn)	–	–	–	–	–	–	–	–	–	–	–	–	–	–	–	–	–	–	+	+	+
Age (years)	8	1.5	3	52	45	4	10	7	13	31	3	12	62	53	5	10	5	16	14	9	4
Developmental delay	+	+	+	+	+	+	+	+	+	+	Mild	Mild	+	–	+	+	+	+	+	+	+
Age at walking	Walks with aid	Not amb	NA	NA walks	3	Not amb	8	3.5	7	2.5	1.5	1.5	4	1	2	4	2.5	7	Not amb	Not amb	Not amb
Oculomotor apraxia	–	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	–	+	+	+
Breathing abnormality	+	+	+	+	+	+	+	+	–	–	–	–	–	–	–	+	–	+	+	+	+
Limb abnormality†	–	+	–	–	–	+	–	–	–	–	–	–	–	–	–	–	–	–	–	+	+
Brain MRI	MTS	MTS	MTS	ND	MTS	MTS	MTS	MTS	MTS	MTS	MTS	MTS	NA	N	MTS	MTS	MTS	MTS	MTS	MTS	MTS
Retinal involvement‡	–(f)	–(e)	–(e)	–(h)	–(h)	–(f)	–(e)	–(e)	–(f)	–(h)	–(e)	–(e)	–(f)	–(e)	–(e)	–(e)	–(e)	–(f)	–(f)	+(f)	+(e)
Renal involvement§	–(us)	–(us)	–(us)	–(h)	–(h)	–(us)	–(us)	–(us)	–(us)	–(h)	–(us)	–(h)	–(h)	–(h)	–(us)	–(us)	–(h)	–(us)	–(us)	+(us)	+(us)

The nucleotide and amino acid positions for *CC2D2A* are based on reference sequence #NM_001080522.2, for *TMEM231* on reference sequence #NM_001077418.1, and for *C5ORF42* on reference sequence #NM_023073.3 except for c.G4690A/p.A1564T that is based on ENSEMBLE transcript ID #ENST00000509849.

†IV-2 from family 406/301 has a 3–4 syndactyly in the left hand, II-1 from family 474 has pre- and postaxial polydactyly of the four limbs, and II-2 from family 387 and II-1 from family 483 have postaxial polydactyly and 4-5-6 syndactyly of the right foot.

‡Retinal involvement was determined by electroretinogram (erg), funduscopy (f) or history (h).

§Renal involvement was determined by renal ultrasound (us) or history (h). Individuals II-2 from family 387 and II-1 from family 483 have renal cysts with normal renal function.

MTS, Molar tooth sign; N, normal; NA, not available; Not amb, Not ambulatory.

