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Delineation of the Clinical, Molecular and Cellular Aspects of Novel *JAM3* Mutations Underlying the Autosomal Recessive Hemorrhagic Destruction of the Brain, Subependymal Calcification and Congenital Cataracts

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Abstract

We have recently shown that the hemorrhagic destruction of the brain, subependymal calcification and congenital cataracts is caused by biallelic mutations in the gene encoding junctional adhesion molecule 3 (JAM3) protein. Affected members from three new families underwent detailed clinical examination including imaging of the brain. Affected individuals presented with a distinctive phenotype comprising hemorrhagic destruction of the brain, subependymal calcification and congenital cataracts. All patients had a catastrophic clinical course resulting in death in 7 out of 10 affected individuals. Sequencing the coding exons of *JAM3* revealed three novel homozygous mutations: c.2T>G (p.M1R), c.346G>A (p.E116K) and c.656G>A (p.C219Y). The p.M1R mutation affects the start codon and therefore is predicted to impair protein synthesis. Cellular studies showed that the p.C219Y mutation resulted in a significant retention of the mutated protein in the endoplasmic reticulum, suggesting a trafficking defect. The p.E116K mutant traffics normally to the plasma membrane as the wild type and may have lost its function

Conflict of interest

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due to the lack of interaction with an interacting partner. Our data further support the importance of JAM3 in the development and function of the vascular system and the brain.

Keywords

brain; subependymal calcification; congenital cataract; JAM3

Introduction

Junctional adhesion molecule 3 (JAM3; MIM# 613730) is a member of the JAM subfamily of proteins which include F11R (JAM1, JAM-A), JAM2 (JAM-B), JAM3 (JAM-C) and IGSF5 (JAM4) (Weber et al., 2007; Ebnet, 2008). Other related members of this subfamily of proteins include endothelial cell selective adhesion molecule (ESAM) and coxasackie virus and adenovirus receptor (CXADR) (Weber et al., 2007). JAMs and related proteins are localize at intercellular contacts and participate in the assembly and maintenance of tight junctions (TJs) and the control of cellular permeability (Ebnet, 2008). TJs are morphologically distinct subcellular structures that are located closer to the apical side (compared to other junctional complexes) and are highly regulated areas of close contact between the plasma membranes of adjacent epithelial and endothelial cells (Balda and Matter, 2008). TJs are important for the formation of polarized epithelial and endothelial barriers by forming intermembrane diffusion barriers and by controlling diffusion along the paracellular pathway (Cereijido et al., 2008). It has also been found that TJs are important for intracellular signaling (Matter and Balda, 2007; Steed et al., 2009; Terry et al., 2010). JAMs have been found to regulate adhesion between leukocytes and endothelial cells and the paracellular transmigration of leukocytes across the endothelium (Bradfield et al., 2007; Weber et al., 2007). Moreover, JAMs play a role in cell polarisation during spermatogenesis (Gliki et al., 2004).

In addition to JAMs, TJs contain other proteins including claudins, occludins and tricellulins (Ebnet et al., 2004; Ebnet, 2008). JAMs are single-span transmembrane proteins whereas claudins, occludins and tricellulins are tetraspan transmembrane proteins (Balda and Matter, 2008; Ebnet, 2008). The tetraspan proteins form the paracellular permeability barrier and therefore determine the selectivity and extent of paracellular diffusion. JAMs on the other hand are postulated to mediate homotypic cell-cell adhesion (Bazzoni et al., 2000; Weber et al., 2007). Several inherited conditions have been shown to be caused by mutations in genes encoding members of TJs protein complexes (O'Driscoll et al., 2010). Hypomagnesaemia has been shown to be caused by mutations in genes encoding claudins 16 and 19 (Simon et al., 1999; Konrad et al., 2006) and familial hypercholanemia to be caused by mutations in *TJP2* and *BAAT* genes (Carlton et al., 2003). Similarly, mutations in claudin 14 and tricellulin have been found to cause hereditary deafness (Wilcox et al., 2001; Riazuddin et al., 2006) and mutations in *NHS* to cause Nance-Horan syndrome (Burdon et al., 2003). The variability in the clinical presentations reflects the variable functions of TJs in various tissues and organs.

