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Developmental and epileptic encephalopathies (DEE) are characterized by pharmacoresistant seizures with concomitant intellectual disability. Epilepsy of infancy with migrating focal seizures (EIMFS) is one of the most severe of these syndromes. De novo variants in ion channels, including gain-of-function variants in *KCNT1*, have been found to play a major role in the etiology of EIMFS. Here, we test a potential precision therapeutic approach in *KCNT1*-associated DEE using a gene silencing antisense oligonucleotide (ASO) approach. We generated a mouse model carrying the KCNT1 p.P924L pathogenic variant; only the homozygous animals presented with the frequent, debilitating seizures and developmental compromise that are seen in patients. After a single intracerebroventricular bolus injection of a *Kcnt1* gapmer ASO in symptomatic mice at postnatal day 40, seizure frequency was significantly reduced, behavioral abnormalities improved, and overall survival was extended compared to mice treated with a control ASO (non-hybridizing sequence). ASO administration at neonatal age was also well-tolerated and effective in controlling seizures and extending the lifespan of treated animals. The data presented here provide proof of concept for ASO-based gene silencing as a promising therapeutic approach in *KCNT1*-associated epilepsies.

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Antisense oligonucleotide therapy for KCNT1 encephalopathy

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ABSTRACT

Developmental and epileptic encephalopathies (DEE) are characterized by pharmacoresistant seizures with concomitant intellectual disability. Epilepsy of infancy with migrating focal seizures (EIMFS) is one of the most severe of these syndromes. *De novo* variants in ion channels, including gain-of-function variants in *KCNT1*, have been found to play a major role in the etiology of EIMFS. Here, we test a potential precision therapeutic approach in *KCNT1*-associated DEE using a gene silencing antisense oligonucleotide (ASO) approach. We generated a mouse model carrying the KCNT1 p.P924L pathogenic variant; only the homozygous animals presented with the frequent, debilitating seizures and developmental compromise that are seen in patients. After a single intracerebroventricular bolus injection of a *Kcnt1* gapmer ASO in symptomatic mice at postnatal day 40, seizure frequency was significantly reduced, behavioral abnormalities improved, and overall survival was extended compared to mice treated with a control ASO (non-hybridizing sequence). ASO administration at neonatal age was also well-tolerated and effective in controlling seizures and extending the lifespan of treated animals. The data presented here provide proof of concept for ASO-based gene silencing as a promising therapeutic approach in *KCNT1*-associated epilepsies.
INTRODUCTION

Epilepsy of infancy with migrating focal seizures (EIMFS) (previously known as malignant migrating partial seizures of infancy) is one of the most severe developmental and epileptic encephalopathy (DEE) syndromes (1). Pharmacoresistant, nearly continuous multifocal seizures begin to occur during the first six months of life and are accompanied by a marked developmental regression or stagnation (1, 2). In addition, major axial hypotonia, as well as pyramidal and extrapyramidal signs become more apparent with the progressive development of athetotic movements and other movement disorders (2–4). Many of these patients also display microcephaly and strabismus (2–4). The prognosis of this condition is very poor with most patients being non-verbal and non-ambulatory (2, 4, 5). Importantly, this syndrome is associated with a high mortality rate (ranging from 17% to 33%) (1, 2, 5), that can occur as a consequence of prolonged status epilepticus and respiratory failure (2).

Pathogenic gene variants have been identified in KCNT1 and account for up to 50% of the etiology of EIMFS (1, 3, 6–10). KCNT1 encodes the sodium activated potassium channel protein KNa1.1 which mediates an outward rectifying K⁺ current related to the slow hyperpolarization that follows repetitive action potential firing (11–16). Functional studies have shown that KCNT1 pathogenic variants associated with epilepsy result in an overall gain of function effect on the channel activity, increasing the current up to 22-fold compared to the wild-type channel (6, 11, 17). For a subset of these pathogenic variants, changes in voltage dependence or Na⁺ sensitivity contribute to the increase in current (18, 19). Gain of function variants in KCNT1 have also been found in patients with other epileptic syndromes including Autosomal Dominant Nocturnal Frontal Lobe Epilepsy, Ohtahara syndrome, and Lennox-Gastaut syndrome (3, 5, 7, 8, 20, 21). The antiarrhythmic drug quinidine has been shown to reduce the pathogenic currents produced by some of the mutant
KCNT1 channels in vitro (11, 17, 22–25) but has had a limited clinical translation due to unwanted side effects (QTc prolongation and increased risk for arrhythmia) thereby resulting in a limited therapeutic window (25–37). To date, there is no effective therapy to treat KCNT1 associated epilepsies.

RNA-targeted therapies have recently received significant attention for the recurrent examples of preclinical and clinical success, as well as fast-tracked development (38, 39). Some of these approaches are proving to be disease-modifying in the treatment of progressive neurological conditions (40, 41). Of the RNA-targeted therapies, which include antisense oligonucleotides (ASOs), short interfering RNA (siRNA), antagonirs, microRNA mimetics, and DNAzymes, ASOs have provided the most impactful clinical benefit (38, 41, 42). ASOs have emerged as a therapeutic option for orphan pediatric neurogenetic conditions including spinal muscular atrophy (43) and Duchenne’s muscular dystrophy (44). ASOs are synthetic, single-stranded nucleic acids typically of 10 to 30 nucleotides in length that bind to a specific complementary mRNA target sequence and modulate the expression of a specific gene at the RNA level (42). This control is achieved by different molecular mechanisms, from the steric block of ribosomal activity to regulation of RNA splicing (39, 42). ASOs that harness endogenous RNase H1 mechanisms are commonly used to specifically reduce the expression of mRNAs. These ASOs are referred to as gapmer ASOs because they contain central block of deoxynucleotide monomers needed to induce the cleavage of target mRNA (39, 42). The major advantage of ASOs as a therapeutic approach is that the interaction with the target is considerably more specific than traditional small-molecule based therapeutics (42).

