A Novel Locus for Generalized Epilepsy With Febrile Seizures Plus in French Families

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Background: Generalized epilepsy with febrile seizures plus (GEFS⁺) is a familial autosomal dominant entity characterized by the association of febrile and afebrile seizures. Mutations in 3 genes—the sodium channel α1 subunit gene (SCN1A), the sodium channel β1 subunit gene (SCN1B), and the γ2 GABA receptor subunit gene (GABRG2)—and linkage to 2 other loci on 2p24 and 21q22 have been identified in families with GEFS⁺, indicating genetic heterogeneity.

Objectives: To localize by means of linkage analysis a new gene for GEFS⁺ in a large family with 11 affected members and to test the new locus in 4 additional families with GEFS⁺.

Design: Family-based linkage analysis.

Setting: University hospital.

Patients: Five French families with GEFS⁺ and at least 7 available affected members with autosomal dominant transmission. All the patients had febrile seizures and/or afebrile generalized tonic-clonic seizures or absence epilepsy.

Main Outcome Measures: We analyzed 380 microsatellite markers and conducted linkage analysis.

Results: In the largest family, a 10-cM-density genome-wide scan revealed linkage to a 13-Mb (megabase) interval on chromosome 8p23-p21 with a maximum pairwise logarithm of odds (LOD) score of 3.00 (at θ = 0) for markers D8S351 and D8S530 and a multipoint LOD score of 3.23. A second family with GEFS⁺ was also possibly linked to chromosome 8p23-p21 and the region was narrowed to a 7.3-Mb candidate interval, flanked by markers D8S1706 and D8S549. We have not, so far, identified mutations in the coding exons of 6 candidate genes (MTMR9, MTMR7, CTSB, SGCG, SG223, and ATP6V1B2) located in the genetic interval.

Conclusions: We report a sixth locus for GEFS⁺ on chromosome 8p23-p21. Because no ion channel genes are located in this interval, identification of the responsible gene will probably uncover a new mechanism of pathogenesis for GEFS⁺.

Arch Neurol. 2008;65(7):943-951
have been reported: febrile convulsions 1 gene (FEB1) on chromosome 8q13-q21,12 FEB2 on 19p13,13 and FEB5 on 6q22-q24.14

Herein, we ascertained 5 French families with GEFS+/H11001 without linkage to FS or GEFS+/H11001 loci previously reported. Conclusive linkage to chromosome 8p23-p21 was obtained in a large multigenerational family, and, in a second family, linkage to the same locus was suggested.

**METHODS**

**FAMILIES**

Five French families with at least 7 affected members for whom DNA was available were identified during a national campaign organized by the Association de Recherche sur la Génétique des Épilepsies and supported by the French Génethon Center. Simplified pedigrees of the families are given in Figure 1. All members of the families underwent clinical assessment using a detailed questionnaire. Information was also obtained retrospectively from medical records. Informed consent was obtained from all the participants or from their legal representatives.

**GENOTYPING**

Genomic DNA was isolated from blood lymphocytes using standard procedures. Genome scans for families 15173 and 12402 were performed at the Centre National de Génotypage. The set consisted of 400 fluorescent microsatellite markers (including 20 for chromosome X that were not tested because of a male-to-male transmission in each family) selected from the Génethon human linkage map that cover the entire human genome with a resolution of approximately 10 cM. Subsequent markers used for fine mapping were genotyped as previously reported.4

**SEQUENCING**

A mutation search by means of direct sequencing was performed in the coding exons and flanking splice sites of 6 candidate genes (MTMR9 [OMIM 606260], MTMR7 [OMIM 603562], CTSB [OMIM 116810], SGCZ [OMIM 608113],