We have recently identified a homozygous mutation in *JAM3* in a consanguineous family from the United Arab Emirates (UAE) with a rare recessive disorder that is characterised by hemorrhagic destruction of the brain, subependymal calcification, and congenital cataracts (MIM# 613730) (Mochida et al., 2010). In this manuscript, we describe three additional novel mutations in *JAM3* in patients from different ethnic backgrounds with similar phenotypes and elucidate the cellular mechanisms underlying some of those mutations.

Materials and Methods

Clinical evaluations

All individuals participating in this study or their guardians provided written informed consents. Clinical assessments were performed by experienced clinical geneticists in Turkey, Spain, Australia and the UAE. Full medical history was obtained. Participating members underwent careful clinical examination including imaging of the brain by magnetic resonance imaging (MRI).

Mutation screening

Genomic DNA was extracted from peripheral leukocytes using the Flexigene DNA kit (Qiagen, Santa Clara, CA). Intronic primer pairs were designed to amplify individual *JAM3* exons and its exon/intron junctions using Primer3 (http://frodo.wi.mit.edu/). Following PCR amplification, the PCR products were cleaned using the ExoSAP-IT reagent (USB Co., Cleveland, Ohio, USA) followed by DNA sequencing using the BigDye Terminator v3.1 Cycle Sequencing kit and run on an ABI PRISM 3130x/ DNA Analyzer (Applied Biosystems, Foster City, CA). Sequencing data were analyzed using clustalw2 referencing the NCBI GeneBank NM_032801.4 for *JAM3* gene.

Site-directed mutagenesis

A transfection-ready *JAM3*-pCMV6 plasmid tagged C-terminally with Myc and Flag was purchased from OriGene Inc. (OriGene, Rockville, USA). The identified missense mutations (p.E116K and p.C219Y) were introduced by site-directed mutagenesis using *Pfu* Ultra high-fidelity DNA polymerase (Stratagene, La Jolla, CA) following the manufacturer's protocol. The primers used for site directed mutagenesis are the following with the mutagenic nucleotide in bold:

1. For generating c.346G>A (p.E116K) we used

Forward:5' CCCTTTATCGCTGTAAGGTCGTTGCTCG 3'

Reverse: 5' CGAGCAACGACCTTACAGCGATAAAGGG 3'

2. For generating c.656G>A (p.C219Y) we used

Forward: 5' CTGGGCAGTACTACTACATTGCTTCCAATG 3'

Reverse: 5' CATTGGAAGCAATGTAGTAGTACTGCCCAG 3'

All constructs were confirmed by DNA sequencing.

Cell culture

Human cervical cancer (HeLa) cell lines were cultured in DMEM (Invitrogen, Carlsbad, CA, USA) supplemented with 10% heat-inactivated fetal bovine serum at 37°C with 5% CO2. Transient transfection of cells was achieved with FuGENE HD transfection reagent (Promega, Madison, Wisconsin, USA) using 0.5 μ g plasmid cDNA for confocal microscopy experiments. Co-transfection with hRas-GFP plasmid (0.2 μ g cDNA) served as a transfection indicator and a plasma-membrane marker. All experiments were performed 24–48 h following transfection.

Confocal fluorescence microscopy

Transfected cells were grown on coverslips (Ibidi, Martinsried, Germany) and fixed (5 min) with methanol (Sigma-Aldrich, St. Louis, MO, USA) at -20 °C. After 3 washes (5 min each) with PBS cells were blocked with 1% bovine serum albumin (BSA, Sigma-Aldrich,

St. Louis, MO, USA) for 30 minutes. Membranes were incubated with primary antibodies at room temperature for 1 hr. Mouse anti-flag (Sigma Aldrich, St. Louis, MO, USA) and Rabbit anti-calnexin (Santa Cruz Biotechnology, Santa Cruz, CA, USA) were used at 1:1000 and 1:100 dilutions, respectively. After the primary antibodies incubations, the slides were washed 3 times (5 min each) then incubated for 1 hour at room temperature with secondary antibodies (fluorescence-labelled 'Alexa-Fluor-568' [anti-mouse] and 'Alexa-Fluor-488' [anti-rabbit] (Santa Cruz Biotechnology, Santa Cruz, CA, USA). Confocal microscopy was performed with a Nikon Eclipse system (Nikon Instruments Inc., Japan). Control experiments without the primary antibodies or with non-transfected cells were carried out and revealed very low-level background staining.

Results

Clinical Features

Three families from different ethnic backgrounds (one Turkish, one Afghani and one Moroccan) have been recruited to participate in this study (Figure 1). Consanguinity is evident in two of the families and the clinical features of all the affected children are listed in Supp. Table S1.