To investigate the therapeutic potential of reducing KCNT1 expression we evaluated the efficacy of a Kcnt1 gapmer ASO in a homozygous mouse model of a KCNT1-DEE pathogenic variant.
Mice homozygous for the *Kcnt1* variant p.P905L (L/L) display spontaneous seizures, abundant interictal activity in the electrocorticogram (ECoG), behavioral abnormalities, and early death. After a single intracerebroventricular (i.c.v.) bolus injection of *Kcnt1* gapmer ASO at postnatal day 40, L/L mice showed a marked knockdown of *Kcnt1* mRNA, resulting in an almost complete abolition of seizures, prolonged survival, and improved performance in behavioral tests. A dose of gapmer ASO that produced >90% *Kcnt1* mRNA knockdown to mimic exaggerated pharmacology in wild-type mice was well tolerated suggesting minimal on-target liability. The preclinical evidence presented here provides proof of concept for ASO based gene silencing as a therapeutic approach in *KCNT1* gain of function epilepsies.

**RESULTS**

The *Kcnt1* L/L model of *KCNT1* encephalopathy

*Kcnt1* p.P905L heterozygous mice (L/) showed no evident epileptic or behavioral phenotype (Figures 1 and 2) and no increased susceptibility to chemical or thermally induced seizures (Supplemental Figure 1, A-D). In contrast, homozygous mice (L/L) were smaller in size (Figure 1 A and B) and had a markedly reduced lifespan compared to their littermates (median survival of 43 days) (Figure 1C). In addition, the yield of L/L mice, born from L/+ breeders, was not consistent with the predicted Mendelian ratio of 1:2:1 (+/+: L/+: L/L) with only 8.8% of pups born found to be L/L by age P8-12 (age at which mice were sampled for genotyping) (Supplemental Figure 1E). Spontaneous tonic-clonic seizures were observed as early as P18. Video monitoring and handling showed diverse seizure phenotypes, with milder seizures lasting approximately 1 to 2 minutes and involving clonic movements of the forelimbs, neck and head while in a seated posture (Video 1), which were consistent with a Racine seizure intensity stage of 3 to 4 (45). More severe seizures including generalized tonic-clonic episodes with Straub tail, wild running and
jumping, loss of postural tone and tonic hindlimb extension (Racine 5 and 6). Isolated atonic seizures (pure postural tone loss) were also observed. Milder seizures, in general, were not followed by an evident postictal state, and the mice quickly recovered (Video 2-4). Longer and generalized seizures were followed by a short period of complete immobility and rapid breathing with later episodes of reduced mobility lasting up to 10 minutes. Status epilepticus (a seizure lasting > 5 minutes) was also observed and in most cases resulted in death. The frequency of tonic-clonic seizures was variable between mice (Figure 1F). ECoG recordings revealed the presence of frequent high amplitude interictal acute spikes in the L/L mice, (Figure 1D) this signal was absent in L/+ and +/+ (Figure 1E) and did not correlate with changes in behavior. Convulsive seizures were associated with ictal ECoG signals characterized by clusters of high-amplitude sharp wave activity followed by electrical suppression at the end of the seizure (Figure 1D, Video 5).

No differences in seizure frequency or survival were observed based on sex (data not shown).

Behavioral abnormalities were also found in the L/L mice. Nest building ability, an identified measure of the general wellbeing in mice (46, 47), has shown to be sensitive to brain lesions, the application of pharmacological agents, and the effects of genetic variants (48). In mouse models, instinctual nest-building performance is used to simultaneously assess general social behavior, cognitive, and motor performance (49). At P40, the nesting behavior in the L/L mice was markedly impaired compared to their +/+ and L/+ littermates (Figure 2, A and B).

Gross motor malfunction and exploratory behaviors were tested on the locomotor cell test and revealed a tendency towards hyperactivity, especially within the first 20 minutes of testing. The light/dark box test showed L/L mice are prone to anxiety-like traits based on the limited time spent on the light chamber (Figure 2D). The elevated plus maze (EPM) was used to further explore anxiety-like traits. Interestingly, the L/L mice showed a marked preference for the open arms.
compared to their littermates (Figure 2E), suggesting a reduced fear for open and elevated spaces. Lastly, alterations in social behavior and cognition were tested in L/L mice using the three-chamber social interaction test and Y-maze test, respectively, but showed no significant difference compared to +/+ mice (Supplemental Figure 1, F-G).

