CUX2 mutations in temporal lobe epilepsy; increased kainate susceptibility and excitatory input to hippocampus in deficient mice

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Abstract

CUX2 gene encodes a transcription factor that controls neuronal proliferation, dendrite branching and synapse formation, locating at the epilepsy-associated chromosomal region 12q24 that we previously identified by a genome-wide association study (GWAS) in Japanese population. A CUX2 recurrent de novo variant p.E590K has been described in patients with rare epileptic encephalopathies and the gene is a candidate for the locus, however the mutation may not be enough to generate the genome-wide significance in the GWAS and whether CUX2 variants appear in other types of epilepsies and physiopathological mechanisms are remained to be investigated. Here in this study, we conducted targeted sequencings of CUX2, a paralog CUX1 and its short isoform CASP harboring a unique C-terminus on 271 Japanese patients with a variety of epilepsies, and found that multiple CUX2 missense variants, other than the p.E590K, and some CASP variants including a deletion, predominantly appeared in patients with temporal lobe epilepsy (TLE). Human cell culture and fly dendritic arborization analyses revealed loss-offunction properties for the CUX2 variants. Cux2- and Casp-specific knockout mice both showed high susceptibility to kainate, increased excitatory cell number in the entorhinal cortex, and significant enhancement in glutamatergic synaptic transmission to the hippocampus. CASP and CUX2 proteins physiologically bound to each other and co-expressed in excitatory neurons in brain regions including the entorhinal cortex. These results suggest that CUX2 and CASP variants contribute to the TLE pathology through a facilitation of excitatory synaptic transmission from entorhinal cortex to hippocampus.

Introduction

Cux2 gene encodes a homeobox transcription factor CUX2 that is predominantly expressed in progenitor cells of the subventricular zone in mouse embryos and pyramidal neurons of the upper neocortical layers (II–IV) in adult mice (Zimmer et al., 2004). CUX2 is also expressed in Reelinpositive neurons distributed throughout the layers II–IV in postnatal day 0 (P0) mice (Cubelos et al., 2008a). CUX2 controls neuronal proliferation, dendrite branching, spine morphology and synapse formation (Cubelos et al., 2008b; Cubelos et al., 2010). We recently reported a genomewide association study (GWAS) on 1,825 Japanese patients with variable epilepsies which identified an associated region at chromosome 12q24.11 – 12q24.13 harboring 24 transcripts including CUX2 gene (Suzuki et al., 2021). In the region, CUX2 is only gene which has been reported to be relevant for epilepsy; a recurrent de novo variant (c.1768G>A, p.E590K) has been identified in patients with rare epileptic encephalopathies (EEs) (Chatron et al., 2018; Barington et al., 2018). CUX2 is therefore one of the most promising candidate genes in this 12q24 epilepsy-associated region, but the mutation reported in rare EEs may not be enough to explain the association detected in the Japanese GWAS study.

To investigate whether *CUX2* and its paralogues' mutations are involved in other types of epilepsies, here we performed targeted sequencing of *CUX2*, its paralog *CUX1*, and *CASP* which is a short isoform of *CUX1* with a unique C-terminus, in Japanese patients with variable epilepsies including idiopathic and symptomatic ones, and identified their variants predominantly in patients with temporal lobe epilepsy (TLE), the most common but intractable form of epilepsy (Téllez-Zenteno *et al.*, 2012). Analyses in human cultured cell and transgenic fly showed that the variants have loss-of-function effects. *CUX2* and *CASP* deficiencies in mice increased their seizure susceptibilities to a convulsant, kainate, which has long been used to

generate TLE animal models (Lévesque *et al.*, 2013). Histological and electrophysiological analyses revealed increases of excitatory neuron numbers in entorhinal cortex and those in excitatory input to hippocampus in both mice, proposing a circuit mechanism for the pathology of TLE.

Materials and Methods

Subjects

Genomic DNAs from 271 Japanese patients with a variety of epilepsies (see Results section for contents) and 311 healthy Japanese volunteers recruited by cooperating hospitals were used for the targeted sequencing analyses for *CUX2*, CUX1 and *CASP*. In addition, DNAs from independent 69 Japanese patients with TLE were used to check the frequency of c.3847G>A (p.E1283K) variant in *CUX2* gene. The patients' DNAs analyzed in our GWAS (Suzuki *et al.*, 2021) were not used in the present study, because their epilepsy subtype information [TLE, etc.] were not available for those materials.

Targeted sequencing

Genomic DNA samples were amplified with the illustra GenomiPhi V2 DNA Amplification Kit (GE Healthcare). We designed PCR primers to amplify all coding regions of *CUX2* (NM_015267), *CUX1* (NM_001202543 and first exon of NM_181552), unique C-terminus region of *CASP* (NM_001913), and amplified genomic DNA by PCR using the PrimeSTAR HS DNA Polymerase (TaKaRa) or KOD-plus Ver. 2 (TOYOBO). The PCR products were purified using ExoSAP-IT PCR product Cleanup (Thermo Fisher Scientific) and analyzed by direct sequencing using the ABI PRISM 3730xl Genetic Analyzer. All novel variants identified in amplified DNA by GenomiPhi were verified by direct-sequencing of patients' genomic DNAs. Primer sequences and PCR conditions are available upon request.

Quantification of mRNA – described in the Supplemental Methods.

Domain search

Domain searches in CUX2, CUX1, and CASP amino acid sequences were performed using the SMART database.

Expression constructs and mutagenesis – described in the Supplemental Methods.

Cell imaging – described in the Supplemental Methods.

Drosophila stocks and crosses – described in the Supplemental Methods.

TUNEL assay in flies – described in the Supplemental Methods.

Mice

Cux2 knockout (KO) mouse was obtained from Texas A&M Institute for Genomic Medicine (TIGM) as cryopreserved sperm of heterozygous Cux2 gene trap mouse (129/SvEv × C57BL/6 background) derived from the gene trapped clone OST440231. Live animals were produced by in vitro fertilization at the Research Resources Division (RRD) of the RIKEN Center for Brain Science. The heterozygous mice were maintained on the C57BL/6J (B6J) background, and the resultant heterozygous mice were interbred to obtain wild-type (WT), heterozygous, and homozygous mice. Genotyping was carried out as described previously (Iulianella et al., 2008). Casp-specific KO mice were generated using the CRISPR/Cas system. Plasmid vector pX330-U6-Chimeric BB-CBh-hSpCas9 was a gift from Dr. Feng Zhang (Addgene plasmid # 42230). A pair of oligo DNAs corresponding to Casp-gRNA (TTTCCATCATCTCCAGCCAA AGG) in exon 17 of Casp (NM 198602) was annealed and ligated into pX330-U6-Chimeric BB-CBhhSpCas9. For knock-out mouse production, Cas9 mRNAs and Casp-gRNA were diluted to 10 ng/µL each. Further, B6J female mice and ICR mouse strains were used as embryo donors and foster mothers, respectively. Genomic DNA from founder mice was extracted, and PCR was performed using gene-specific primers (CRISPR check F: 5'-

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GGAGCTATTGTAGGACATCACAGA-3' and CRISPR check R: 5'-

CCCCAGTGTTCTTTACTTTGAGTT-3'). PCR products were purified using ExoSAP-IT PCR product Cleanup and analyzed by direct sequencing using the ABI PRISM 3730xl Genetic Analyzer. The heterozygous mutant mice (c.1514^1515ins.TT, p.S506fs) were crossbred with B6J mice, and the resultant heterozygous mice were interbred to obtain WT, heterozygous, and homozygous mutant mice. The frame-shift mutation was confirmed by sequence analysis of cDNA from mutant mouse brains.