Family 1—The parents of the affected child in this family are second cousins of Turkish origin. The parents are normal. Their first child (II1) was a preterm baby girl who had intracranial bleeding and died in the neonatal period. The second child (II2) of the couple was a female who was the product of normal pregnancy and term delivery. Her birth weight was 2480 gm ($<10^{th}$ centile). She was admitted to the neonatal intensive care unit on day 5 due to tonic seizures which were resistant to treatment. Physical examination revealed a head circumference of 30 cm, slopping forehead, prominent, bulging anterior fontanelle, bitemporal grooving and bilateral nuclear cataracts (Figure 2A-1). Cranial ultrasound showed a prominent expansive bleeding. Brain computerized tomography showed calcification of the subependymal region and multifocal intraparenchymal hemorrhage (Figure 2A-2). Serological tests were negative for toxoplasmosis, rubella, and cytomegalovirus. Cerebrospinal fluid PCR for cytomegalovirus and toxoplasma were also negative. Platelets levels were normal. The infant developed post hemorrhagic hydrocephalus and died at 2 months of age.

Family 2—The parents are non-consanguineous and healthy of Afghani origin. Patient 1 (II1), was the first child of the family born at term by emergency caesarean section. His birth weight was 2480 gm (<10th centile). He was noted to have bilateral cataracts and had feeding problems from birth. He was discharged home on day 4. At 2 weeks of age he presented with vomiting, irritability and a high-pitched cry. Brain MRI at the age of 15 days showed Grade IV intraventricular haemorrhage, severe ventricular dilatation and white matter abnormalities in both cerebral hemispheres and multiple subependymal cysts (Figure 2B). Renal ultrasound showed horseshoe kidneys with echogenic cortex and altered corticomedullary differentiation. He had progressive neurological deterioration and died at two weeks of age. Investigations were all normal and included respiratory chain enzymes on muscle and liver, 7 DHC, very long chain fatty acids, transferrin isoforms, blood karyotype, Affy 250K array, and coagulation study including platelets. Mitochondrial point mutations (MELAS, NARP, Leigh syndrome) were all negative.

Patient 2 (II3), was the third child of the family, with the family's second child being a healthy male. Patient 2's pregnancy was normal and delivery was by elective caesarean section at term. His birth weight was 3315 gm (25^{th} centile), length was 52 cm ($50-90^{th}$ centile), and head circumference was 35 cm (50^{th} centile). At birth he was noted to have

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bilateral cataracts. His tone and reflexes were normal and there was no clinical evidence of seizures. Cranial ultrasound showed periventricular echogenicity but there was no bleed. However, he deteriorated on day 5 and his haemoglobin dropped from 180g/L to 96d/L. He required ventilation. EEG on the third day of life showed bilateral epileptiform activity. Cranial ultrasound revealed extensive bilateral intracerebral haemorrhage. He died on the fifth day of life. Abdominal ultrasound showed pelvicalyceal dilatation of the right kidney. Full metabolic work up was normal.

Family 3—The parents are first cousins of Moroccan origin, who had a total of five pregnancies. The family history is significant in that the mother's brother (I3) died on the seventh day of life without any known diagnosis. This couple's first child was a male born at 36 weeks gestation who had cerebral haemorrhage, hydrocephalus and died at 10 days of age. They then had two healthy children. The fourth pregnancy ended with a stillborn child who had hydrocephalus and the cerebral tissues showed necrotic degeneration. The index case (II5) is 28 days old female, the product of normal pregnancy. Fetal tachycardia was detected at 37 weeks gestation and therefore the baby was delivered by caesarean section. Her birth weight was 2520 gm (10-25th centile), length 46.5 cm (25th centile), and head circumference 34.5 cm (50th centile). Bilateral cataracts were detected at birth both central and nuclear (Figure 2C-1). On the second day of life she presented with irritability, crying and a bulging anterior fontanelle. Brain ultrasound showed dilated lateral ventricles, intraventricular haemorrhage in the right ventricle and frontal periventricular cysts. On the 8th day of life she was referred to another hospital for neurosurgical evaluation. Head CT revealed multifocal intraparenchymal hemorrhages, subependymal calcification and diffusely hypodense brain parenchyma (Figure 2C-2). MRI of the brain revealed intraventricular and intraparenchymal hemorrhage, edema and cystic changes of the brain parenchyma, dilated ventricles and thin corpus callosum with haemohrragic damage (Figure 2C-3). There was normal circulation in the circle of Willis arteries. The EEG showed electrical status epilepticus, however there were no seizures clinically. Metabolic screening was normal. Platelets levels were normal. It was decided that the child was not a candidate for ventricular shunt. She had severe anemia. Brain CT scan demonstrated bilateral subependymal calcification in the lateral ventricles. On the 18th day of life her head circumference was 41 cm (+4 SD) and she was discharged for palliative care at home. She died at home the age of 39 days.