**ASO-mediated Kcnt1 reduction rescues the seizure phenotype in L/L mice and improves overall behavior and survival.**

To determine if the levels of Kcnt1 mRNA could be reduced in a dose-dependent fashion we tested a mouse-specific Kcnt1 ASO in adult +/+ mice. At P40, Kcnt1 ASO was delivered by i.c.v. injection with a range of doses (10, 30, 100, 300 and, 500 μg) and vehicle (PBS)-treated animals received an injection of 10 μl of sterile Ca²⁺ and Mg²⁺ free PBS. Two weeks after treatment, the level of Kcnt1 mRNA was determined using quantitative reverse transcription PCR (RT-qPCR) and compared to that of vehicle (PBS)-treated mice. The administration of Kcnt1 ASO produced a dose-dependent knockdown of Kcnt1 mRNA in the cortex and the spinal cord (Figure 3, A and B). We then determined the specificity of the Kcnt1-targetted knockdown by evaluating the expression of the highly homologous gene Kcnt2 in +/+ mice treated with a bolus i.c.v. injection of 250 μg Kcnt1 ASO and control ASO (non-hybridizing sequence). Kcnt1 ASO produced a robust knockdown of Kcnt1 mRNA in the cortex compared to untreated mice (Figure 3C) but did not affect the expression of Kcnt2 mRNA (Figure 3D). No significant differences in Kcnt1 gene expression were found between control ASO treated and untreated mice, confirming that the knockdown observed in Kcnt1 ASO treated mice resulted from on-target hybridization. A consistent reduction of the protein was observed 2 weeks after injection with ED50, ED80 and 500 μg of Kcnt1 ASO (Figure 3, E-F). Using a pan-ASO antibody, which recognizes the
phosphorothioate (PS) backbone of the ASO (50), we confirmed broad ASO distribution in the mouse brain after a single bolus injection of 75 μg of Kcnt1 ASO (Figures 3G).

**ASO mediated Kcnt1 mRNA reduction rescues the seizure phenotype in L/L mice and improves overall behavior and survival**

Survival, seizure frequency, and behavioral markers (nesting, exploratory behavior, anxiety like [light/dark box test, and EPM]) were examined to determine the efficacy of Kcnt1 knockdown. Adult L/L mice were treated with a single bolus of Kcnt1 ASO at either ED50 (35μg), ED80 (75μg), or 500 μg. The control group received a dose of 500 μg of control ASO. Treating adult symptomatic L/L mice with a single dose of Kcnt1 ASO resulted in an increase in survival. This effect was dose-dependent and statistically significant across all the tested doses (median survival of 209 days for ED50, 256 for ED80 and >300 days for 500 μg Kcnt1 ASO) (Figure 4B). In contrast, the animals treated with control ASO displayed a survival consistent with that of untreated L/L mice (median survival of 73 days for control ASO and 65 days for untreated mice). Mortality was often related to seizure and presented as status epilepticus.

L/L mice treated with a dose of ED80 and 500 μg of Kcnt1 ASO had a significant reduction in the total seizure frequency; while the difference in the ED50 group was not statistically significant, a trend towards reduction was observed. In the group that received the control ASO, a significant increase in seizure frequency was seen (Figure 4D). Further, the presence of acute spikes was also reduced in Kcnt1 ASO treated animals three weeks after treatment (Figure 4C). Age-matched untreated L/L mice were used as controls for ECoG, as the severity of the Kcnt1 phenotype prohibited meaningful electrical recordings to be made in mice treated with control ASOs. During ECoG recording, none of the ASO treated mice presented seizures, while there were 5 seizures recorded in the untreated group (representative traces of ECoG signal in treated and untreated
animals are presented in Supplemental Figure 3). Altogether, these data indicate that the selective knockdown of Kcnt1 mRNA has an anticonvulsant effect in our mouse model. In addition, mice treated with the Kcnt1 ASO became less likely to exhibit seizures during routine handling, with no mice presenting seizures during the period of behavioral testing. In contrast, control ASO-treated mice continued to present seizures during routine handling.

Nesting behavior was evaluated 10-13 days after i.c.v. injection. 48-hour after receiving new nesting material, an improvement in nesting behavior was noticed in all animals treated with Kcnt1 ASO, independent of the dose received (Figure 4, E-F). Control treated mice displayed poor nesting behavior, similar to what we observed in untreated L/L mice.

L/L mice treated with Kcnt1 ASO ED50 and ED80 showed an exploratory behavior similar to that of +/-, while mice that received 500 μg showed a markedly reduced exploratory behavior (Figure 5A). In the light/dark box test, we found no difference in the time spent in the light compartment between mice treated with Kcnt1 ASO and control ASO, showing an increase in this parameter for all mice that received an i.c.v. injection (Figure 5B). The EPM test showed that independent of the dose received, all Kcnt1 ASO treated mice spent less time in the open arms of the maze compared to control ASO treated mice (Figure 5C). These data indicate that the ASO silencing of Kcnt1 not only reduces seizure burden but also normalizes behavioral markers of disease. Treatment with Kcnt1 ASO did not alter the behavior of the L/L mice in the Y-maze compared to untreated mice (Figure 5D).

