A silent mutation, C924T (G308G), in the L1CAM gene results in X linked hydrocephalus (HSAS)

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Abstract
The L1 cell adhesion molecule (L1CAM) is a neuronal gene involved in the development of the nervous system. Mutations in L1CAM are known to cause several clinically overlapping X linked mental retardation conditions: X linked hydrocephalus (HSAS), MASA syndrome (mental retardation, aphasia, shuffling gait, adducted thumbs), spastic paraplegia type I (SPG1), and X linked agenesis of the corpus callosum (ACC). In an analysis of a family with HSAS, we identified a C→T transition (C924T) in exon 8 that was initially thought to have no effect on the protein sequence as the alteration affected the third base of a codon (G308G). Extensive analysis of the other 27 exons showed no other alteration. A review of the sequence surrounding position 924 indicated that the C→T transition created a potential 5′ splice site consensus sequence, which would result in an in frame deletion of 69 bp from exon 8 and 23 amino acids of the L1CAM protein. RT-PCR of the RNA from an affected male fetus and subsequent sequence analysis confirmed the use of the new splice site. This is the first report of a silent nucleotide substitution in L1CAM giving rise to an alteration at the protein level. Furthermore, it shows that as mutation analysis plays an ever more important role in human genetics, the identification of a synonymous base change should not be routinely discounted as a neutral polymorphism.

Keywords: silent mutation; L1CAM; restriction endonuclease fingerprinting (REF); cryptic splice site

Rosenthal et al. were the first to present evidence for the involvement of a mutation of the L1 cell adhesion molecule (L1CAM) in X linked hydrocephalus. The finding was consistent with linkage analyses which placed both L1CAM and X linked hydrocephalus resulting from stenosis of the aqueduct of Sylvius (HSAS) in Xq28, and studies that had implicated L1CAM in neurite outgrowth. Subsequently, mutations in this neuronal glycoprotein were found to cause other disorders clinically related to X linked hydrocephalus, such as MASA syndrome (mental retardation, aphasia, shuffling gait, adducted thumbs), X linked spastic paraplegia type I (SPG1), and X linked agenesis of the corpus callosum (ACC). At present, there are over 50 L1CAM mutations listed on the L1 Mutation Web Page. Approximately 75% of the mutations are contained within 19 exons that represent 65% of the coding sequence. These mutations include frameshifts, missense mutations, stop codons, and alterations in splice site junctions. In almost every case, a mutation appears limited to a single family. In a few instances, the same mutation will give rise to more than one of the above clinically related genetic disorders, so that one can observe both inter- and intrafamilial variability in families with L1CAM mutations.

As part of our interest in X linked mental retardation, we have been investigating families with HSAS and MASA. As expected, we have observed the range of mutations in L1CAM mentioned above. However, in one family with a history of HSAS over four generations, we were unable to detect any mutation other than a presumed silent alteration at position 924 in which a C→T transition affected the third position in a glycine codon (G308G). We present evidence that this neutral mutational event is, in fact, the cause of HSAS as a result of the creation of an alternate 5′ splice site.

Materials and methods

CLINICAL FINDINGS
Case report
The proband (IV.1) had severe congenital hydrocephalus associated with apparent aqueductal stenosis; he received a shunt at 4 days of age. The remaining cortex was very thin. His head circumference was within the normal range at 3 months and was almost 1 SD below the mean at 4 years 8 months. At the age of 7 years, a head MRI showed agenesia of the corpus callosum and hydrocephalus. His hands were held tightly clenched with the thumbs and the third finger overlapping the second finger. All fingers could be straightened passively except the index fingers which were held tightly contracted in the palms. The thumbs could be extended more easily than the index fingers. Therefore, he appeared to have adducted/contracted index fingers rather than the adducted thumb position that is sometimes associated with HSAS.

FAMILY HISTORY
Four maternal great uncles died at birth or in early infancy with congenital hydrocephalus; they were brothers of the maternal grandmother (fig 1A). Clinical information is not available, but they were described by the family
Silent mutation in LICAM and hydrocephalus

as looking like the proband. This family is family 7 in the report by Willems et al.2

GENOMIC DNA
Blood or amniocytes were obtained with informed consent. Genomic DNA was isolated from peripheral blood using a high salt method.11 DNA from amniocytes was prepared using routine procedures.12