Mutation analysis

DNA sequencing revealed three different homozygous missense mutations in *JAM3* gene in the three studied families (Table 1; Supp. Figure S1). In family 1 we detected a non-synonomous mutation (c.656G>A) in exon 6 of *JAM3* gene, which led to an amino acid substitution at position 219 (p.C219Y). In family 2 we identified a single nucleotide substituation (c.346G>A) in exon 4 of *JAM3* leading to a missense mutation at position 116 (p.E116K). The third mutation (c.2T>G) changed the initiator codon (methionine) of JAM3 leading to amino acid substitution to arginine (p.M1R). The three mutations were not reported in dbSNP or the 1000 genome project. All the amino acids affected by these mutations were highly conserved across species (Table 1) and were predicted to be disease-causing by Polyphen2 (http://genetics.bwh.harvard.edu/pph2/), SIFT (http://sift.jcvi.org/) and Mutation taster (http://www.mutationtaster.org/). Furthermore, the detected mutations segregated with disease status in the three families; the parents were heterozygous, consistent with carrier status and the available unaffected siblings were homozygous for the wild type or heterozygous for the mutation. All the detected mutations were submitted to the LOVD database (www.lovd.nl/JAM3).

C219Y mutation results in JAM3 trafficking defect

In order to further understand the pathophysiological significance of the two detected missense mutations (p.E116K and p.C219Y), we employed confocal fluorescence microscopy to establish their subcellular localization relative to the wild type (WT) JAM3 protein. HeLa cells, known to lack endogenous expression of JAM3 (Betanzos et al., 2009), were either transfected with individual *JAM3* constructs (WT, p.E116K or p.C219Y) or co-transfected with GFP-hRas. The cells were then stained with anti-flag antibodies to visualize *JAM3* constructs. The cells transfected with *JAM3* only constructs were co-stained with anticalnexin antibodies to visualize the endoplasmic reticulum (ER) network. As shown in panels A–F of Figure 3, the wild type JAM3 is predominantly localized to the plasma membrane and co-localised largely with the GFP-hRas. Similarly, the p.E116K is predominantly localized to the plasma membrane as shown in panels G–L of Figure 3. On the other hand, the p.C219Y mutant co-localized predominantly with the ER marker (calnexin) and not with GFP-hRas indicating its retention in the ER (Figure 3 panels M–R, Supp. Figure S2).

Discussion

The impact of JAM3 Mutations on the protein product

Arrate et al. (2001) and Santoso et al. (2002) cloned and characterized JAM3 as an approximately 43KDa transmembrane glycoprotein. This type I integral membrane glycoprotein consists of two immunoglobulin (Ig)-like domains with intrachain disulfide bonds at the extracellular region. By sequence analysis Arrate et al. (2001) concluded that the N-terminal domain is a V-type while the membrane proximal domain is a C2-type. Also they predicted that the intrachain disulfide bonds that stabilize each Ig domain are between C53–C115 and C160–C219. Ig-like domains are implicated in a range of vital functions, including cell-cell recognition, cell-surface receptors, muscle structure and the immune system (Teichmann and Chothia, 2000). The C-terminal region of the protein is composed of a short transmembrane domain followed by an intracellular domain containing a PDZ binding motif (Supp. Figure S1A).

The p.C219Y mutation detected in family 1 alters one of the four highly conserved cysteine residues of JAM3 disrupting one out of two disulfide bonds known to stabilize the Ig-folds of the protein (Arrate et al., 2001; Santoso et al., 2002). Several functional studies demonstrated that cysteine-substitutions disrupt the structure of cell surface proteins causing their misfolding, instability and ER retention (Rudarakanchana et al., 2002; Fukuda et al., 2011). In consistent, comparison of the subcellular localization of flag-tagged C219Y mutants transiently expressed in HeLa (Figure 3; Supp. Figure S2A) and COS7 (Supp. Figure S2C) clearly showed that most of the mutated protein failed to reach the cell membrane. ER retention is a well established disease causing mechanism in many human syndromes caused by loss-of-function mutations (Chen et al., 2005; Aridor, 2007; Ali et al., 2011).