**ASO-mediated Kcnt1 reduction improves general health and allows mating behavior, pregnancy and parental behavior of L/L mice**

Due to the severity of the L/L mice phenotype, early mortality and behavioral abnormalities impaired normal mating behavior. To test if mating behavior could be restored, L/L mice received
a single dose of 250 μg of Kcnt1 ASO at P40. Two weeks after injection, male and female mice were set up as breeding pairs or trios. ASO treated dams gave birth to litters of 5 to 8 homozygous pups. Although the first litter was often neglected, both male and female mice displayed effective parental behavior after the second pregnancy. The resulting L/L offspring displayed a phenotype as severe as the homozygous mice obtained from heterozygous breeding if left untreated.

**Neonatal administration of Kcnt1 ASO is efficacious and well tolerated**

To determine if the levels of Kcnt1 mRNA could be reduced in a dose-dependent fashion at the neonatal period, we tested the Kcnt1 ASO in newborn +/- mice. At P2, the Kcnt1 ASO was delivered by ICV injection with a range of doses (0.5, 1, 1.5, 3, 6 and 30 μg) and vehicle (PBS)-treated animals received an injection of 2 μl of sterile Ca²⁺ and Mg²⁺ free PBS. Two weeks after treatment, the level of Kcnt1 mRNA was determined and compared to that of vehicle (PBS)-injected mice. The early administration of Kcnt1 ASO produced a dose-dependent knockdown of Kcnt1 mRNA in the mouse cortex (**Figure 6A**).

Then we examined whether the early administration of Kcnt1 ASO could prevent the development of a disease phenotype in the L/L model. At P2-P3, L/L mice received a single i.c.v. bolus injection of 3.4 μg (ED80), while control ASO mice received an injection of 50 μg. Kcnt1 ASO treated mice showed extended survival, with the first deaths being observed at P135 (**Figure 6B**) and displaying approximately 25% mortality by P150. Similarly, control ASO treated mice also showed a shorter yet significant improvement in survival (median survival of 63 days compared to 46.5 for untreated mice). Seizure frequency improved significantly in mice treated with Kcnt1 ASO (**Figure 6E**). Bodyweight was measured at P40 and was increased for Kcnt1 ASO L/L mice compared to that of controls but overall, body weight was still lower compared to +/- mice (**Figure 6C**). Similar to adult treated L/L mice, behavioral markers including nesting and hyperactive exploratory activity
(locomotor cells) and EPM showed improvement compared to untreated mice (Figure 6, D, G, H).

Considering the likelihood that an ASO therapy for KCNT1-DEE in patients would require chronic ASO administration, we tested if a second dose was tolerated. A small group of +/- mice received a 3.4 μg dose of Kcnt1 ASO at P2 and a second injection of either Kcnt1 ASO (35 or 75 μg) or control ASO (500 μg) at P30. The procedure was tolerated in all treatment groups. By P42, the administration of a second dose of 75 μg Kcnt1 ASO showed a further reduction of Kcnt1 mRNA compared to mice receiving control ASO on the second injection (Figure 6I).

In a small subset of L/L mice (n=4) treated at P2, we tested if a second dose of Kcnt1 ASO (75 μg) could be effective after the animals have become symptomatic (i.e. nesting behavior was deficient and seizures at handling were observed in 1 mouse). In these cases, a further extension in survival was observed (Figure 6J) and nesting behavior was rescued (Figure 6K).

ASO-mediated Kcnt1 reduction in +/- mice is well tolerated

To evaluate the consequence of Kcnt1 reduction, we treated adult +/- mice with 500 μg Kcnt1 ASO and performed an array of behavior tests. ASO-mediated knockdown of Kcnt1 mRNA in +/- mice did not affect their nesting behavior (Supplemental Figure 2A) and was consistent with a previously reported study on a Kcnt1 knockout (KO) mouse model. The exploratory behavior in the locomotor cell test was similar to that of control-ASO treated mice and although the total ambulatory distance explored displayed a tendency towards reduced activity, this difference did not reach statistical significance (Supplemental Figure 2B). Similarly, a tendency to spend less time in the light chamber was observed (Supplemental Figure 2C).
On the EPM, +/- Kentl ASO treated mice showed a behavior similar to that observed in L/L Kentl ASO treated mice, with a reduction in the time spent in the open arms (Supplemental Figure 2D). In the three-chamber social interaction test, no difference was found between control and Kentl ASO treated mice in the time spent with an intruder mouse (Supplemental Figure 2E). Finally, to further explore changes in memory, the mice were tested in the Y maze. A reduction in the median time spent in the novel arm was observed in Kentl ASO treated mice compared to control (Supplemental Figure 2F).
DISCUSSION

In this study we investigated the potential of an ASO-based gene silencing approach as a potential therapy for KCNT1-DEE. We have shown a gene-specific and dose-dependent knockdown of Kcnt1 mRNA achieved with the Kcnt1 ASO in L/L mice which significantly improved survival, epilepsy, behavioral comorbidities and complete rescue of mating.

We first developed a rodent model and established disease biomarkers in homozygous animals that could be used for therapeutic screening. While heterozygosity for a gain of function variant in KCNT1 is sufficient to result in a disease state in patients with gain of function KCNT1 variants, this was not replicated in our mouse model. L/+ mice did not display an epileptic phenotype, increased susceptibility to seizures or pathologic behavioral changes. This is not completely unexpected for mouse models of genetic epilepsy demonstrate considerable phenotypic variability based on factors including strain (51) and sub-strain (52) used. In contrast, homozygous mice presented a robust epilepsy phenotype, behavioral deficits and a progressive deteriorating course resulting in reduced survival, thus presenting a valuable tool for drug screening.