Seizure susceptibility in mice – described in the Supplemental Methods.

CUX2 antibody generation

To generate a rabbit polyclonal antibody to CUX2, a fusion protein was prepared, in which the glutathione-S-transferase (GST) protein was fused to the a.a. residues 356 to 415 of mouse CUX2 which has been used in the previous study's antibody generation (Gingras *et al.*, 2005). Rabbits were injected with 0.5 mg of purified GST fusion protein in Freund's complete adjuvant, boosted five times with 0.25 mg of protein, and serum collection at 1 week following the last boost. Polyclonal antibody was purified by affinity chromatography. The serum was passed through a GST affinity column ten times, and the flow-through was then applied to a GST-CUX2 (356-415 a.a.) affinity column to isolate antibodies.

Histological analyses – described in the Supplemental Methods.

In vitro electrophysiology – described in the Supplemental Methods.

Co-immunoprecipitation – described in the Supplemental Methods.

Statistical analysis

In the *in vitro* and *in vivo* experiments, data are presented as Box-and-whisker plots or means ± s.e.m. The boxes show median, 25th and 75th percentiles, and whiskers represent minimum and maximum values. *P*-value for p.E1283K was calculated using the Cochrane-Armitage trend test.

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One-way or two-way ANOVA, Tukey's multiple comparison test, Chi-square test, or Kolmogorov-Smirnov test were used to assess the data as mentioned in the figure legends. Statistical significance was defined as P < 0.05.

Study approval

Human study: All subjects and control individuals provided written informed consent to participate in this study in accordance with approval by ethical committees at the RIKEN Centers for Brain Science.

Animal study: All experimental protocols were approved by the Animal Experiment Committee of the RIKEN Center for Brain Science.

Results

CUX2 variants predominantly appeared in Japanese TLE patients

We carried out a targeted sequencing of CUX2 in 271 Japanese patients with variable epilepsies consisting of 116 idiopathic and 155 symptomatic ones. Symptomatic epilepsy samples contained 68 TLEs, which were further divided to 57 mesial TLE (mTLE) and 11 lateral TLE (ITLE). We identified five CUX2 heterozygous missense variants in nine unrelated patients (Figure 1A, Table 1 and Supplemental Note). Notably, eight of the nine patients with CUX2 variants had TLE (one ITLE and seven mTLE patients). All of the mTLE patients showed hippocampal sclerosis. Three (p.R34W, p.P454L, and p.W958R) out of the five variants were absent or rare (< 0.5%) in the in-house Japanese control individuals (in-house controls) and databases and were also predicted to be damaging (Table 1). The p.E1283K variant, a frequent variant predicted to be less damaging, appeared in Japanese TLE patients at a significantly high ratio [$P = 5.93 \times 10^{-3}$, OR = 6.94, 95% CI = 1.39-34.61 calculated in 137 (above-mentioned 68 + additional independent 69) TLE patients vs 311 in-house controls] and therefore we hypothesized it may also be a genetic contributor for TLE. The p.D337N variant appeared in one case with TLE and controls with a similar allele frequency.

Loss-of-function effects of CUX2 variants

To investigate the functional impacts of *CUX2* variants appeared in patients with epilepsy (Figure 1A, Table 1), we transfected HeLa.S3 cells with expression constructs of wild-type (WT) and the five variants (p.R34W, p.D337N, p.P454L, p.W958R, and p.E1283K). It revealed that, while CUX2-WT protein was limited to and well distributed in the nuclei, variant proteins (except for p.R34W) leaked out to the cytoplasm or aggregated (Figure 1B, C and Figure S1A).

CUX2 is an ortholog of Drosophila melanogaster cut, which promotes dendritic arbor morphological complexity (Jinushi-Nakao et al., 2007). We generated transgenic fly lines to express human WT CUX2 or variants (p.R34W, p.D337N, p.P454L, p.W958R, and p.E1283K) and analyzed their dendritic arbor morphology in Drosophila larvae (Figure 1D-G and Figure S1B-D). Similar to its Drosophila orthologue (Jinushi-Nakao et al., 2007), ectopic expression of CUX2 WT strongly increased dendritic arbor complexity (branch number and length). However, activities to drive arbor complexity in the variants were significantly decreased, except for p.R34W and p.D337N. RT-qPCR assays in the adult transgenic flies revealed that expression levels of the CUX2 variants, p.P454L and p.W958R, were significantly lower (Figure 1H). All CUX2 constructs were inserted into the same genomic site; therefore, the lower expression levels of transgenes are not likely to be due to position effects but most likely due to variant-dependent degradation. TUNEL of adult fly brains showed that the alleles did not promote apoptotic cell death (Figure S1E). Together, these observations suggest that the CUX2 variants present in patients with epilepsy cause loss-of-function of the protein.

Cux2-deficient mice show increased susceptibility to kainate

Because of the loss-of-function nature of epilepsy-associated CUX2 variants, we next investigated Cux2-KO mice (Iulianella $et\ al.$, 2008). The body weight of 2-month-old mice was comparable among genotypes (Figure S2A). In electrocorticogram analysis, the median of the poly spike and wave discharges frequency was slightly higher in the primary somatosensory cortex forelimb region of Cux2(-/-) than WT mice, however the difference did not reach statistical significance (data not shown). No obvious epileptic behaviors or changes in local field potential recordings in the hippocampus were noted in Cux2(+/-) or Cux2(-/-) mice. Although

patients with TLE often have past histories of febrile seizures (Maher *et al.*, 1995), *Cux2*-KO mice did not show any seizure susceptibility to increased body temperature (data not shown). Seizure susceptibility to pentylenetetrazole (PTZ), a GABA-A receptor antagonist, remained unchanged in *Cux2*-KO mice (Figure S2B-F). Importantly, however, *Cux2*-KO mice had a high susceptibility to kainate, which is commonly used to generate TLE animal models (Lévesque *et al.*, 2013), in frequencies of generalized convulsive seizures (GS) (Figure 2A) and lethality (Figure 2B). The latencies to onset of GS and death were also significantly decreased in *Cux2*(-/-) mice (Figure S2G, H). Seizure severity was also significantly higher in *Cux2*-KO female mice (Figure 2C). These results support the notion that *CUX2* loss-of-function mutations cause TLE.