RESTRICTION ENDONUCLEASE FINGERPRINTING (REF)
Exons 7-10 were amplified as a 1088 bp fragment using the following primers: ex7-10f: 5’CTGCGGTTGAGGAAGAGGTG3’, ex7-10r: 5’CCAGTTGGTGACGGAGCAGA3’. The PCR reaction mixture contained 10 mmol/l Tris-HCl, pH 8.3, 50 mmol/l KC1, 1.5 mmol/l MgCl2, 50 μmol/l of each dATP, dCTP, dGTP, dTTP, 1 μmol/l of each primer, 2.5 U of Taq DNA polymerase (Boehringer-Mannheim), 0.55 μg TaqI polymerase start antibody (Clontech), and 125 ng genomic DNA in a 50 μl total volume. The initial denaturing temperature was 94°C for five minutes. This was followed by 35 cycles of denaturation at 94°C for 30 seconds, annealing at 70°C for one minute, and extension at 72°C for 1.5 minutes, followed by a final extension at 72°C for 10 minutes. The PCR products were checked by 1.5% agarose gel electrophoresis before doing REF.

A total of 100 ng of the exons 7-10 PCR product were digested separately by four different restriction endonucleases in a total volume of 10 μl. The four digestion reactions contained 4 U HhaI in NEB buffer 4, or 4 U DdeI in NEB buffer 3, or 4 U HinfI, or 4 U MspI in NEB buffer 2, and 0.4 U CIAP at 37°C for 12 hours.
After digestion, all the endonucleases were inactivated at 96°C for 10 minutes. The several reactions were mixed together and reheated at 96°C for five minutes. Digested fragments (10 ng) were incubated with 6 μCi γ-32P ATP and 1 U T4 kinase in a buffer of 70 mmol/l Tris-HCl, pH 7.6, 10 mmol/l MgCl2, 5 mmol/l DTT at 37°C for three hours. After the labelling reaction, 10 μl stop buffer (95% formamide, 10 mmol/l NaOH, 0.25% bromophenol blue, 0.25% xylene cyanol) were added to each tube and 4 μl were loaded onto either a 6% acrylamide gel or a MDE (FMC) gel separately. The 6% acrylamide gel contained 10% glycerol and 0.6 × TBE and was run at 8 W, 4°C for 24 hours. The MDE gel was prepared according to the FMC instruction manual and run at 8 W at room temperature for 12 hours. The gels were dried and exposed to Biomax film (Kodak, Rochester, NY, USA) at ~80°C for 1.5 hours.

SEQUENCING OF PCR PRODUCTS
PCR products (50 μl) were run on 1% agarose gels containing 1 × TBE. The DNA bands were cut out and DNA was purified using QIAquick gel extraction kit (Qiagen). DNA sequencing was performed using the fmol DNA sequencing system (Promega). α-33P dATP or α-35S dATP were used in sequencing reactions and 3 μl of sequencing reaction was run on 6% polyacrylamide gel containing 7 mol/l urea at 85 W for three to six hours.

DETECTION OF HphI POLYMORPHISM
Template DNA (100 ng) from peripheral blood was PCR amplified in a 40 μl volume using primers G10 and G11 (23), which flank the SSCP bands altered within the fingerprint which corresponds to the 364 bp DdeI fragment (arrows 1, 2) and the 221 bp HinfI fragment (arrows 3) containing exon 8. Lanes with enzyme headings represent amplified control DNA digested with that particular enzyme. (B) Sequence of exon 8 from a control (left panel) and IV1 (right panel). Primer sequences for amplifying exons 7–10 are given in Materials and methods. The C→T transition at position 924 is indicated in bold.
Silent mutation in LICAM and hydrocephalus

A
Normal

Exon 8
Genomic sequence
TGAGGGCAAGT
Protein sequence
QLKVGEEDDGGEYRCGLAESNLGRHAYYTYVEAAPYWHLKPSHLL

Exon 9
Genomic sequence
GTGCCGCCG
Protein sequence

B
K8715

Exon 8
Genomic sequence
GATGATGAGT
Protein sequence
QLKVGEEDDGGEYRCGLAESNLGRHAYYTYVEAAPYWHLKPSHLL