Recently, Shore et al reported on a gain of function rodent model of KCNT1 epilepsy. This model was based on the KCNT1 p.Y796H variant, which has been identified as causing both inherited and de novo severe, early-onset Autosomal Dominant Nocturnal Frontal Lobe Epilepsy (ADNFLE) (25, 53). Patients with ADNFLE present with frequent, mostly nocturnal seizures and neuropsychiatric features (3, 7, 25, 27). Similar to our model, the Kcnt1 p.Y777H heterozygous mice did not exhibit spontaneous epileptiform activity while the homozygous mice showed generalized tonic-clonic seizures and tonic seizures. In vivo electrocorticography in pups localized these seizures to the somatosensory cortex and electrophysiology showed changes in membrane excitability with inhibitory-neuron-specific impairments inaction potential firing. Furthermore, the
Kcnt1 p.Y777H homozygous mice exhibited a hyperactivity phenotype in the open field and poor nesting behaviour, similar to what we observed in our model. Interestingly, the p.Y777H model did not show alterations on the elevated plus maze, opposite to what we observed with our homozygous model.

Within 2 weeks of bolus administration, ED80 and 500 μg doses of Kcnt1 ASO produced a significant reversal of the epileptic phenotype. While dosing at ED50 showed only a trend towards seizure reduction, which could have been influenced by the sample size on this treatment group. A clear dose-dependent effect was observed for survival of adult L/L treated mice. Overall, the effectiveness of the ASO approach in controlling seizure activity confirms that brain hyperexcitability in this disease model is driven by Kcnt1 gain of function and therefore by reducing the excessive channel activity the seizure threshold can be modified.

A critical concern shared among neurodevelopmental conditions and neurodegenerative disorders is whether disease progression can be stopped and the cognitive capacity preserved or improved once the pathological process has begun. Importantly, we have shown here that the downregulation of Kcnt1 can be safe and effective in both neonatal (pre-symptomatic) and adult (symptomatic) mice by not only in controlling spontaneous seizures but also providing disease modifying effects as indicated by improvements in cognition and behavior.

We also asked if the down-regulatory ASOs had any harmful effects derived from excessive knockdown. Studies on Kcnt1 KO mice have reported some mild behavioral and cognitive alterations (54, 55). Here we show that the administration of a high dose of Kcnt1 ASO to adult (500 µg) and neonatal (30 µg) +/+ mice did not result in serious adverse events. Treated adult mice showed a nesting behavior similar to that of untreated mice, and consistent with the previously reported phenotype of Kcnt1 KO (55).
A reduced exploratory behavior in the open field was reported in the *Kcnt1* KO model characterization published by Bausch *et al* (54). Interestingly, while the initial exploratory activity of wild-type mice treated with *Kcnt1* ASO was similar to that of the control treated group, we identified a trend that suggests an overall reduced exploratory activity. Similarly, in L/L mice treated with 500 μg of *Kcnt1* ASO, we observed a significant reduction in locomotor activity when compared to +/+ . This behavior completely contrasted the hyperactivity observed in the untreated L/L mice and further supports that *Kcnt1* plays a role in the exploratory response to new environments. While seizures precluded us from testing the behavioral battery before and after treatment in our model, the overall improvement in survival and the observed improvement in behavior compared to untreated and ASO control treated mice are consistent with significant improvement in cognition.

RNA-targeted ASO technology is an innovative therapeutic modality that has already achieved clinical success as an effective treatment for rare neurogenetic conditions like spinal muscular atrophy (43), Duchene’s muscular dystrophy (44) and hereditary transthyretin amyloidosis (56). The FDA approval of nusinersen marked an important milestone for ASO technology. Nusinersen has demonstrated the disease-modifying properties and fulfilled the promise of precision medicine, opening a path for the further development of other RNA based therapies for neurogenetic diseases (38). Although CNS targeting ASOs require direct intrathecal delivery to be effective, this invasive approach is compensated by the extended half-life in the target tissue, and the wide brain distribution and high cellular uptake (57–60) which allows for less frequent dosing. In addition, direct delivery to the CSF compartment precludes the ASO from being distributed to the rest of the body, limiting the occurrence of adverse effects (57).
There is growing evidence that supports the development of ASO as a therapy for DEE. Successful use of ASOs for severe epilepsies has been reported in preclinical models of SCN1A (61), SCN2A (62) and SCN8A encephalopathy (63). Targeted Augmentation of Nuclear Gene Output (TANGO) ASO targets naturally occurring, non-productive alternative splicing events to specifically reduce nonproductive mRNA and increase productive mRNA and protein of a target gene (64). Therefore, this approach can be used to address conditions with monogenic loss-of-function as a pathomechanism (64). Huat Lim showed that upregulation is not limited by the size of the gene and the effect is titratable. Further, Hu et al have shown that the TANGO ASO modality can be used to upregulate the WT allele and compensate for the defective allele in a model of Dravet Syndrome (SCN1A encephalopathy). A significant reduction of the incidence of seizures and mortality related to sudden unexpected death in epilepsy were observed after a single i.c.v. administration of a TANGO ASO at P2 and P14 in a mouse model of Dravet syndrome (61). Furthermore, Lenk et al have demonstrated the efficacy of a downregulating ASO targeting Scn8a in preclinical rodent models of both SCN8A and SCN1A DEE (63). Recently, Li et al tested a PS gapmer ASO targeting Scn2a mRNA in a mouse model of early onset SCN2A DEE. The Scn2a gapmer was shown to reduce spontaneous seizures, improve the behaviour (making it indistinguishable from wild-type animals) and significantly extend the lifespan of the treated mice, suggesting that ASO treatment can be effective and well tolerated (62).