Cux2-deficient mice show increased cell number in entorhinal cortex and glutamatergic input to hippocampus

Cux2(-/-) mice have been reported to show overgrowth of the neocortical upper layers (Cubelos et al., 2008b). In a Nissl staining, we also found a significantly increased cell number in entorhinal cortex layers II-III, which projects to hippocampal dentate granule cells and CA3 pyramidal cells (Figure 2D). In the slice-patch recordings, we further found that perforant pathevoked excitatory postsynaptic currents (eEPSCs) in the dentate granule cells were significantly higher in Cux2(-/-) mice (Figure 2E), indicating that glutamatergic synaptic transmission from the entorhinal cortex layers II–III onto the hippocampus was significantly facilitated in Cux2-KO mice.

At a glance, hippocampal structures in *Cux2*-KO mice were comparable to 2- and 10-month-old WT mice (Figure 2D, Figure S3). We generated an anti-CUX2 antibody similarly to a previous study (Gingras *et al.*, 2005) and confirmed the presence of CUX2 immunosignals in

WT and absence in Cux2(-/-) mice (Figure S3A, B). CUX2 immunosignals were dense in the neocortical upper layers (II-IV) as previously reported (Zimmer et al., 2004) and also dense in the entorhinal cortex upper layers (II-III) (Figure S3C). In WT hippocampus, we only observed CUX2 immunolabeling signals in inhibitory interneurons, specifically somatostatin (SST)positive, reelin (RLN)-positive and parvalbumin (PV)-positive inhibitory, but not in excitatory neurons (Table S1, Figure S4). We found that there were no significant differences in interneuron cell numbers between genotypes (Figure S5A-D). Timm staining and immunohistochemistry for c-Fos (Figure S5E, F), Doublecortin, phospho-Histone H3, Ki67, NeuN, GFAP, and ZnT-3 (data not shown) did not show differences in the hippocampus between genotypes. There are five subtypes of kainate receptors (KARs), GLUK1–GLUK5, in primates and rodents. We investigated KARs expression in the hippocampus of 2-month-old Cux2-KO female mice. RT-qPCR assays revealed that the expression of GluK1 (formerly named GluR5) was significantly higher in Cux2(-/-) mice (Figure 2F), which is presumably a homeostatic compensatory reaction to epileptic seizures (see Discussion). The baseline frequencies of spontaneous inhibitory postsynaptic currents (sIPSCs) in hippocampal dentate granule cells were significantly higher at 6–7 weeks old, and this difference was suppressed after bath-application of GYKI and AP5, which are AMPA and NMDA receptor antagonists, respectively (Figure 2G). Frequency of sIPSC in dentate PV-positive interneurons was also increased in Cux2-KO mice (Figure S6). These results suggest that, in Cux2(-/-) mice, the function of hippocampal inhibitory neurons remained intact, but the increased excitatory input from the entorhinal cortex to the hippocampus could facilitate firing activities of inhibitory neurons, which itself would also be a compensatory action to epileptic activities in mice. In CA3 pyramidal neurons of Cux2-KO mice, EPSCs were not significantly affected (Figure S6), suggesting that the increased excitatory input

in the upstream dentate granule cells may be neutralized by the increased inhibitory input in those cells.

Taken together, these results suggest that the increase in entorhinal cortex layers II–III cell numbers and the resultant facilitation of glutamatergic synaptic transmission from the entorhinal cortex layers II–III onto hippocampi are causal factors leading to the increased susceptibility to kainate of *Cux2*-KO mice. Increases in GLUK1 and facilitated firing of inhibitory neurons in the mouse hippocampus would rather be compensatory reactions.

CASP variants in TLE patients

CUX1 is a paralog of CUX2, and CASP is an alternatively spliced short isoform of CUX1 harboring a unique C-terminus (Lievens et al., 1997) (Figure 3A, Figure S7). CUX1 and CUX2 proteins have four DNA binding domains (three CUT repeats and one homeodomain), but CASP lacks all of these domains and instead contains a transmembrane domain. We performed targeted sequencing analyses of CUX1 and CASP in the 271 Japanese patients with epilepsy and identified nine nonsynonymous variants (Figure 3A, Table 1, and Supplemental Note). Among those, one variant in CUX1, c.4172C>T (p.T1391I) and three variants in CASP, c.1433C>T (p.A478V), c.1524delG (p.R509fs), and c.1868_1870delTCT (p.F623del), were completely absent or very rare in in-house controls and databases (Table 1). No other truncation variants of the CASP-specific sequence were found in these databases, suggesting that CUX1 and especially CASP variants contribute to epilepsy. Although epilepsies observed in patients with CUX1 and CASP variants were rather heterogeneous, CASP-p.G563S and p.F623del variants appeared in mTLE and lTLE patients, respectively (Table 1). The mTLE patient SIZ-060 showed hippocampal sclerosis (Supplemental Note).

CASP and CUX2 proteins are co-expressed in excitatory neurons of entorhinal cortex upper layer and physiologically bind to each other

Immunohistochemistry with CUX1 antibodies in 2-month-old WT mice revealed CUX1 immunosignals in excitatory neurons at the neocortical upper layers (II-IV), as previously reported (Cubelos *et al.*, 2008b), and those at the entorhinal cortex upper layers (II-III) (Figure 3B), similar to CUX2 (Figure S3C). In contrast in the hippocampus, CUX1 was expressed in SST-positive, RLN-positive, and PV-positive interneurons, but not in excitatory neurons (Figure 3B and Figure S8), similar to CUX2 (Figure S4). Using a CASP-specific antibody recognizing 400–650 a.a., we found that CASP was rather widely expressed in neurons of multiple brain regions, but still dense in the neocortical and entorhinal cortex upper layers, similar to CUX1 and CUX2 (Figure 3C). In the hippocampus, CASP was dense in hilar and stratum-oriens SST-positive cells that expressed CUX2 (Figure 3C, D), and more specifically, within the cytoplasm (Figure S9), which is consistent with CASP expression in the Golgi apparatus (Gillingham *et al.*, 2002).