Exon 9
Genomic sequence
GTGCCGCCG
Protein sequence

Figure 3  Schematic representation of exon 8, intron 8, exon 9 organisation in LICAM. (A) Normal genomic sequence near the exon 8/intron 8 and intron 8/exon 9 boundaries. Protein sequence of the LICAM gene corresponding to this coding region is given below the genomic sequence. (B) Genomic sequence for the same region in the X chromosome in K8715 which carries the C924T alteration. The predicted amino acid sequence for the altered LICAM protein is given below the genomic sequence. The new 5' splice site is indicated by an arrow. In both (A) and (B), position 924 is indicated by *

exon 8 of the LICAM gene. The reaction contained 10 mmol/l Tris-HCl, pH 8.3, 50 mmol/l KCl, 1.5 mmol/l MgCl2, 50 μmol/l of each dATP, dCTP, dGTP, dTTP, 1 μmol/l of each primer, 2 U Taq DNA polymerase, and 0.44 μg TaqI Polymerase. Start antibody. The initial denaturation was performed at 94°C for five minutes and then followed by 35 cycles of 94°C for 30 seconds, 68°C for 30 seconds, and 72°C for one minute; the final extension was at 72°C for seven minutes. PCR product (16 μl) was digested using 8 U HpaI in a 24 μl total volume containing 50 mmol/l potassium acetate, 20 mmol/l Tris-acetate, 10 mmol/l magnesium acetate, 1 mmol/l dithiothreitol (pH 7.9) at 37°C for 12 hours. The digested samples were run on a mini non-denaturing gel containing 8% acrylamide and 15% glycerol at 170 volts and room temperature for one hour. The gel was stained in a 0.5 μg/ml ethidium bromide solution for five minutes and a gel picture was taken using an AlphaImager 2000 (Alpha Innotech Corporation, California).

Results

MUTATION ANALYSIS

The LICAM gene was screened for mutations using DNA from the proband in family K8715 (IV.1, fig 1A). Restriction endonuclease fingerprint (REF) analysis was used for 19 of the 28 exons. This method involved amplifying the 19 exons in five clustered groups: 5-6, 7-10, 11-14, 16-18, 20-24. REF analysis showed a fingerprint pattern consistent with an alteration in exon 8 (fig 2A). Sequence analysis of exon 8 detected a C→T transition at position 924 (fig 2B). This base substitution occurred in the third base in codon 308 which encodes a glycine, thereby creating a neutral mutation.

CELL CULTURE AND RNA ISOLATION

Amniocytes were maintained in MEM (Sigma) supplemented with 20% fetal bovine serum and 1% penicillin/streptomycin (Gibco). When the number of the cultured cells reached 5 million, isolation of the total RNA was accomplished using Rneasy mini kit (Qiagen).

CDNA PREPARATION AND RT-PCR

The total RNA was treated with DNase I before the reverse transcription (RT) reaction. The RT reaction was performed using 5 μg DNase I treated total RNA containing 20 mmol/l Tris-HCl (pH 8.4), 50 mmol/l KCl, 2.5 mmol/l MgCl2, 0.5 mmol/l of each dNTP, 0.01 mol/l DDT, 250 ng random hexamer primers, and 200 units of SuperScript II reverse transcriptase (Gibco BRL). The reaction conditions were according to the manufacturer's instructions.

cDNA mix (2 μl) from the RT reaction was used for RT-PCR in a 50 μl volume. The PCR reaction contained 10 mmol/l Tris-HCl (pH 8.3), 1.5 mmol/l MgCl2, 50 mmol/l KCl, 0.2 mmol/l each dATP, dCTP, dGTP, dTTP, 1 μmol/l primers of L5-10f (5'CGCTCGACACAACCTACAGACTACA3') and L5-10r (5'CGGTGTGGCGGCTTACACATT3'), 0.66 μg Taq polymerase start antibody, and 3 U Taq polymerase (Boehringer Mannheim). The PCR conditions were as follows: denaturation at 90°C for five minutes followed by 35 cycles of denaturation at 94°C for 30 seconds, annealing at 62°C for one minute and extension at 72°C for two minutes; the final extension was at 72°C for 10 minutes. The PCR products were run on 1.2% agarose gel containing 0.5 × TBE buffer and 0.5 μg/ml ethidium bromide.

CDNA PREPARATION AND RT-PCR

The total RNA was treated with DNase I before the reverse transcription (RT) reaction. The RT reaction was performed using 5 μg DNase I treated total RNA containing 20 mmol/l Tris-HCl (pH 8.4), 50 mmol/l KCl, 2.5 mmol/l MgCl2, 0.5 mmol/l of each dNTP, 0.01 mol/l DDT, 250 ng random hexamer primers, and 200 units of SuperScript II reverse transcriptase (Gibco BRL). The reaction conditions were according to the manufacturer's instructions.