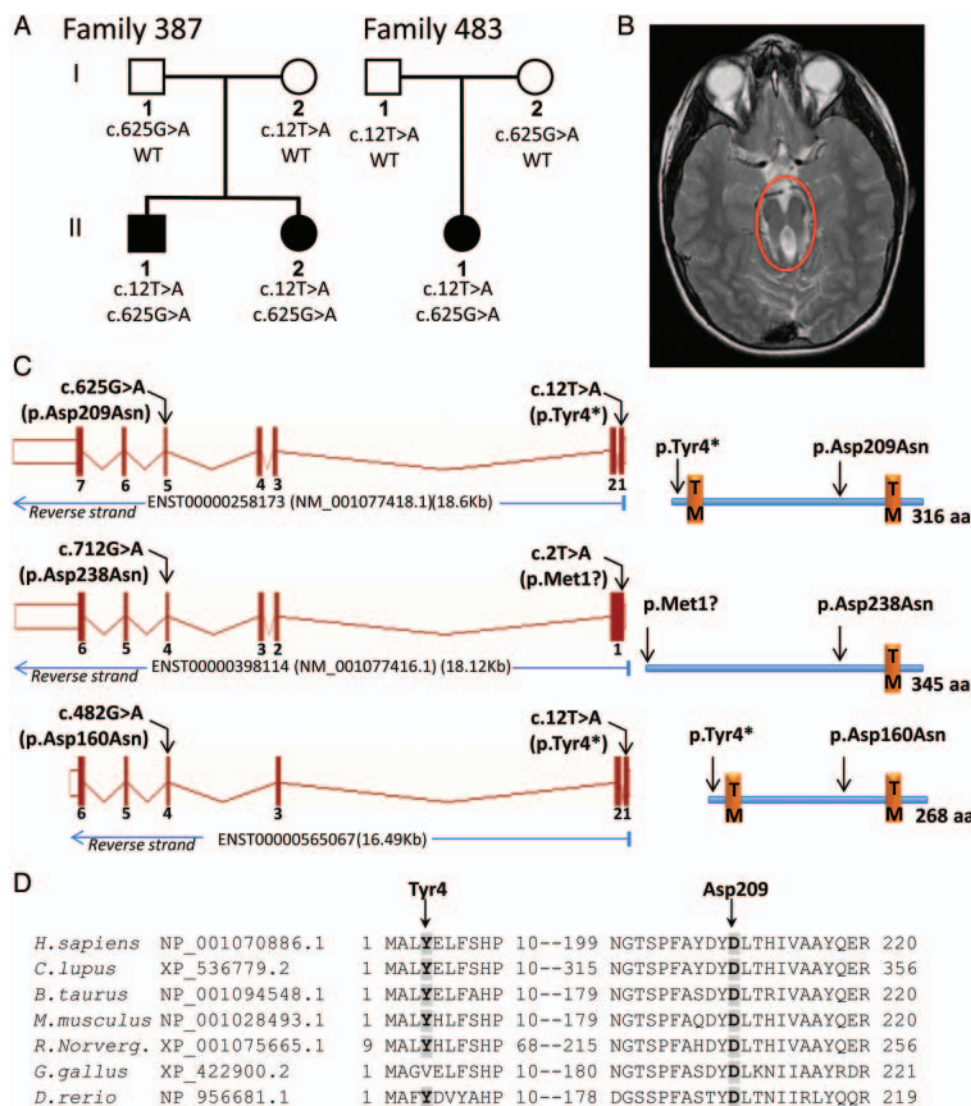


Figure 1 (A) Segregation of mutations in *TMEM231* in JBTS families. (B) Brain MRI from individual II-1 from family 387 showing the 'molar tooth sign'. (C) Left panel, scheme showing the presence of the mutations with respect to the different *TMEM231* Ensemble-annotated transcripts predicted to produce proteins; right panel, the corresponding *TMEM231* proteins are depicted in the right panel. TM, denotes the presence of a transmembrane domain, as predicted by *SMART* algorithm. (D) Amino acid conservation of the residues affected by the p.Tyr4* and p.Asp209Asn mutations in *TMEM231*. Amino acid alignments were generated using homologue (NCBI).

cohort. They are dependent in all activities of daily living, are non-verbal, and can take steps only if assisted with a walker. Both siblings from family 387 had significant aggressive and self-mutilating behaviour consisting of head banging and biting, requiring treatment with antipsychotic agents, mouth guard and protective helmet. Individuals II-2 from family 387 and II-1 from family 483 show extraneural manifestations consisting of retinopathy, bilateral macroscopic renal cysts (but normal renal function), and postaxial polysyndactyly of one foot (table 1 and see online supplementary figure S1, D-F).