A protein interaction between CUX1 and CASP has been previously reported (Lievens *et al.*, 1997). Here we newly found that the CASP protein physically interacts with CUX2 (Figure S10). All three tested CUX2 rare variants bound to CASP, and all three tested CASP rare variants bound to CUX2 (Figure S10), suggesting that the variants did not affect protein binding between CASP and CUX2.

Casp-deficient mice also show increased cell number in entorhinal cortex and glutamatergic input to hippocampus

It has been reported that the number of cortical neurons was significantly increased in Cux1(-/-); Cux2(-/-) double-mutant mice, but this increase was no greater than that in the Cux2(-/-) single mutant; therefore, regulation of the upper layer neuronal number was assumed to be a unique function of CUX2 and not redundant with CUX1 activities (Cubelos et al., 2008b). Because of the low survivability of Cux1-KO mice (Luong et al., 2002) and our observation that the TLE variants appeared in the CASP-unique sequence but not in CUXI itself, we decided to investigate Casp-specific KO rather than Cux1-KO mice for analysis. We generated a Caspspecific KO mouse by targeting exon 17 at the unique C-terminus (Figure S7C). Casp(+/-) and Casp(-/-) mutant mice were born at a Mendelian ratio, grew normally, and were fertile. RTqPCR analyses revealed that the Casp mRNA became half and diminished in Casp(+/-) and Casp(-/-) mice, respectively, whereas Cux1 and Cux2 mRNA levels remained unchanged (Figure 3E). CASP immunosignals well disappeared in Casp(-/-) mice (Figure S9A), confirming the specificity of the CASP antibody. At a glance, there were no abnormal localizations and intensities of CUX1 and CUX2 proteins in Casp-KO mice (Figure S9B, C). The median body weight was comparable among genotypes at 2 months of age (Figure S9D). RT-qPCR assays of KARs mRNA in the hippocampi of 2-month-old *Casp*-KO mice did not show significant change in KARs expression levels (Figure S9E).

In a Nissl staining of 2-month-old *Casp*-KO mice, no increase of neuron number was observed in the entorhinal cortex (Figure S9F). However, immunohistochemical staining using the anti-CUX1 antibody as a marker of neurons at upper layers of the neocortical (II–IV) and the entorhinal cortex (II–III) showed a tendency of increase in both the neocortex and entorhinal cortex ($P = 7.57 \times 10^{-2}$ and $P = 4.50 \times 10^{-1}$, respectively) (Figure 3F). Furthermore, *Casp*-KO mice also showed high susceptibility to kainate (Figure 3G, H, Figure S9G-I). After

intraperitoneal application of kainate, a larger number of *Casp*-KO mice showed GS and lethality (Figure 3G, H). Onset latencies of GS and death were significantly decreased in *Casp*-KO mice (Figure S9H, I). Seizure severity in *Casp*(-/-) mice was also significantly higher (Figure S9G). Kainate susceptibility was higher in male *Casp*-KO mice, contrary to *Cux2*-KO mice in which the susceptibility is higher in female. Notably again, perforant path-evoked EPSCs (eEPSCs) in the dentate granule cells were significantly higher in *Casp*-KO mice (Figure 3I), which is similar to *Cux2*-KO mice (Figure 2E).

All of these observations propose that facilitation of glutamatergic synaptic transmission from the entorhinal cortex onto hippocampal dentate granule cells is a common mechanism for TLE caused by *CUX2* and *CASP* variants.

Discussion

In this study, we performed targeted sequencing analyses of CUX2, CUX1 and CASP on 271 Japanese patients with a variety of epilepsies, and found that CUX2 missense variants predominantly appear in TLE patients, in that eight of 68 TLE patients (12%) had CUX2 variants. Most of the CUX2 variants showed functional impairments. CASP variants also appeared in two TLE patients. Most of these patients showed hippocampal sclerosis, suggesting that CUX family variants may contribute to TLE, especially drug-resistant mesial TLE. Our recent GWAS analysis of Japanese patients with variable epilepsies identified a region with genome-wide significance at chromosome 12q24 which harbors CUX2 (Suzuki et al., 2021). Although a recurrent de novo CUX2 variant p.E590K has been described in patients with EEs (Chatron et al., 2018; Barington et al., 2018), a previous whole exome sequencing study for Japanese patients with EEs (Takata et al., 2019) did not find the CUX2 pathogenic variant, and therefore the CUX2 recurrent variant in EEs may not be enough to explain the genome-wide association with epilepsy at the 12q24 region in Japanese population. In our GWAS study (Suzuki et al., 2021), sub-analyses for subtypes of epilepsies further revealed that a polymorphic marker at the 12q24 epilepsy-associated region showed genome-wide significant association with symptomatic epilepsy. TLE is the major entity for symptomatic epilepsy, and therefore the CUX2 variants in patients with TLE would contribute to the association with epilepsy at 12q24 in Japanese population.

Human cell culture and fly dendritic arborization analyses revealed loss-of-function properties of the CUX2 variants appeared in TLE patients. *CASP* also showed variants in epilepsy patients including TLE at the unique C-terminus and we further found that the CASP physically binds to CUX2. Although all tested CASP variants did not affect the binding activity

to CUX2, the CASP protein has been reported to play a role in intra-Golgi retrograde transport (Malsam *et al.*, 2005) and therefore the variants in CASP may still affect the subcellular transport or protein modification of CUX2.

Cux2- and Casp-KO mice did not show spontaneous seizures but showed significantly elevated seizure susceptibility to kainate, an agent which has been used to establish TLE animal models (Lévesque et al., 2013). We previously reported a nonsense mutation of the KCND2 gene encoding a voltage-gated potassium channel Kv4.2 in a patient with TLE (Singh et al., 2006). Similar to Cux2- and Casp-KO mice, Kcnd2-KO mice did not show spontaneous epileptic seizures but showed increased susceptibility to kainate in seizure and mortality (Barnwell et al., 2009). These observations suggest that increased seizure susceptibility to kainate correlates with genetic susceptibility to TLE and that other additional modifying, genetic, or environmental factors influence full expression of TLE phenotypes.