The p.E116K mutation found in family 2 did not seem to affect the subcellular localization or trafficking of the protein (Figure 3). However, its position in the Ig-like V-type domain suggests an interfering role of this mutation with the dimerization or with the interaction(s) of this protein with one or more of its interacting partners. JAM3 controls tight junction maintenance by engaging in homophilic and heterophilic interactions with neighboring JAM molecules (Ebnet et al., 2008).

Arrate et al. (2001) and Santoso et al. (2002) predicted the first ATG by homology alignments between *JAM3*, *JAM1*, *JAM2* and mouse *Jam3*. Arrate et al. (2001) found using

SignalP V1.1 the signal sequence of JAM3 which should be cleaved after glycine 30 from the putative ATG. The mutated methionine in family 3 is predicted to be the first methionine of JAM3 protein and such mutations almost invariably interfere with the translation of the protein. However, there are two in-frame ATG's upstream of this start codon and the improbable use of one of them or for the next in-frame ATG would encode a non-functional or unstable protein.

Clinical consequences of JAM3 deficiency

Mouse Jam3 is expressed at the apical junctions of endothelial cells, smooth muscle cells, fibroblasts, Schwann cells, spermatids and hematopoietic stem cells. More recently, Jam3 expression was detected in the tight junctions of neural stem cells in the embryonic ventricular zone and the adult ependymal cell layer of mouse brain (Stelzer et al., 2012). More than half of mice with homozygous mutation in *Jam3* exhibit postnatal lethality. *Jam3* deficient mice exhibit growth retardation, muscle weakness, abnormal spermatogenesis, granulocytes homeostasis defects, electrophysiological defects, and hypersensitivity to mechanical stimuli (Scheiermann et al., 2007; Colom et al., 2012). Furthermore, mouse Jam3 apical expression was found on the embryonic retinal neuroepithelia, and its deficiency caused nuclear congenital cataracts (Daniele et al., 2007). However, the brains of mice lacking Jam3 are relatively unaffected (Stelzer et al., 2012).

In human, Mochida et al. (2010) demonstrated that a homozygous mutation in JAM3 is the cause of a severe autosomal recessive syndrome segregating in a consanguineous family. The affected members in this family were born with congenital cataracts and developed severe brain abnormalities from progressive hemorrhagic destruction of the brain tissue, including the cerebral white matter and basal ganglia. Three out of eight affected individuals died early in infancy. Brain imaging in all the affected showed calcification in the subependymal region of the brain, multifocal intraparenchymal hemorrhage with associated liquefaction and massive cystic degeneration resulting in large ventricles (Al-Gazali et al., 1999; Mochida et al., 2010). The phenotype of the children in this report is very similar to the phenotype of the affected children in this previously reported family. All the affected children in this report had bilateral cataracts and develop progressive neurological impairment, associated with seizures in some individuals. In addition neuro-imaging in all patients showed intraparanchymal hemorrhage with evidence of brain destruction and subependymal, basal ganglia and white matter calcification. These changes are very similar to those found in the original family. Some of the other features described in the original family, such as renal anomalies and liver enlargement, were more variable in this patient group. The disorder was lethal in 7 out of the 10 cases. Death occurred in the first few weeks of life. Those who survived (3 in total) are severely retarded with severe spasticity and seizures. Other conditions which could be considered in the differential diagnosis include the syndrome of band-like intracranial calcification, simplified cerebral gyrai and polymicrogyria (Abdel-Salam et al., 2008; O'Driscoll et al., 2010). This syndrome is caused by mutations in another TJ gene, occludin (OCLN; MIM# 602876). However, patients with this syndrome do not have intracranial hemorrhage or congenital cataract (Mochida et al., 2010). Furthermore, our patients did not show any evidence of developmental malformation of the cerebral cortex and polymicrogyria. Similarly mutations in collagen type IV alpha (COL4A1; MIM# 120130), can cause overlapping clinical presentations which include antenatal intracranial hemorrhage (de Vries et al., 2009), brain small vessel disease with hemorrhage or vascular leukoencephalopathy, porencephaly (Gould et al., 2006; Sibon et al., 2007), and hereditary angiopathy with nephropathy, aneurysms and muscle cramps (Plaisier et al., 2007). All these disorders are inherited as autosomal dominant and the clinical presentations is different from the cases in this report. Proliferative vasculopathy and hydranencephaly-hydrocephaly syndrome (Fowler syndrome) is an autosomal recessive