Genetic neurodevelopmental conditions, including DEE are devastating conditions that lack effective therapies. The unravelling of the genetic architecture of DEE has provided an exciting opportunity to develop precision medicines. Therapeutics that use RNA targeting molecules are well positioned to directly address the underlying cause of DEE including seizures and comorbid pathologies. In this study, we demonstrated that ASO-mediated reduction of Kcnt1 is safe and has
a disease-modifying effect in a homozygous mouse model of a *KCNT1*-DEE pathogenic variant. These findings provide crucial evidence to support the development of ASO based therapies for refractory epilepsies and developmental disorders that can contribute to reducing the impact of neurogenetic disorders.
METHODS

*Kcnt1* knock-in mouse model

Mice were housed in temperature-controlled rooms (≈22°C) on a 12 h dark/light cycle, weaned at ≈ postnatal day 21 (P21), and maintained with rodent diet and water available *ad libitum*. Male and female mice were used for all experiments. Mice were housed in individually ventilated cages (IVC, Tecniplast, Sealsafe plus mouse IVC green line) in groups of up to five animals. Mice that underwent ECoG electrode implantation, seizure monitoring, and nesting behavior assessment were housed individually during recovery of surgical procedures and for the duration of the test.

*Mouse model generation*

The L/L knock-in mouse line was generated at the University of Adelaide using CRISPR/Cas9 to insert a single nucleotide variant at position c.2714, changing C>T. The *Kcnt1* p.P905L variant is homologous to the *KCNT1* p.P924L heterozygous pathogenic variant which has been found in two patients with EIMFS (65, 66). *Kcnt1* p.P905L founders were generated as described previously (67). C57BL/6J zygotes were injected with 50 ng/μl of sgRNA (TGCATGAACCGCATGTTGGA), 100 ng/μl of Cas9 mRNA and 100 ng/μl of a single-stranded oligonucleotide repair template (Ultramer DNA from Integrated DNA Technologies). The oligonucleotide repair template sequence was:

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GTTTGGAAAGAGCCAGAGAGTAGCTGTCCCTGGCACCGAACTGCATGAACCGCATG
TTGAaGCTTGTGAGCTCCGTTGGGATGCTGAGACTGGGGAAAAGCCTGAGGGA
GGATGATCG.
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Injected embryos were transferred to pseudopregnant females for further development. Pups were genotyped by PCR and the intended C>T variant was confirmed by Sanger sequencing. The colony
was maintained on the C57BL/6J background and initially backcrossed to wild-type (+/+) mice to expand the colony with subsequent heterozygous intercross carried out to obtain wild-type (+/+), heterozygous (L+/+) and homozygous (L/L) mice. Where possible +/+ littermates were used as controls. When unavailable or as “intruders” for behavioral paradigms (specifically the three chamber social interaction test), C57BL/6J wild-type mice were purchased from the Animal Resources Centre (Canning Vale, WA, Australia).

For experiments with neonatal mice: C57BL/6J wild-type pregnant mice were purchased from the Animal Resources Center (Canning Vale, WA, Australia). L/L litters were obtained from the crossing of Kcnt1 ASO-treated L/L breeders.

Identification and genotyping

Animals were toe-clipped at postnatal days 8 to 12 (P8 - P12) for identification and the tip of the tail was biopsied for genotyping. DNA extraction was performed using the REDExtract-N-Amp Tissue PCR Kit (Sigma, St Louis, MO, USA).

The following PCR primers were used to amplify exon 24 of Kcnt1:

Forward: 5’-CCACCCAGTTATGACCACAG- 3’
Reverse: 5’-GCTGTTAGGTATCTGTTAGCAG- 3’

PCR products were digested with the restriction enzyme BstBI (New England BiolLabs Inc, Ipswich, MA, USA) and the products were separated through electrophoresis on a 2% agarose gel stained with GelRed (Biotium, Fremont, CA, USA). The wild-type allele generated a single fragment of 460 bp and the mutant allele generated two fragments of 276 and 184 bp.

Mouse model phenotypic validation

Spontaneous behavioral seizures video recording
Mice older than P21 were housed individually in a 19.56 x 30.91 x 13.34 cm Thoren # 9 Small mouse II cage (Thoren Caging System, Hazleton, PA, USA) and monitored continuously for up to 5 days. Video recording was done using the Vivotek video server (VS8102) connected to an infrared day and night digital color camera (EVO2; Pacific Communications). Following recording, the videos were played back on a computer at 8-13x speed using the VLC media player (Paris, France). Seizures that presented with any of the following behaviors were counted as an event: a) clonic seizure in a sitting position; b) clonic and/or tonic–clonic seizures while lying on the belly c) pure tonic seizures; d) clonic and/or tonic–clonic seizures while lying on the side and/or wild jumping. A tonic clonic seizure lasting more than 3 minutes was considered to be prolonged, and a seizure with a duration of more than 5 minutes was considered to be a status epilepticus.