Cux2-KO mice showed a significant and Casp-KO mice showed a tendency of, increases in entorhinal cortex layer II–III excitatory neuron cell number. Although Casp-KO mice did not show any significant changes in Cux2 mRNA expression levels and histological or cytological distributions of CUX2 protein, co-expression of CASP and CUX2 proteins in neurons including entorhinal cortex projection neurons, and the physiological interaction between CASP and CUX2 proteins still suggest that CASP deficiency may impair CUX2 function through an as-yet unknown mechanism, consequently leading to increased entorhinal cortex excitatory neuron cell number in Casp-KO mice. Furthermore, both Cux2- and Casp-KO mice revealed significant increases in perforant path-evoked EPSCs in dentate granule cells. These results suggest that facilitation of glutamatergic synaptic transmission from the entorhinal cortex onto hippocampal dentate granule cells is a causal basis for the significant increase in seizure susceptibilities to

kainate in Cux2- and Casp-KO mice. In contrast, the observed changes in the hippocampus of Cux2-KO mice are assumed to be homeostatic compensatory reactions to the epileptic causal changes. In the hippocampus of WT mice, CUX2 immunolabelling was only observed in inhibitory interneurons such as SST-positive, RLN-positive, and PV-positive inhibitory, but not in excitatory neurons. In the Cux2-KO mice, although no changes were observed in interneuron cell numbers, a significant increase in sIPSC frequency was observed in dentate granule cells similar to patients with TLE (Li et al., 2010), which would presumably be a compensatory reaction to the increased excitatory input from the entorhinal cortex to the hippocampus. The increased GLUK1 expression in the hippocampus of Cux2-KO mice may also be suppressive for epileptic seizures because GluK1 is expressed in inhibitory neurons (Bureau et al., 1999), and GLUK1 expression has been assumed to be protective at least for kainate-induced epilepsy (Fritsch et al., 2014). Taken together, the changes within the hippocampus of Cux2-KO mice may be homeostatic compensatory responses rather than causal actions to epileptic seizures, and these changes in the hippocampus themselves also support the occurrence of epileptic causal changes in these mice.

In summary, our results of mutation analyses of *CUX* family genes in patients with epilepsies including TLE and the functional and mouse model analyses suggest that *CUX* family gene deficiency is one of the bases for TLE and that increase of cell number in the entorhinal cortex projection neurons and resultant increase of glutamatergic synaptic transmission to hippocampus is a possible pathological mechanism for TLE.

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Disclosure of Conflicts of Interests

None of the authors has any conflict of interest to disclose.

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Figure Legends

Figure 1. Loss-of-function effects of TLE variants in *CUX2*. (A) CUX2 protein structure (NP_056082) with variants appeared in patients with epilepsy. (B) Abnormal subcellular localization of CUX2 variant proteins. CUX2-WT protein (arrows) was limited to, but well distributed within, nuclei stained with DAPI (cyan), whereas variants showed abnormal aggregates in nuclei (W958R) or leaked-out to the cytoplasm (E1283K) (arrowheads). Scale bars = 20 μm. (C) Ratio of abnormally localized CUX2 proteins (> 200 cells counted). n = WT: 545, R34W: 282, D337N: 363, P454L: 239, W958R: 565, and E1283K: 323 cells. (D-H) CUX2 WT accelerated arborization of fly neurons and TLE variants lowered its activity and expression. Representative images of neurons without CUX2 (D), WT control (E), and W958R (F). Scale bars = 50 μm. (G) Shortened dendrite length in transgenic fly with mutants (n = 11–25 neurons per genotypes) and (H) lowered expression of mutants (n = 6). One-way ANOVA Tukey's multiple comparison test (C, G, H). *P < 0.05, **P < 0.01, ***P < 0.001.

Figure 2. Increased kainate susceptibility, entorhinal cortical cell number, and excitatory input to hippocampal granule cells in Cux2-KO mice. (A-C) Seizure-related events in mice after intraperitoneal injection of kainate (KA). Ratio of animals exhibiting generalized convulsive seizure (GS) (A), mortality rate (B), and seizure severity scores (C) was significantly higher in Cux2(-/-) female and combined gender mice. (D) Number of entorhinal cortex layer II-III excitatory neurons was significantly increased in Cux2(-/-) mice (2-month-old). Scale bar = $100 \mu m$. (E) Slice-patch analyses showed that perforant path-evoked EPSCs in dentate granule cells were significantly increased in Cux2(-/-) female. (F) RT-qPCR analyses revealed that GluK1 mRNA was significantly increased in Cux2(-/-) mice. (G) Basal frequency of sIPSC in

dentate granule cells of *Cux2*(-/-) female was significantly increased, and it was suppressed with subsequent applications of antagonists for AMPA receptor (GYKI) and NMDA receptor (AP5). Kainate (KA) increased the sIPSC frequency, which was then suppressed by the GABA-A receptor antagonist picrtoxin. DG; dentate gyrus, Ent; entorhinal area. Yates' correction after Pearson's Chi-square (A, B), one-way ANOVA Tukey's test (C, F, G), one-way ANOVA (D), or two-way ANOVA Tukey's test (E). n: mouse numbers. *P < 0.05, **P < 0.01, ***P < 0.001.

Figure 3. CASP variants in epileptic patients, CASP distribution, and increases in kainate susceptibility and excitatory input to hippocampal granule cells in Casp-KO mice. (A) Locations of CUX1 and CASP variants in patients with epilepsy (see Table 1). Dashed lines define the common region. (B) CUX1 immunosignals (brown) in neocortical and entorhinal cortex upper layer excitatory neurons and hippocampal interneurons. (C) CASP (brown) expressed more widely in neurons, and intensely expressed in neocortical and entorhinal cortex upper layer excitatory neurons. (D) In hippocampus, CASP (brown) was dense in SST-positive (blue) interneurons at hilus and stratum oriens (arrows). d2-d5; magnified images outlined in d1. Scale bars = $100 \mu m$ (B, C and d1), $20 \mu m$ (d2-d5), so; stratum oriens, sp; stratum pyramidale, sg; stratum granulosum, h; hilus. (E) RT-qPCR analyses revealed that Casp mRNA was decreased, while Cux1 and Cux2 mRNAs remained unchanged, in Casp-KO mice. (F) Thickness of the CUX1-positive neocortical layer (left), density of CUX1-positive cell in neocortex (middle), and CUX1-positive cell density in entorhinal cortex (right). CUX1-positive cell density tended increase at neocortex and entorhinal cortex in Casp(-/-) mice (2-month-old) but not statistically significant. (G, H) Casp-KO mice showed significantly higher susceptibility to kainate in seizure rate (G), mortality (H). (I) Perforant path-evoked EPSCs in dentate granule

cells were significantly increased in Casp(-/-) male (6–7-week-old). One-way ANOVA Tukey's test (E), Yates' correction after Pearson's Chi-square (G, H), or two-way ANOVA Tukey's test (I). n: mouse numbers. *P < 0.05, **P < 0.01, ***P < 0.001.