perinatally lethal disorder characterized by hydrocephalus associated with progressive destruction of the central nervous system (CNS) tissue as a result of an unusual and characteristic proliferative vasculopathy (Fowler et al., 1973). The hallmark of the syndrome is the microvascular proliferation which is associated with extensive necrosis and calcification of the CNS tissue. There are no visceral malformations (Lalonde et al., 2010). It is caused by mutations in the *FLVCR2* gene (MIM# 610865). Clinically affected fetuses present with fetal akinesia deformation sequence with muscular neurogenic atrophy (Fowler et al., 1973; Meyer et al., 2010) which is different from the clinical presentation of the cases in this report.

Our study revealed several missense mutations scattered throughout the coding sequence of *JAM3* affecting different domains yet the affected children shared similar phenotype indicating homogeneity of this disorder at the clinical and molecular level. This is likely due to total loss of the protein function as a result of those mutations.

Conclusion

Our study confirms the importance of JAM3 as a component of the junctional complexes and its deficiency leading to a distinctive and catastrophic neonatal presentation of cataracts and hemorrhagic destruction of the brain.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1. Pedigree Drawings of the studied Families

Three families with different ethnic origin were recruited in this study. Family 1 is Turkish, family 2 is Afghani, while family 3 is Moroccan. Filled symbols denote affected individuals with the syndrome of hemorrhagic destruction of the brain, subependymal calcification, and congenital cataracts. Sequenced individuals are indicated with an asterisk.



Figure 2. Clinical data

A-1) Facial picture of the affected individual in Family 1 shows bilateral nuclear cataracts.
A-2) Head CT image of the patient in Family 1 reveals multifocal intraparenchymal hemorrhages (arrows) and subependymal calcification (arrowheads). B) T2-weighted axial brain MRI of Patient 1 in Family 2 reveals multifocal intraparenchymal hemorrhages of varying ages (arrows) as well as intraventricular hemorrhage (asterisks).

C-1) Facial picture of the affected individual in family 3 shows bilateral cataracts, enlarged frontal part of the cranium and prominent scalp veins. **C-2)** Head CT of the patient in family 3 shows multifocal intraparenchymal hemorrhages (arrows) and subependymal calcification (arrowheads). Diffusely hypodensity of brain parenchyma is also noted, suggesting severe edema. **C-3)** T2-weighted coronal MRI image of the same patient shows intraventricular hemorrhage with associated enlargement of the ventricles.



Figure 3. p.C219Y mutation disrupts normal intracellular JAM3 protein localization HeLa Cells transiently transfected with C-terminal Flag-tagged *JAM3*-pCMV5 vector constructs wild type (WT), **A**–**F**; p.E116K, **G**–**L**; and p.C219Y, **M**–**R**) and stained with anti-Flag Ab (red). WT vector localizes to the plasma membrane (PM) with the hRas-GFPtagged marker (**A**–**C**) and not with the endoplasmic reticulum (ER) marker calnexin (green; **D**–**F**). JAM3–p.E116K–Flag construct localization was similar to the WT with hRas-GFPtagged marker (**G**–**I**) and the calnexin (**J**–**L**). P.C219Y construct do not localize at the PM like hRas-GFP-tagged marker (**M**–**O**) and appears to be accumulated or clustered within the ER With calnexin (**P**–**R**).