Susceptibility to induced seizures

Thermogenic seizure assay: Mice between the ages of P18-P21 were placed in a container heated to constant 42°C for a maximum of 20 minutes. The latency to a first tonic-clonic seizure was recorded.

Pentylenetetrazol induced seizures: Mice between the ages of P30-P40 were injected subcutaneously (s.c. injection) with the GABA antagonist pentylenetetrazol (PTZ; 80mg.kg⁻¹; Sigma, St. Louis, MO, USA) dissolved in 0.9% sterile saline solution (Pfizer, NY, USA). The animals were then monitored for a maximum of 45 minutes and latencies to a minimal (first tonic-clonic) and maximal (tonic hindlimb extension) seizures were recorded.

Loxapine induced seizures: Mice between the ages of P30-P40 received a peritoneal injection (i.p. injection) with the antipsychotic drug Loxapine (LOX; 100mg.kg⁻¹; Sigma, St. Louis, MO, USA) dissolved in 0.9% sterile saline solution (Pfizer, NY, USA) and 25% Dimethyl Sulfoxide (DMSO, Sigma, St. Louis, MO, USA). The animals were then monitored for a maximum of 1 hour and the
latency to a first tonic-clonic seizure was recorded.

In the three assays, animals were euthanized at the end of the experiment.

*Behavioral profile*

Behavioral tests were conducted between 9:00 A.M. and 6:00 P.M., under similar lighting conditions for each task. The mice were transferred to the behavior testing room at least 30 min before the test. All mice were subjected to each behavioral test with a 5-7-day interval between each task.

Nesting behavior: Two facial tissues (18 x 18.5 cm ;1.75-1.9 g) (Austwide Paper products, VIC, Australia) were provided as nesting material two hours prior to the onset of the dark phase (5-6pm). The quality of the nest was assessed 48h after. A score from 0 to 5 was given by adapting the scoring system of Hess et al (46).

Locomotor cells: Mice were placed in the center of a 27.31 x 27.31 x 20.32 cm covered chamber (Med Associates Cat# ENV-510S-A) for 30 min and their activity was monitored by a 48 channel infrared (IR) controller in 5 min bins and analyzed with the Activity Monitor Software Version 7 (Med Associates, Fairfax, VT, USA).

Light/Dark Box: The light–dark apparatus consisted of a 27.31 x 27.31 x 20.32 cm chamber (Med Associates Cat# ENV-510S-A) divided into dark and light compartments of equal size by the insertion of a black Perspex box. The box contained a small opening in the middle allowing the mouse to move between the compartments. The light chamber was brightly illuminated to 750 Lux by LED light lamps. Each mouse was placed in the center of the dark compartment and allowed to freely explore the box for 10 minutes. The latency to move into the light compartment as well as the amount of time spent in each side were automatically recorded and analyzed with the
Activity Monitor software version 7 (Med Associates, Fairfax, VT, USA).

Elevated plus maze (EPM): The elevated plus-maze apparatus was made of light-colored Perspex and consisted of two open arms (5 x 30 cm) and two enclosed arms (5 x 30 x 14 cm) extending from a central area (5 x 5 cm). A raised lip (2.5 mm high, 5 mm wide) around the open arms was placed to reduce the likelihood of mice falling off the maze. The maze was elevated 60 cm above the floor. Each mouse was placed in the central area, facing an open arm. Their activity in the maze was video recorded for 10 min. The time spent on each arm of the maze and in the center area was measured, and arm entries were counted using the TopScan software (CleverSys, Reston, VA, USA).

Three chamber social interaction: The test was conducted in a box (43 x 39 x 11 cm) made of transparent plastic and divided into three chambers; the middle section was 8 x 39 cm and the two lateral sections were 16 x 39 cm each. The chambers were connected by rectangular openings in the middle. Metal mesh cages of 16 x 10 x 11 cm were placed in the two side chambers. The test consisted of two phases: habituation and trial phase. During the habituation period, the mouse was placed in the central chamber and allowed to explore freely all chambers for 10 min. After that, a preference for a chamber (left vs right) was assessed and an unfamiliar mouse of similar age and size was placed in the metal cage of the side chamber of least preference. For the trial period, the mouse was again allowed to explore all chambers for 10 min. The activity during both habituation and trial period was video-recorded and the amount of time spent in each chamber and near the cages was analyzed using TopScan software (CleverSys, Reston, VA, USA).

Y-maze: The Y-maze was made of light-colored Perspex and consisted of three 7.5 x 30 x 14 cm arms separated by a 120° angle and containing a visual cue at the end. The test was performed in two phases: a training session and a trial session. During the training session the mouse was placed
in the far end of one arm (home arm), facing away from other arms, and was allowed to explore this and one other arm (familiar arm) for 10 min. During this period, the third arm (novel arm) was not accessible. After a 1-hour interval, the mouse was again placed in the home arm but allowed to explore all three arms for 5 min (trial session). The test was video recorded, and the amount of time spent in each arm was analyzed using TopScan software (CleverSys, Reston, VA).