Supplementary Figure Legends

Figure S1. CUX2 mutants show abnormal subcellular localizations in human. cultured cells and decreased branching effects but no apoptosis in fly. (A) In HeLa.S3 cells, CUX2 mutant proteins (R34W, D337N, and P454L) show aggregates or abnormal leakage-out into cytoplasm (arrowheads) and some WT-like distribution (arrows). Nuclei were stained with DAPI (cyan). (B-D) Mutations lowered the neurite arborization activity of CUX2 in fly neurons. Representative images of CUX2-R34W (B) and CUX2-P454L (C). Terminal point numbers were significant decreased in CUX2-P454L, CUX2-W958R and E1283K (n=13 ~ 25) (D). Results of one-way ANOVA followed by Tukey's test. (E) TUNEL assay revealed no apoptosis in adult transgenic flies of CUX2-WT and mutants. Scale bars = 20 μ m (A) or 50 μ m (B, C, and E). * P < 0.05, ** P < 0.01, *** P < 0.001.

Figure S2. Cux2-deficient mice show seizure susceptibility to kainate but not to PTZ. (A) Body weight was similar among genotypes of 2-month-old mice. (B-F) Unchanged seizure susceptibility of Cux2-KO mice to PTZ. There are no significant differences in latency to generalized convulsive seizure (GS) (B), latency to death (C), percentage of animals exhibiting GS (D), mortality rate (E), and seizure severity score (F) among genotypes. (G, H) Cux2-KO mice show increased seizure susceptibility to kainate. Latency to onset of GS (G) and that to death (H) were significantly decreased in Cux2(-/-) female and combined gender mice. One-way ANOVA (A-C, F-H) or Pearson's Chi-square test (3×2 contingency table) (D, E). mean

(horizontal bars) \pm s.e.m. (B, C, G, H). n or numbers in round brackets =mouse numbers. * P < 0.05, ** P < 0.01.

Figure S3. CUX2 antibody specifically recognizes CUX2 protein. (A) Sagittal brain sections from 10-month-old adult WT mice were stained with an antibody to CUX2. Immunosignals (arrows) were observed widely in the brain including in neurons at the hippocampus and cerebral cortex. A2-A5: magnified images outlined in A1. (B) CUX2 immunosignals were not observed in *Cux2*(-/-) mouse. B2-B5: magnified images outlined in B1. (C) In 2-month-old wild-type mouse, CUX2 (brown) is densely expressed in excitatory neurons at neocortical (II–IV) and entorhinal cortex (II–III) upper layers, but in hippocampal observed in inhibitory but not excitatory neurons. (D) In hippocampus at P15, CUX2 (brown) is expressed in interneurons but not in excitatory neurons. Scalebars=500 μm (A1 and B1), 100 μm (C, D) or 20 μm (A2-A5 and B2-B5). CTX; cerebral cortex, DG; dentate gyrus, so; stratum oriens, h; hilus.

Figure S4. CUX2 is expressed in hippocampal SST-positive, RLN-positive or PV-positive inhibitory neurons. Tissue sections from P15 (A-C) or 2-month-old (E-G) WT mouse brains were stained with antibodies to CUX2 (brown) and somatostatin (SST, blue) (A, E), reelin (RLN, blue) (B, F) or parvalbumin (PV, blue) (C, G). CUX2 expression was observed in SST-positive, PV-positive and RLN-positive (arrow) interneurons at both stages. Some of the intense CUX2-positive cells were SST-negative, RLN-negative or PV-negative (white arrow head), and some of the intense SST-positive or PV-positive cells were CUX2-negative (black arrow head). CUX2-positive / PV-positive cell number increased at 2-months compared to P15. A2-A4, B2-B4, C2-C4, E2-E4, F2-F4, G2-G4, A5 and B5: magnified images outlined in A1, B1, C1, E1, F1, G1, A4 and B4, respectively. (D, H) Tissue sections from at P15 (D) or 2-month-old (H) WT mouse brain were stained with antibodies to SST (green) or RLN (magenta) and DAPI (cyan). The SST

expression was observed in RLN-positive interneurons (arrows) at the hilus of hippocampus. SST/RLN-double positive cells were more frequent in P15 (D) than 2-month-old sections (H). RLN signals became weaker at 2-months (H) compared to P15 (D). Some of intense SST-positive cells were RLN-negative (arrow head), and some of intense RLN-positive cells were SST-negative (asterisk). Scale bar=100 µm (A1, B1, C1, E1, F1, and G1) or 20 µm (A2-A4, B2-B4, C2-C4, D, E2-E4, F2-F4, G2-G4 and H). h; hilus.

Figure S5. Unchanged cell numbers of hippocampal inhibitory neurons, mossy fibers and cFos expression in Cux2-deficient mice. (A) Hippocampus was separated in 7 regions and immunoreactive cells were counted. Cell densities were determined by average number of cells in 4 sections / area. (B-D) There were no differences in the number of SST-positive (B), RLN-positive (C) or PV-positive (D) neurons between genotypes. Statistical analyses were performed using one-way ANOVA (p > 0.05) (n = WT: 6, Cux2(+/-): 6, Cux2(-/-): 6). so; stratum oriens, sp; stratum pyramidale, sr; stratum radiatum, lm; lacunosum-moleculare, sm; stratum moleculare, sg; stratum granulosum, h; hilus, w; whole hippocampus. (E, F) Timm staining (E) and c-Fos immunohistochemistry (F) did not show differences in Cux2-KO mice. Scale bar = $100 \mu m$ (A and E).

Figure S6. IPSCs in PV-positive cells at dentate gyrus were increased, but no differences in EPSCs in CA3 pyramidal cells of *Cux2*-deficient mice. (A-D) In PV-positive cells at dentate gyrus, IPSCs were measured at 6~7-week-old. Amplitude histograms of sIPSC (A) and mIPSC (C) from WT mice and *Cux2*(-/-) mice. Cumulative probability plots and average values (inset) for sIPSC (B) and mIPSC (D) show significant differences in inter-event intervals of sIPSC and mIPSC and amplitude of mIPSC populations derived from *Cux2*(-/-) and significantly higher average of frequency of sIPSC, but unchanged averages of amplitude of sIPSC and both

frequency and amplitude of mIPSC in Cux2(-/-). (**E-H**) In pyramidal neurons at CA3 region, EPSCs were measured at $6\sim7$ -week-old. Amplitude histograms of sEPSC (E) and mEPSC (G) from WT, Cux2(+/-) and Cux2(-/-) mice. Cumulative probability plots and average values (inset) for sEPSC (F) and mEPSC (H) in WT, Cux2(+/-) and Cux2(-/-) show no differences in EPSCs among genotypes. (I) No significant difference on MF-evoked EPSCs in pyramidal neurons at CA3 region was observed across genotypes at $6\sim7$ -week-old. Statistical analyses were performed using one-way ANOVA followed by Tukey–Kramer Multiple Comparison Test (B, D, F, H, I) or Kolmogorov-Smirnov (K-S) Test (B, D). * P < 0.05, *** P < 0.001.