**LINKAGE ANALYSIS**

We calculated parametric pairwise logarithm of odds (LOD) scores using the MLINK program and multipoint LOD scores using the Allegro program, assuming an autosomal dominant trait with a disease allele frequency of 0.0001, penetrance of 0.60, equal recombination fractions in males and females, and equal frequencies for the alleles observed in the families. These variables were also used to calculate theoretical maximum LOD scores using the “affected only” method. At-risk individuals I:4 (67 years old), II:8 (40 years old), III:3 (21 years old), and III:6 (11 years old) of family 15173 and at-risk individuals I:2 (60 years old), I:6 (69 years old), II:3 (33 years old), II:5 (40 years old), and II:6 (38 years old) of family 12402 were clinically normal and were, thus, considered to be unaffected. All individuals who experienced 1 or more FSs or afebrile seizures were considered to be affected.
SG223, and ATP6V1B2 [OMIM 606939]). The genomic organization of the candidate genes and the primer sequences were obtained from the University of California Santa Cruz Web site (http://genome.ucsc.edu/) and are available on request. Polymerase chain reaction products were sequenced on both strands on an ABI3730 automatic sequencer using the Big Dye Terminator Cycle sequencing kit (Applied Biosystems, Foster City, California).

RESULTS

CLINICAL CHARACTERISTICS

Family 15173

We identified a 3-generation kindred with 22 family members: 11 affected individuals with a history of at least 1 FS, 3 of whom also had afebrile seizures, 2 unaffected obligate carriers, 5 unaffected individuals, and 5 married-in individuals (Table 1 and Figure 2). No blood sample was available for patient II:7, who experienced 1 FS.

The FSs began at age 8 months to 1.5 years and persisted beyond age 6 years in 4 patients (II:1, 11 years; II:3, 10 years; II:4, 11 years; and III:7, 6.5 years). The number of FSs was particularly high in 3 patients in generation II despite the prescription of antiepileptic drugs (20 episodes of FS in patient II:1 and 15 in patient II:3). In addition, individual II:4 had an association of FSs and afebrile seizures; he experienced a total of 50 seizures between ages 1 to 11 years. Two individuals had complex FSs: II:4 had 1 prolonged FS followed by transitory hemiplegia at 1 year of age and another prolonged FS without a motor deficit 6 months later, and III:7 had several FSs lasting more than 30 minutes, 1 of which was followed by transitory hemiplegia.

Three individuals (II:1, II:4, and III:5) also had afebrile seizures. Individual II:1 had 1 generalized tonic-clonic seizure (GTCS) during a period of sleep deprivation and alcohol intake, another during treatment with antidepressant.

Table 1. Clinical Features of Families 15173 and 12402 With Linkage to Chromosome 8p23-p21

<table>
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<tr>
<th>Patient</th>
<th>Pedigree/Sex/Age, y</th>
<th>First Occurrence Last Occurrence No. Complex</th>
<th>First Occurrence Last Occurrence Pattern No. Interictal EEG</th>
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<td>30 y 40 y</td>
<td>GTCSs 2</td>
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<tr>
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<td>1 y 40 y</td>
<td>GTCSs</td>
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<td>Not app</td>
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<td>15 y 50 y</td>
<td>Partial (right frontotemporal)</td>
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<td>Not app Not app Not app Not app</td>
<td>13 y Until death</td>
<td>Partial (right temporal) 1/mo</td>
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</table>

Abbreviations: EEG, electroencephalogram; GTCS, generalized tonic-clonic seizure; NA, not available; Not app, not applicable; SW, spike wave.

One seizure occurred with sleep deprivation and alcohol intake, another during treatment with antidepressant.

Mixture of febrile and afebrile seizures.