cDNA mix (2 μl) from the RT reaction was used for RT-PCR in a 50 μl volume. The PCR reaction contained 10 mmol/l Tris-HCl (pH 8.3), 1.5 mmol/l MgCl2, 50 mmol/l KCl, 0.2 mmol/l each dATP, dCTP, dGTP, dTTP, 1 μmol/l primers of L5-10f (5'CGCTCGACACAACCTACAGACTACA3') and L5-10r (5'CGGTGTGGCGGCTTACACATT3'), 0.66 μg Taq polymerase start antibody, and 3 U Taq polymerase (Boehringer Mannheim). The PCR conditions were as follows: denaturation at 90°C for five minutes followed by 35 cycles of denaturation at 94°C for 30 seconds, annealing at 62°C for one minute and extension at 72°C for two minutes; the final extension was at 72°C for 10 minutes. The PCR products were run on 1.2% agarose gel containing 0.5 × TBE buffer and 0.5 μg/ml ethidium bromide.
Du, Dickerson, Aylsworth, et al

Figure 4  RT-PCR analysis of exons 6-10 of the L1CAM cDNA from amniocytes from the affected fetus (IV3, fig 1A). (A) Agarose gel electrophoresis of RT-PCR products generated from a control (lane 1), the fetus (lane 3), and a cDNA library (lane 5), using primers given in Materials and methods. Lane 7 is the PCR product generated using genomic DNA as a template. Lane 8 is a product generated using primers and RNA supplied with the SuperScript II reverse transcriptase kit (Gibco BRL) which was used as a positive control for the RT-PCR reactions. (B) Sequence analysis of the RT-PCR product from a control sample (left panel) and the affected fetus (right panel). The arrow in the right panel indicates the location of the 69 bp deletion in the mRNA from the affected fetus.

(G308G). The base substitution, however, did create a new HphI site. Analysis of the family showed that the presence of the site segregated with the HSAS (fig 1B). This was consistent with the previous haplotype analysis (data not shown).

The remaining nine exons (1-4, 19, 25-28) with their flanking intronic sequences were examined using SSCP. No sequence alterations were detected after duplicate analyses (data not shown).

THE C924T (G308G) ALTERATION AS A MUTATION

While the family was still being investigated, III.2 became pregnant. Amniocentesis showed the fetus to be a male (IV.3, fig 1A). Analysis of DNA isolated from the amniocytes indicated the X chromosome of the fetus carried the at risk haplotype (data not shown) and the HphI restriction site (fig 1B). Ultrasound at 23 weeks' gestation indicated borderline to mild ventriculomegaly. Physical examination of the fetus after delivery at 24 weeks showed relative
Silent mutation in LI CAM and hydrocephalus

macrocephaly and clapsed thumbs. Necropsy was refused. Based on the clinical features and the absence of any other identified sequence alteration in the LI CAM gene, the DNA sequence encompassing the C924T alteration was re-examined. Re-examination of the sequence showed that the C→T transition appeared to create a potential 5′ splice site 69 bp upstream from the normal exon 8/intron 8 splice site (fig 3). The new consensus sequence (TGttgagt) had a consensus value of 0.88 while the normal splice site (GGGttagg) had a value of 0.76. In theory, the new 5′ splice site would possibly result in 69 bp being eliminated from the mRNA (fig 3B). Furthermore, the absence of the HphI polymorphism in 200 X chromosomes indicated that this alteration was not a common polymorphism in the normal population.

Based on these findings and assumptions, RT-PCR was performed on total RNA isolated from amnioncytes of IV.3 using primers designed to amplify exons 6-10 in LI CAM mRNA. The results, presented in fig 4A, showed a band estimated to be about 559 bp while the band amplified from control LI CAM mRNA contained the expected band of 683 bp. Thus, 69 bp were apparently missing from the LI CAM transcript of IV.3 as predicted. Direct sequencing of the amplified fragment from IV.3 showed that the deletion involved the last 69 bp of exon 8 (fig 4B).

Discussion

Sequence analysis of exon 8 of LI CAM in a family with HSAS indicated a single difference between the proband's exon and the normal LI CAM exon 8, a thymine substituted for a cytidine (C924T). Both the normal codon 308 (GGC) and the proband's codon 308 (GGT) encode for glycine. This silent nucleotide substitution was not found to be a common variant in the normal population. In fact, this neutral mutation resulted in the activation of a new splice site 69 bp 5′ to the normal exon 8/intron 8 donor splice site (fig 3B), and segregated exclusively with HSAS in affected subjects and carriers in the family. The alternatively spliced mRNA codes for a LI CAM protein in which 23 amino acids (residues 308-330) are missing from the third Ig domain. This effectively eliminates the third near β sheet of the domain, undoubtedly severely affecting the activity of the LI CAM protein and its ability to induce neurone-neurone adhesion. 18 17

This mutation is notable for two reasons. First, the mutation differs from other splice site mutations in LI CAM and other genes that give rise to a pathogenic phenotype as it neither destroys the function of a normal splice site nor creates a new splice site within an intron. 18 A review of published reports found that mRNA splicing mutations account for 15% of all point mutations causing human genetic disease. 14 Of these, 87% (88/101) are point mutations within either the 5′ or 3′ splice site consensus sequence. The remaining point mutations (13%) lead to creation of novel splice sites. 18 Thus, the creation of a novel splice site resulting from the C924T transition in exon 8 in our family is representative of a very rare mutational event that accounts for about 2% of all point mutations.