Figure S7. Genome structures of human and mouse CUX1 and CASP in WT and Casp-specific knock-out mice. Exons 1~14 are commonly found in human CUX1 and CASP in human (A) and in the corresponding genes in mice (B). In Casp-KO mice, a mutation (c.1514^1515insTT, p.S506fs) was inserted in Casp-specific exon 17 (C). The diagrams were reconstructed from that of UCSC Genome Browser. Vertical lines indicate exons.

Figure S8. CUX1 is expressed in hippocampal SST-positive, RLN-positive, or PV-positive interneurons. CUX1 (brown) is expressed in SST-positive (A), RLN-positive (B) and PV-positive (C) interneurons (arrows). Some of intense CUX1-positive cells were SST-negative, RLN-negative or PV-negative (white arrowheads), and some of SST-positive, RLN-positive or PV-positive cells were CUX1-negative (black arrowhead and not shown). Most of CUX1-positive cells were PV-negative (white arrowhead) in hilus (C). A2-A4, B2-B4, C2-C4: magnified images outlined in A1, B1, C1, respectively. Scale bars = 100 μm (A1, B1 and C1) or

20 μm (A2-A4, B2-B4 and C2-C4). sp; stratum pyramidale, so; stratum oriens, sg; stratum granulosum, h; hilus.

Figure S9. CASP, CUX1 and CUX2 expression and increased seizure susceptibility to kainate with a tendency of increase in excitatory cell number in entorhinal cortex of Casp**deficient mice.** (A) CASP immunosignals were observed in both excitatory and inhibitory neurons in the hippocampus, though the signals in inhibitory neurons were especially intense such as those at hilus (A3) and stratum oriens of CA3 (A5), while signals in excitatory neurons are tiny such as those at stratum granulosum (A4) or pyramidal of CA3 (A6). CASP signals well disappeared in Casp(-/-) mice (A7, A8). Nuclei were stained with hematoxylin (blue). (B, C) Unaltered expression and subcellular localization of CUX1 and CUX2 in cerebral cortex (B) and hilus of hippocampus (C) of Casp-KO mice. (D) Body weights were similar among Casp-KO mice and WT littermates at 2-months. (E) qPCR experiments showed that mRNA expression levels of kainate receptor subunits were not altered in Casp-KO mice at 2-month-old. (F) In a Nissl staining, number of entorhinal cortex layer II-III neurons was comparable between WT and Casp(-/-) mice (2-month-old). (G) Seizure severity scores were significantly higher in Casp(-/-) male and combined gender mice. (H) Latencies to onset of generalized seizures (GS) were significantly decreased in male and combined gender of Casp(-/-) mice. (I) Latencies until death were also significantly decreased in combined gender of Casp(-/-) mice. Mice without GS or death within 3,600 sec were plotted at "no GS" or "no death", respectively. Circles represent individual mice. One-way ANOVA followed by Tukey-Kramer Multiple Comparison Test (D,

E, G-I), or one-way ANOVA test (F). mean (horizontal bars) \pm s.e.m. (H, I). n=mouse number. Scale bars= 50 μ m (A1, A2, A7, A8, B and C) or 10 μ m (A3-A6). * P < 0.05, ** P < 0.01.

Figure S10. CASP interacts with CUX2. mRFP-tagged CUX2 was co-immunoprecipitated with FLAG-tagged CASP. Mutations did not affect the binding. Endophilin: negative control.

Table 1: CUX2, CUX1, and CASP gene nonsynonymous variants in patients with epilepsy.

Patient ID	Gene	Nucleotide changes	Amino acid substitutions	SNP ID	Onset age (year)	Sex	Diagnosis	Variant allele count in						Mutation	D 1 D1 -	nn oven a v	CHET	
								case	JP	J-HGVD	EVS	1kGP	ExAC	gnomAD	Taster	PolyPhen-2	PROVEAN	SIFT
SIZ-220	CUX2	c.100C>T	p.R34W	rs199531850	10	M	mTLE	1 / 542	0 / 622	1 / 1,900	1 / 11,778	1 / 5,008	21 / 119,874	62 / 273,306	++	-	++	++
SIZ-016		c.1009G>A	p.D337N	rs201601231	16	M	mTLE	1 / 542	5 / 622	4 / 2,192	1 / 12,344	2 / 5,008	18 / 117,772	34 / 279,160	++	++	-	-
SIZ-296		c.1361C>T	p.P454L	rs768144991	2	M	Doose syndrome	1 / 542	0 / 622	5 / 2,184	NR	NR	1 / 24,448	8 / 167,304	++	+	++	-
SIZ-014		c.2872T>C	p.W958R	NA	3	M	mTLE	1 / 542	0 / 622	NR	NR	NR	NR	NR	++	++	++	++
SIZ-004		c.3847G>A	p.E1283K	rs61745424	11	F	mTLE	5/542	2 / 622	9 / 2,126	227 / 12,676	143 / 5,008	3,893 / 120,384	8,188 / 280,456	-	-	-	
SIZ-022					8	F	mTLE											
SIZ-073					19	M	mTLE											-
SIZ-079					13	M	ITLE											
SIZ-190					16	F	mTLE											
SIZ-784	CUXI	c.3161C>T	p.S1054L	rs146486358	16	M	JME suspected	2 / 542	NT	9 / 1,912	NR	13 / 5,008	101 / 120,970	231 / 282,774	++	++	-	-
SIZ-891					NA	M	GEFS											
SIZ-575		c.3281C>T	p.A1094V	rs184337744	3	F	FLE	1 / 542	NT	NR	1 / 13,006	1 / 5,008	38 / 120,936	38 / 250,814	-	+	-	-
SIZ-669		c.3815G>A	p.R1272Q	NA	0	F	SGE	1 / 542	NT	2 / 2,152	NR	NR	NR	NR	++	++	-	++
SIZ-456		c.4172C>T	p.T1391I	NA	2	F	CAE	1 / 542	0 / 620	NR	NR	NR	NR	NR	-	-	-	-
	CASP	c.1390G>A	p.A464T	rs803064				247 / 542	NT	991 / 2,210	NR	2,800 / 5,008	68,923 / 121,236	159,104 / 282,310	-	-	-	-
SIZ-127		c.1433C>T	p.A478V	NA	14	F	JME	1 / 542	0 / 622	NR	NR	NR	NR	NR	++	+	-	-
SIZ-063		c.1524delG	p.R509fs	rs782400087	6	F	CAE	1 / 542	0 / 622	NR	NR	NR	1 / 121,412	1 / 249,562	++	NA	NA	NA
SIZ-068		c.1687G>A	p.G563S	rs187131238	12	F	JME	2 / 542	1 / 622	3 / 2,164	NR	2 / 5,008	4 / 120,438	7 / 247,922	-	-	-	-
SIZ-060					18	M	mTLE											
SIZ-638		c.1868_1870delTCT	p.F623del	NA	15	M	ITLE	1 /542	0 / 622	NR	NR	NR	NR	NR	++	NA	++	NA