(Reprinted) Arch Neurol/Vol 65 (No. 7), July 2008 www.archneurol.com

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his last FS at age 8 years. He also had the highest number of or complex FSs, with late childhood onset and a 3-Hz spike wave pattern recorded by means of EEG. In addition, in patient I:4, afebrile GTCS occurred during the same period as FSs and stopped by age 9 years. Three family members who died before the study had epilepsy with no reported FSs: the first had absence epilepsy (I:10), and the other 2 had temporal lobe seizures beginning in adolescence (I:11 and II:9). The cause of death was undetermined for patient I:11. Absence epilepsy in this family was of late onset (I:10, 7 years; II:2, 10 years; and II:4, 16 years) and highly photosensitive. No lesions were observed by means of magnetic resonance imaging (MRI) in patients with temporal lobe epilepsy. Psychomotor development was normal in all affected family members. Performance in school of patient II:2 declined from age 12 years; she was oriented toward a profession at age 16 years.

**Family 12402**

In this second 4-generation family, 11 individuals were affected (data were available for only 10 individuals) (Table 1 and Figure 3). Four individuals were dead at the time of this study; DNA was, therefore, available for only 7 individuals. Six of these 7 patients had FSs, with ages at onset ranging from 9 to 24 months. All FSs stopped before or at age 6 years except in 1 patient (I:8) who had his last FS at age 8 years. He also had the highest number of episodes (>10 FSs). The FSs were simple in 4 patients, consisting of brief GTCS, but in 2 patients (I:4 and II:7) they lasted up to 15 to 20 minutes (complex FSs).

Two patients (II:2 and II:4) also had absence epilepsy, with late childhood onset and a 3-Hz spike wave pattern recorded by means of EEG. In addition, in patient I:4, afebrile GTCS occurred during the same period as FSs and stopped by age 9 years. Three family members who died before the study had epilepsy with no reported FSs: the first had absence epilepsy (I:10), and the other 2 had temporal lobe seizures beginning in adolescence (I:11 and II:9). The cause of death was undetermined for patient I:11. Absence epilepsy in this family was of late onset (I:10, 7 years; II:2, 10 years; and II:4, 16 years) and highly photosensitive. No lesions were observed by means of magnetic resonance imaging (MRI) in patients with temporal lobe epilepsy. Psychomotor development was normal in all affected family members. Performance in school of patient II:2 declined from age 12 years; she was oriented toward a profession at age 16 years.

**Families 16923, 17516, and 15635**

In family 16923 (8 affected individuals), FSs were simple, infrequent (n = 1-4), and stopped before age 6 years. Two patients had afebrile GTCS at ages 3 and 6 years and were fully controlled with antiepileptic drug monotherapy.

In family 17516, 10 patients had FSs, which were simple, recurred 2 to 3 times except in 1 patient (n = 8), and stopped before 6 years of age. Three patients had afebrile GTCS that occurred once in 1 patient (age at oc-
currence unknown) but recurs 3 times in the other 2 patients, at ages 4 and 9 years. Seizures in these 2 patients were fully controlled by antiepileptic drug monotherapy.

In family 15635, 9 patients had simple FSs that stopped before age 6 years. One patient had 1 afebrile GTCS at 12 years of age, and 2 half-brothers developed rare partial seizures in adolescence. Their brain MRIs were normal. They were not pharmacoresistant at the time of this study and were 17 and 20 years of age. Their mother had isolated FSs. All affected members of these 3 families had normal psychomotor development.

**LINKAGE ANALYSIS**

**Exclusion of Known FS and GEFS+ Loci**

We first examined whether any of the 5 families presented linkage to previously reported loci for GEFS+ and FS. Negative pairwise LOD scores were obtained in all 5 families, excluding the following loci as the cause of the disease: 19q13 (SCN1B), 2q24 (SCN1A), 3q34 (GABRG2), 2p24 (GEFS+), 21q32 (GEFS+5), 8q13-21 (FEB1), 19p13 (FEB2), and 6q22-24 (FEB5) (data not shown).