The second feature of the C924T mutation in LI CAM is perhaps more important. The point mutation in codon 308 in our family is a classic case of a neutral or silent mutation since the normal codon (GGC) and the variant codon (GGT) both encode for glycine. To our knowledge, there have been at least six other reports of silent mutations giving rise to a pathogenic phenotype. Goldsmith et al 20 reported a T→A substitution in codon 24 of the β globin gene that did not alter the amino acid as both normal and variant codon encoded for glycine. Using a gene expression system, the authors proved that the substitution caused abnormal mRNA splicing in a patient with β thalassaemia. 19 The second report was of an A→G substitution in the third position of a lysine codon of the factor IX gene in a patient with haemophilia B. 20 Although the altered sequence had a splice donor consensus value of 1.00, the authors did not prove that alternate mRNA splicing occurred in their patient. As no other mutation was found, the authors presumed the silent A→G substitution to be pathogenic. 20 Baumbach et al 21 reported a neutral mutation (C766T) in the third position of codon 236 of the growth hormone receptor (GH-R) gene. The mutation was shown to create a cryptic splice site 63 bp 5′ to the normal exon 7 splice site, thereby eliminating 21 amino acids from the GH-R. Homozygosity for this mutation results in Laron's syndrome in a genetic isolate in the Bahamas. 22 A neutral G to A transition in the third position of the alanine at codon 344 (A344A) in FGFR2 (fibroblast growth factor receptor II) has been observed in four unrelated Crouzon syndrome patients. 23 24 Reardon et al 25 thought this base substitution would result in a new acceptor splice site. Jabs et al 26 suggested the mutation could create either an acceptor or donor splice site. Recently, Li et al. 24 using RT-PCR and RNA isolated from cell cultures of their affected subject, showed that the A344A mutation created a new donor splice site. This finding was confirmed independently by Del Gatto and Brethnach 27 using cloned genes in cultured HeLa cells. Richard and Beckman 28 found a silent C→T substitution in codon 624 of the calpain gene (CANP3) in a homozygous patient from one family with limb girdle muscular dystrophy (LGMD2A). Analyses of illegitimate transcripts showed an abnormal product which, upon sequencing, was found to be missing the last 44 bases of exon 16. Thus, this neutral mutation results in the gain of a new donor splice site. Lastly, a synonymous substitution (G2142A; S649S) in exon 11 presumably creates a A→G acceptor splice site giving rise to a truncated RET proto-oncogene protein in members of a family with Hirschsprung disease. 27

Most certainly the C924T mutation in LI CAM would have appeared as an altered transcript using RT-PCR. Indeed, at least two neutral mutations have been detected in this manner, both involving the XNP gene in two
unrelated male patients with the ATR-X syndrome.\(^ 28^\)\(^ 29\) In neither paper, however, was the "neutral" nature of the mutation stressed since the method of detection involved RT-PCR. Unfortunately, in many instances, use of genomic DNA rather than mRNA is better suited for mutational analysis. For one thing, a cell line is not always available for a dead proband while DNA may have been obtained at some time before death. Second, not all genes are expressed in lymphocytes which are the most readily accessible source of mRNA.\(^ 30\) Finally, some genes are quite large, making it difficult if not problematical to obtain full length mRNA for analysis.

The silent mutation in LI-CAM that was identified in our family with HSAH has enormous implications for mutation screening in any gene. The concept of a synonymous or neutral mutation should be altered when one is screening a candidate gene for disease causing mutations. A neutral base substitution can clearly have a profound effect on mRNA splicing or even on codon usage, even though the nucleotide replacement is irrelevant with respect to the amino acid encoded by the codon. Thus, it would be wise to remember the words of Sherlock Holmes: "We must fall back upon the old axiom that when all other contingencies fail, whatever remains, however improbable, must be the truth."\(^ 32\)

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\(\textit{Du, Dickerson, Aylworth, et al.}\)
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Y Z Du, C Dickerson, A S Aylsworth and C E Schwartz

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