mTLE; mesial temporal lobe epilepsy, ITLE; lateral temporal lobe epilepsy, JME; juvenile myoclonic epilepsy, CAE; childhood absence epilepsy, SGE; symptomatic generalized epilepsy, FLE; frontal lobe epilepsy, GEFS; generalized epilepsy with febrile seizure plus, M; Male, F; Female. JP; in-house Japanese control individuals, J-HGVD; Japanese Human Genetic Variation Database, EVS; Exome Variant Server NHLBI GO Exome Sequencing Project, 1kGP; The 1000 Genomes Project, ExAC; Exome Aggregation Consortium, gnomAD; Genome Aggregation Database. ++; Disease causing, Probably damaging, Deleterious, or Damaging, +; Possibly damaging, Genome Aggregation Database, The CUX2 reference sequence (NM_015267) has an error at c.4414, and the correct nucleotide is C. CUX2 nucleotide change c.4414G>C (p.V1472L) (rs6490073 in dbSNP, NCBI) was observed in all sequences in databases and in our subjects, suggesting that the CUX2 reference sequence (NM_015267) has an error at this position (the correct nucleotide is c.4414C).

Figure 1

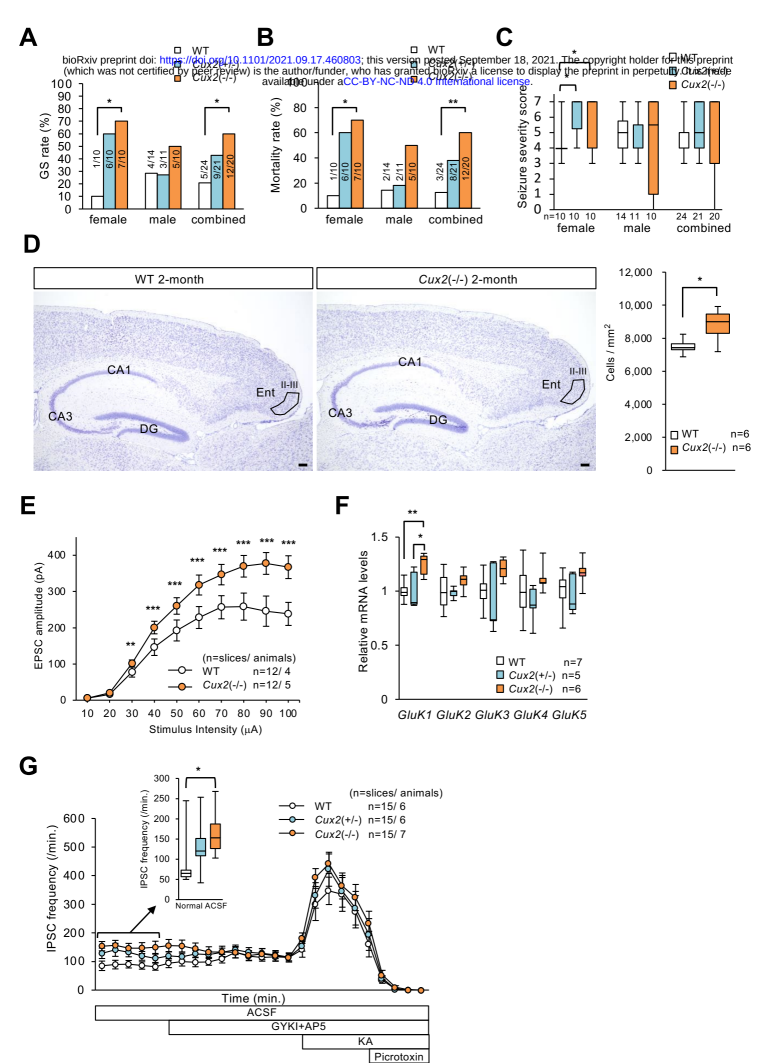


Figure 2

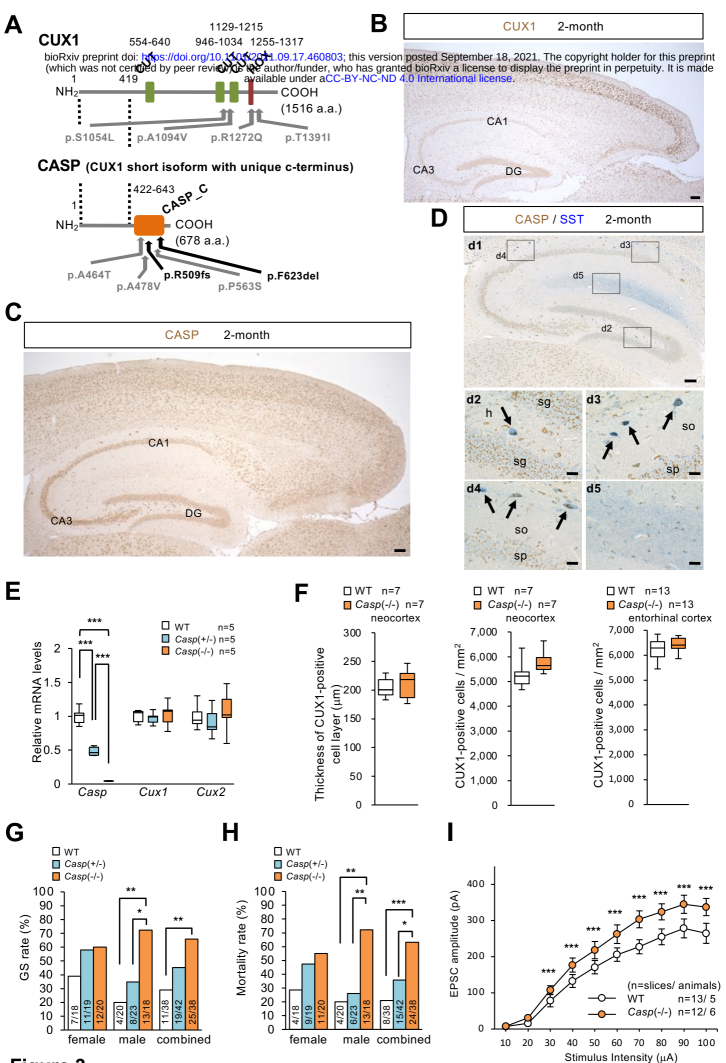


Figure 3