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Table 1

Mutations detected in JAM3

ICTC: ICTC:	CAGGCACTTTGCTAAGATCTC CAGGCACTTTGTTAAGATCTC CAGGCACTTTGCTAAGATCTC CAGGCACTCTGCTAAGATCTC GGCACTCTGCTAAGATCTC GGCACTCTGCTAAGATCTC CAGGCACTCTGCTAAGA	Normal CAGGCACTTTGCTAGGATCTC Mutated CAGGCACTTTGCTAAGATCTC Human CAGGCACTTTGCTAAGATCTC Chimp CAGGCACTCTGGTAAGATCTC Cat CAGGCACTCTGGTAAGATCTC Cat CAGGCACTCTGGTAAGATCTC Mouse CAGGCACTCTGGTAAGA Normal FTAVHKDDSGQYYCIASNDAGS Mutated FHAVHKDDSGQYYCIASNDAGS M.musculus FNAVHKDDSGQYYCIASNDAGF	Communication Construct Mutated CAGGCACTTTGGTAAGATCTC Human CAGGCACTTTGGTAAGATCTC Human CAGGCACTTTGGTAAGATCTC Cat CAGGCACTTTGGTAAGATCTC Cat CAGGCACTTTGGTAAGATCTC Cat CAGGCACTTTGGTAAGATCTC Cat CAGGCACTTTGGTAAGATCTC Cat CAGGCACTTTGGTAAGATCTC Mouse CAGGCACTCTGGTAAGATCTC Mouse CAGGCACTCTGGTAAGATCTC Mouse CAGGCACTCTGGTAAGATCTC Mutated FTAVHKDDSGQYYCIASNDAGS Mutated FHAVHKDDSGQYYCIASNDAGS M.musculus FNAVHKDDSGQYYCIASNDAGS R.norvegicus FNAVHKDDSGQYYCIASNDAGS B.taurus FNAVHKEDSGQYYCIASNDAGS B.taurus FSAVHKEDSGQYYCIASNDAGS	Splite All Common Normal CAGGCACTTTGGTAAGATCTC Mutated CAGGCACTTTGGTAAGATCTC Human CAGGCACTTTGGTAAGATCTC Chimp CAGGCACTTTGGTAAGATCTC Cat CAGGCACTTGGTAAGATCTC Anore CAGGCACTTTGGTAAGATCTC Anore CAGGCACTTTGGTAAGATCTC Anore CAGGCACTTGGTAAGATCTC Anore FAAVHKDDSGQYYCIASNDAGA Anore FAAVHKDDSGQYYCIASNDAGA Anore
	CAGGCACTTTGGTAAGATC CAGGCACTCTGGTAAGATC GGCACTCTGGTAAGATC CAGGCACTCTGGTAAGA	HumanCAGGGCACTTTGGTAAGATCChimpCAGGCACTCTGGTAAGATCCatGGGCACTCTGGTAAGATCGGCACTCTGGTAAGATCGGGCACTCTGGTAAGAMouseCAGGCACTCTGGTAAGANormalFTAVHKDDSGQYYCIASNDMutatedFTAVHKDDSGQYYCIASNDG.gallusFHAVHKGDTGRYSCIATNDM.musculusFNAVHKDDSGOYYCIASND	Human CAGGCACTTTGGTAAGATC Chimp CAGGCACTCTGGTAAGATC Cat CAGGCACTCTGGTAAGATC Cat CAGGCACTCTGGTAAGATC Mouse CAGGCACTCTGGTAAGATC Mutated FTAVHKDDSGQYYCIASNDF M.musculus FNAVHKDDSGQYYCIASNDF R.norvegicus FSAVHKEDSGQYYCIASNDF B.taurus FSAVHKEDSGQYYCIASNDF	Human CAGGCACTTTGGTAAGATC Chimp CAGGCACTTTGGTAAGATC Cat CAGGCACTCTGGTAAGATC Cat CAGGCACTCTGGTAAGATC Cat CAGGCACTCTGGTAAGATC Cat CAGGCACTCTGGTAAGATC Mouse CAGGCACTCTGGTAAGATC Normal FTAVHKDDSGQYYCIASNDF Mutated FHAVHKDDSGQYYCIASNDF M.musculus FHAVHKDDSGQYYCIASNDF M.musculus FNAVHKDDSGQYYCIASNDF R.norvegicus FSAVHKEDSGQYYCIASNDF B.taurus FSAVHKEDSGQYYCIASNDF
7777		Normal FTAVHKDDSGQYYCIASN Mutated FTAVHKDDSGQYYZIASN G.gallus FHAVHKGDTGRYSCIATN M.musculus FNAVHKDDSGOYYCIASN	<pre> Isilia C1-1979 Normal FTAVHKDDSGQYYCIASN Mutated FTAVHKDDSGQYYCIASN Mutated FTAVHKDDSGQYYCIASN G.gallus FHAVHKDDSGQYYCIASN M.musculus FSAVHKEDSGQYYCIASN R.norvegicus FSAVHKEDSGQYYCIASN B.taurus FSAVHKEDSGQYYCIASN b.taurus for the fo</pre>	Non-symmetries IgeNetC1-type Normmal FTAVHKDDSGQYYCIASN Mutated FTAVHKDDSGQYYCIASN Mutated FHAVHKDDSGQYYCIASN G.gallus FHAVHKGDTGRYSCIATN M.musculus FNAVHKDDSGQYYCIASN R.norvegicus FSAVHKEDSGQYYCIASN C.lupus FSAVHKEDSGQYYCIASN B.taurus FSAVHKEDSGQYYCIASN