We then calculated the maximal theoretical pairwise LOD scores to determine which families were sufficiently informative to map a new gene. Using the affected only method, at Θ = 0.0, family 15173 (DNA available for 10 affected members) had a \( Z_{\text{max}} \) of 3.06, family 12402 (DNA available for 7 affected members) had a \( Z_{\text{max}} \) of 1.7, family 16923 (DNA available for 8 affected members) had a \( Z_{\text{max}} \) of 2.7, family 15635 (DNA available for 7 affected members) had a \( Z_{\text{max}} \) of 1.8, and family 17516 (DNA available for 7 affected members) had a \( Z_{\text{max}} \) of 1.8.

**Genomewide Scan in Family 15173**

A genomewide scan of all autosomes was conducted with 380 microsatellite markers in family 15173, the only family in which the theoretical maximal pairwise LOD score reached the threshold value of 3.00 in linkage analyses. Calculation of pairwise LOD scores revealed 20 of 380 noninformative markers, which were excluded by means of haplotype reconstruction (data not shown). In addition, positive pairwise LOD scores were obtained for 6 markers (D10S208, D15S130, D16S423, and D18S61 and the 2 adjacent markers D8S351 and D8S550). Fine mapping with additional markers (D10S183, D10S199, D15S331, D15S207, D16S3030, D16S3184, D18S1125, and D18S386) excluded the regions on chromosomes 10, 15, 16, and 18. In contrast, genotyping of 14 additional markers (D8S264, D8S262, D8S1742, D8S277, D8S1706,
Table 2. Pairwise LOD Scores for 15 Markers on Chromosome 8p23-p21 From Telomere (Top) to Centromere (Bottom) for Family 15173 and Family 12402

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</table>

Abbreviations: AO, “affected only” method; AR, all-at-risk individuals; LOD, logarithm of odds.

Maximum LOD scores.

D8S351, D8S503, D8S552, VNTR22TG, D8S602, D8S639, D8S261, VNTR25CA, and D8S258) confirmed linkage on chromosome 8p23-p21. Maximal pairwise LOD scores were 3.00 at \( \Theta = 0.0 \) for D8S351 and D8S550 calculated using the affected only method (Table 2). The scores recalculated with inclusion of all-at-risk individuals yielded pairwise LOD scores slightly below 3.00 (Table 2) because of incomplete penetrance. Under these conditions, multipoint analysis with 15 markers generated a maximal LOD score of 3.23 in the interval flanked by D8S351 and VNTR25CA; the remaining genome was excluded (data not shown).

Haplotype reconstruction showed that all patients shared a common haplotype encompassing markers D8S351 (chromosome 8: 8 714 310-8 914 565) to VNTR25CA (chromosome 8: 19 887 367-19 887 416) (Figure 2). The boundaries of this interval were defined by a recombination between markers D8S1706 and D8S351 in patient III:9 and a recombination between markers VNTR25CA and D8S258 in patient II:1. This genetic interval (D8S1706-D8S258) responds to a 13-Mb (megabase) large region located on chromosome 8p23.1-p21.3, according to the University of California Santa Cruz Genome Browser (http://genome.ucsc.edu/).

Genotyping of the Other Families

After establishing linkage at chromosome 8p23-p21 in family 15173, the 4 smaller families with GEFS\(^+\) were analyzed. We obtained pairwise LOD scores less than –2 for 8p23-p21 markers in families 16923, 17516, and 15635 (data not shown) but positive pairwise LOD scores in family 12402. The maximum pairwise LOD score was 2.03 for D8S262 calculated using the affected only method (Table 2). Multipoint analysis gave maximal LOD scores of 2.14 in the interval D8S262 to D8S1706 when all at-risk individuals were included (data not shown). Haplotype reconstruction showed that the 7 affected patients and the obligate carrier I:6 had a common haplotype of 12 Mb (D8S262-D8S352), defined by a recombination between markers D8S264 and D8S262 in patient I:4 and a recom-
We recruited 5 French families with a phenotype compatible with GEFS+. The disorder segregated as an autosomal dominant trait with incomplete penetrance in all families. Most patients in the 5 families experienced simple FSs (93%) and some afebrile seizures (34%), mostly GTCSs or absence seizures. None of the families were linked to the previously reported GEFS+ and FS loci.