The presence of rare and potentially deleterious mutations in *TMEM231*, which segregate with the disease in two unrelated FC families, strongly suggests that disruption of this gene causes JBTS in our subjects. The fact that the three individuals with the mutations in *TMEM231* show a similar form of JBTS also supports the involvement of this gene. Furthermore, several observations indicate that *TMEM231* plays a key role in the cilia, and physically interacts with known JBTS genes. *TMEM231* encodes a transmembrane protein that localises at

the base of the ciliary axoneme at the transition zone.¹⁹ Recently, *TMEM231* was shown to be part of the B9 complex, which is required for a diffusion barrier between the cilia and plasma membranes that maintains the integrity of the cilia as a privileged membrane domain.¹⁹ The B9 complex includes at least 13 proteins (BD91, BD92, TCTN1, TCTN2, TCTN3, CC2D2A, *TMEM216*, *TMEM67*, *TMEM237*, *TMEM231*, *MKS1*, *AHI1*, *TMEM17*), all of which, with two exceptions (*TCTN3*, *TMEM17*), are involved in JBTS and/or Meckel-Gruber syndrome (MKS), a related ciliopathy.^{19 20} Proteomic studies using the MKS proteins BD91 or BD92 as baits have shown that *TMEM231* is in a complex with *TCTN1*, *TCTN2*, *MKS1*, *AHI1* and *CC2D2A*.¹⁹ siRNA knockdown of *TMEM231* disrupts the integrity of the ciliary barrier and the localisation of components of the B9 complex at the transition zone, resulting in a reduction of cilia formation and ciliary localisation of signalling receptors.¹⁹ *TMEM231* knockout mice die at E15.5 with severe vascular defects, and display typical features of ciliopathy, namely microphthalmia, polydactyly and

abnormalities in patterning of ventral spinal cord.¹⁹ Altogether, these observations indicate that autosomal-recessive mutations in *TMEM231* are a cause of JBTS.

JBTS in FCs show both locus and allelic heterogeneity. We identified three JBTS genes in this population with a total of 14 different alleles. Three mutations in *C5ORF42*, two mutations in *CC2D2A* and two mutations in *TMEM231* were found in at least two unrelated affected individuals (table 1). Our analysis indicates that each of these mutations is located within a distinct haplotype in these individuals, suggesting the existence of multiple founder effects.¹² Founder effects are typically associated with an increase in the frequency of a specific autosomal recessive allele, which is often accompanied by other alleles that remain at their usual background frequency. Interestingly, for each of these three JBTS genes, we found at least two founding mutations. It is likely that more of these complex founder effects will be unravelled with the use of genomic sequencing.

In summary, combining this study and our previous one, we were able to explain the underlying genetic cause of JBTS in 21/24 FC individuals using exome sequencing. In the course of this work, we identified *TMEM231* as a novel JBTS gene. This discovery gives further support to the concept that JBTS results from disruption of the barrier at the ciliary transition zone.

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Contributors MS, FFF, JM, JLM: study design, data analysis and interpretation and manuscript writing and revision. JS: data analysis and manuscript writing and revision. GM, EL, LP, SD: laboratory follow-up of candidate variants and segregation studies. MS, JLM, LHO, MIS, VD, DA, EA, GS, BM: patient recruitment, examination and counselling. DL, GAR: contribution of control samples. CM: coordination of samples and patient consents.

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Competing interests None.

Ethics approval Ethics Committee of Sainte Justine Research Center.

Provenance and peer review Not commissioned; externally peer reviewed.

WEB RESOURCES 1000 Genomes Project, <http://browser.1000genomes.org/index.html>
dbSNP, <http://www.ncbi.nlm.nih.gov/projects/SNP/>
Ensemble Genome Browser: <http://www.ensembl.org>
ESP Exome Variant Serve (EVS): <http://evs.gs.washington.edu/EVS/>
Gene Ontology, <http://www.geneontology.org/>
Mutation Taster: <http://www.mutationtaster.org/>
NCBI homologue, <http://www.ncbi.nlm.nih.gov/homologue>
NCBI Nucleotide database, <http://www.ncbi.nlm.nih.gov/nucleotide>
Online Mendelian Inheritance in Man (OMIM), <http://www.omim.org>
Polyphen-2: <http://genetics.bwh.harvard.edu/pph2/>
SIFT: <http://sift.jcvi.org/>
SMART sequence analysis: <http://smart.embl-heidelberg.de/>

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