ΪX	FTAVHKDDSGQYY C IAS FTAVHKDDSGQYY Y IAS	G.gallus FHAVHKGDTGRYSCIAT M.musculus FNAVHKDDSGOYYCIAS	G.gallus FHAVHKGDTGRYSCIAT M.musculus FNAVHKDDSGQYYCIAS R.norvegicus FSAVHKEDSGQYYCIAS C.lupus FSAVHKEDSGQYYCIAS B.taurus FSAVHKEDSGQYYCIAS	G.gallus FHAVHKGDTGRYSCIAT M.musculus FNAVHKDDSGQYYCIAS R.norvegicus FSAVHKEDSGQYYCIAS C.lupus FSAVHKEDSGQYYCIAS B.taurus FSAVHKEDSGQYYCIAS
INS	FHAVHKGDTGRYSCIA FNAVHKDDSGQYYCIA FSAVHKEDSGQYYCIA FSAVHKEDSGQYYCIA FSAVHKEDSGQYYCIA FTAVHKDDSGQYYCIA FTAVHKDDSGQYYCIA FTAVHKDDSGQYYCIA FTAVHKDDSGQYYCIA	R. norvegicus FSAVHKEDSGQYYCIA C. lupus FSAVHKEDSGQYYCIA B. taurus FSAVHKEDSGQYYCIA H. sapiens FTAVHKDDSGQYYCIA P. troglodytes FTAVHKDDSGQYYCIA D. rerio FSAVKEDAGEYYCRA	P. troglodytes FTAVHKDDSGQIYCIA D. rerio FSAVRKEDAGEYYCRA D. rerio FSAVRKEDAGEYYCRA	H.sapiens FTAVHKDDSGQYYCIA P.troglodytes FTAVHKDDSGQYYCIA D.rerio FSAVRKEDAGEYYCRA
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ARNDRI ARNDRI	TRRDSALYRC E VV. TRRDSALYRC K VV.	Normal TRRDSALYRCEVV Mutated TRRDSALYRCEVV	Le ^{lkeV-type} Normal TRRDSALYRCEVV Mutated TRRDSALYRCEVV	Nersymmetries lystev-type Normal TRRDSALYRCEVV Mutated TRRDSALYRCEVV
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Ш	$\Psi_{ m cDNA}$ change	$\Psi_{ m Protein}$ change	Geno type	Mutation type	Domain		© Conservation	Likelihood of Pathogenicity	Ref.
-	e.215.G	p.MI.R.	Homorygous	Non-synanous	N-terminus	Normal Mutated G.gallus M.musculus R.norvegicus C.lupus B.taurus H.sapiens P.troglodytes D.rerio D.rerio	MALRRPPRLRLCARLPDFFLLLLLFRGC RALRRPPRLRLCARLPDFFLLLLFRGC MSCAISYSLLFVFPGC MALSRRLRLYARLPDFFLLLLFRGC MALSRLRLRLYARLPDFFLLLLFRGC MALSRLRLLCARLPDFFLLLLFRGC MALRRPSLVLMAVLQGC MALRRPFLLCARLPDFFLLLLFRGC MALRPPRLRPRLRCARLPDFFLLLLFRGC MALRPPRLRLCARLPDFFLLLLFRGC MALRPPRLRLCARLPDFFLLLLFRGC	Translation aberrations leading to loss or abrormal expression	This paper
[₩] Mu	ation nomenclature 8	are based on NM_03	2801.4 and	NP_116190.3	starting froi	m A of the ATG as 1			

The nomenclature used by Mochida et al (2010) based on $NM_032801.3$

© Conservation alignments for all the missense mutations are based on homoloGene results for JAM3 (ID: 83700), but the splice site mutation alignments were adopted from Mutationtaster (http://www.mutationtaster.org/) results.

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