A 10-cM-density genomewide scan in the most informative family (15173) revealed a unique region with significant maximum pairwise LOD scores of 3.00 and multipoint LOD scores of 3.23 on chromosome 8p23-p21, which strongly suggests that we have identified a new locus for GEFS+. Because all other regions of the genome were excluded, we believe that the responsible gene is probably localized in this interval. Furthermore, when the 4 other families were then tested for linkage to this novel locus, we obtained evidence of probable linkage in 1 family (12402) to a region of 7.3 Mb that overlapped the locus on chromosome 8p23-p21 in family 15173. Although a linkage in family 12402 might have been obtained by chance, the absence of any other positive region is in favor of a conclusive linkage. Furthermore, all affected family members shared the same haplotype, and no polymorphisms or were found in 100 matched French controls.

**COMMENT**

We described in detail the clinical features of the 2 families, 15173 and 12402, linked to the new locus on chromosome 8p23-p21. In family 15173, all affected individuals experienced FSs, and 3 of 11 also experienced tonic I:8 of family 12402 (Table 3). Some sequence variants were identified in each gene, but they were already mentioned in databases as polymorphisms or were found in 100 matched French controls.

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### Table 3. Candidate Genes Explored in the 8p23-p21 Interval

<table>
<thead>
<tr>
<th>Gene Symbol</th>
<th>Position in Bases at Chromosome 8</th>
<th>Protein</th>
<th>Potential Role in Epilepsy</th>
</tr>
</thead>
<tbody>
<tr>
<td>SG223</td>
<td>8 212 676-8 276 667</td>
<td>Tyrosine-protein kinase</td>
<td>Homology with rat pragma of Rnd2, which is involved in the migration of pyramidal neurons</td>
</tr>
<tr>
<td>MTMR9</td>
<td>11 179 410-11 223 064</td>
<td>Myotubularin-related protein 9</td>
<td>Brain binding partner of MTMR7</td>
</tr>
<tr>
<td>CT5B</td>
<td>11 739 236-11 744 571</td>
<td>Cathepsin B</td>
<td>Upregulated after seizures provoked by hippocampal kindling</td>
</tr>
<tr>
<td>SG52</td>
<td>13 991 744-15 140 163</td>
<td>Sarcoglycan zeta</td>
<td>Abundant in the brain</td>
</tr>
<tr>
<td>MTMR7</td>
<td>17 203 534-17 250 961</td>
<td>Myotubularin-related protein 7</td>
<td>Brain binding partner of MTMR9</td>
</tr>
<tr>
<td>ATP6V1B2</td>
<td>20 098 984-20 123 487</td>
<td>ATP synthase subunit B</td>
<td>High levels of expression in the brain</td>
</tr>
</tbody>
</table>

*a The MTMR7 and ATP6V1B2 genes are located outside of the common interval of both families but were initially screened because they were included in the interval segregating in family 15173.

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**Genomewide Scan in Family 12402**

To confirm that FSs in family 12402 were caused by a mutation in a gene located in this 8p23-p21 locus and that it did not segregate with the disease by chance, we performed a whole genome scan using 380 microsatellite markers. Calculation of pairwise LOD scores showed that 55 of 380 markers were not informative; they were excluded by means of haplotype reconstruction. Two adjacent noninformative markers, D16S3062 and D16S3046, were excluded after saturation of the interval with markers D16S3062, D16S405, D16S3014, D16S3017, and D16S412. In addition, positive pairwise LOD scores were obtained for 11 of 380 markers (D15S226, D3S1534, D5S422, D6S462, D8S552, D13S171, D17S87, D19S902, D19S418, and D22S427). Areas on chromosomes 1, 4, 6, and 17 were excluded by means of haplotype reconstruction. Additionally, microsatellite markers (D5S142, D13S260, D13S1493, D13S1293, D13S1246, D19S8597, D19S556, D19S545, D19S892, D19S879, and D22S427) were further genotyped to exclude the regions on chromosomes 5, 13, 19, and 22. Because the remaining genome was excluded, we concluded that family 12402 was likely linked to this new locus on 8p23-p21.

### SEQUENCING OF CANDIDATE GENES

According to Ensembl (http://www.ensembl.org), the genetic interval of interest spans 7.3 Mb and contains approximately 80 known genes. The region contains no ion channel or neurotransmitter genes. We prioritized the genes according to their patterns of expression in the brain and putative functions and sequenced 6 candidate genes—MTMR9, MTMR7, CT5B, SG52, SG223, and ATP6V1B2—in patient II:4 of family 15173 and in patient I:8 of family 12402 (Table 3). Some sequence variants were identified in each gene, but they were already mentioned in databases as polymorphisms or were found in 100 matched French controls.
Afebrile seizures. Four patients reported the occurrence of FSs after age 6 years and experienced many episodes of FS, or a combination of FSs and afebrile seizures, which were defined as “FS plus.” Family 12402 had 11 affected members. Febrile seizures were present in 6 patients, complex in 2, and lasted until 8 years of age in 1 patient. The FSs recur more than 5 times in half of these patients, with more than 10 episodes in patient 18. In addition, 3 patients also developed afebrile seizures: GTCSs in 1 and absence epilepsy in 2. Three patients had epilepsy with no history of FSs (partial epilepsy in 2 and absences in the third). Frequent absences have already been reported in other families with GEFS. Both families corresponded to the description of the familial GEFS context.

In the literature, phenotype-genotype correlations have been reported in families with GEFS. For example, the proportion of patients with FS+ was higher in families with SCN1A mutations than in families with GABRG2 mutations. In the present study, families with linkage to the 8p23-p21 locus had a higher proportion of patients with FS+ than did families without linkage (28% vs 7%). Furthermore, there was a high rate of recurrence of FSs in 53% of patients from families linked to chromosome 8p23-p21, whereas frequent recurrence was reported in only 1 of 27 patients (4%) from families excluded for this locus. Complex FSs were observed in only the 2 families linked to chromosome 8p23-p21 (4 of 17 patients). Afebrile seizures co-occurred with FSs in at least 1 patient per family except for family 15635. Because none of the patients had FS+ in this family, it was considered to be a family with epilepsy and FSs rather than typical GEFS.

The new locus contains approximately 80 known genes, none of which are known or predicted to encode ion channels, neurotransmitter receptors, or proteins homologous to others involved in epilepsy. Twenty of the genes encode unknown proteins. To identify the causative gene, we sequenced the coding regions and flanking splicing sites of genes encoding proteins expressed in brain or with a putative role in epilepsy (Table 3). So far, no causative mutations have been identified in the coding regions of the following genes: CTSSB, SGC2, SG223, MTMR9, MTMR7, and ATP6V1B2. We hope that elucidation of the function of the unknown proteins encoded by genes located in the interval will provide new candidates to screen for FSs and epilepsy. In parallel, we will search for rearrangements that are reported to be frequent in the 8p23 region.

Identification of additional families with GEFS+ linked to the 8p23-p21 region will reinforce linkage to this new locus, the sixth to be identified for this disorder. This might also narrow the candidate interval and consequently decrease the number of candidate genes to explore. So far, 3 genes have been implicated in GEFS+: SCN1A, SCN1B, and GABRG2. Except for leucine-rich glioma inactivated 1 (LGI1) and EF-hand domain (C-terminal) containing 1 (EFHC1), all other genes involved in idiopathic monogenic epilepsies encode ion channel or neurotransmitter receptor genes. It is, thus, interesting that the novel locus reported herein does not contain genes with functions similar to those already implicated in epilepsy. Identification of the responsible gene in 8p23-p21 might bring to light a new mechanism involved in epileptogenesis.