**Surgical procedures**

_ECoG Electrode implantation surgery:_ Mice were anaesthetized with 3-4 % Isoflurane (IsoFlo; Abbott Laboratories, Abbott Park, IL, USA) for induction and 2% for maintenance. A subcutaneous injection of meloxicam (1 mg/kg) dissolved in 0.9% saline was given prior surgery. Then, the mouse was placed in a stereotaxic apparatus (Kopf Instruments, Tujunga, CA, USA), after the scalp was shaved, sterilized with 80% ethanol and infiltrated with Lignocaine (1% ampules, Pfizer). A 1 cm long incision was made on the scalp and the skull was cleaned with 3% hydrogen peroxide solution (Sanofi). Four burr holes were drilled into the skull and three screws were placed into the holes and used as epidural electrodes (Cat #8403, Pinnacle Technology, Lawrence, KS, USA) (1 reference electrode and 2 recording electrodes). A ball of silver wire was used as ground electrode. All electrodes were connected to a headmount (Cat # 8201, Pinnacle Technology, Lawrence, KS, USA) and affixed to the skull with methyl methacrylate dental cement (Cat # 1255710; Henry Schein Inc, Indianapolis, IN; Lang Jet Denture Repair Acrylic). Then, mice recovered in a warming pad at 30 °C until fully awake.

_ECoG recording and analysis:_ Mice were allowed to recover for 3-5 days before recording. The mice were connected while awake to a mouse pre amplifier (Cat # 8406-SE) and an amplifier (Cat #8204 and 8206, Pinnacle Technology, Lawrence, KS, USA). Brain cortical activity was sampled on Sirenia (Pinnacle Technology, Lawrence, KS, USA) at 250 Hz for 25 hours, signal was band-
pass filtered at 0.5 to 40Hz. ECoG signal was analyzed post acquisition using ClampFit 10.7 (Molecular Devices), a spike was identified when the amplitude was 2.5 times greater than the baseline activity and duration was shorter than 80 ms. An investigator blinded to the treatment groups/genotype reviewed and counted spike activity.

*Intracerebroventricular (i.c.v.) injections*

**Adult mice:** P40-P45 mice, weighing at least 10g, were anesthetized using Isoflurane (IsoFlo; Abbott Laboratories, Abbott Park, IL, USA) at a concentration of 4-5% mixed in O₂ (vol/vol) for induction, and 2% for maintenance. A subcutaneous injection of meloxicam (1 mg/kg) dissolved in 0.9% saline was given prior surgery. The mice were positioned in a stereotaxic apparatus (myNeuroLab, Leica, Germany) and the scalp was cleaned with 80% Ethanol. Then, the skin was infiltrated with Lignocaine (1% ampules, Pfizer) and a 1 cm incision was performed. The skull was cleaned with 3% hydrogen peroxide solution to expose the bone sutures and a burr hole was drilled at 0.8 mm lateral and 0.3 mm posterior to Bregma. The tip of a 33G internal infusion cannula (Plastics One Cat #C315I/SPC) was advanced to −3.0 mm from the skull surface to reach the right lateral ventricle. The cannula was connected to a 0.5 ml glass syringe (SDR, Sydney, Australia) and an infusion pump (legato 210/210p syringe pump, Kd Scientific, Holliston, MA, USA). A total volume of 10 µl of ASO or vehicle (sterile Ca²⁺ and Mg²⁺ free PBS) was delivered at a rate of 0.5 µl/s. One minute after completing the infusion, the cannula was withdrawn and the skin was closed with 4-0 polyglactin 910 absorbable suture (coated Vycryl, Ethicon Inc, Cornelia, GA, USA). Following surgery, the animals recovered on a Thermacage (Datesand, Ltd, Manchester, UK) until active and were then returned to their home cage.

**Neonatal mice:** P2-P3 pups were cryo-anesthetized for 3 minutes. The scalp was cleaned with 80% ethanol and free hand injections were performed using a 32 G needle Hamilton syringe (10µl,
Hamilton, Reno, NV, USA). Using lambda and the right eye as anatomical references, the needle was inserted midway between these two points and advanced -2.0 mm ventral from the skin surface (technique adapted from Kim et al (68)). A total volume of 2 μl of ASO or vehicle (sterile Ca\textsuperscript{2+} and Mg\textsuperscript{2+} free PBS) was injected into the right lateral ventricle. Pups were gently warmed until skin color returned to pink and rolled over dirty bedding before being returned to their home cage. Maternal behavior was monitored for the following 10 minutes to ensure the pups were attended and returned to the nest.

**ASO synthesis**

The *Kcnt1* ASO and control ASO (non-hybridizing sequence) were synthesized and screened in vitro by Ionis Pharmaceuticals (Carlsband, CA, USA) as previously described (69, 70). Both molecules are 20bp long and have the following structure: 5 modified nucleotides with 2’- MOE modifications at the 5’ and 3’ end, and a central gap (gapmer) of 10 unmodified oligodeoxynucleotides. A PS backbone was used to enhance nuclease resistance. *Kcnt1* ASO targets the mRNA 3’ untranslated region at position corresponding to 3981-4000 in transcript variant 1 (NM_175462.4).

The specific sequence for both ASOs is listed below:

*Kcnt1* ASO: 5’-GCTTCATGCCACTTTCCAGA-3’

Control ASO: 5’-CCTATAGGACTATCCAGGAA-3’

ASOs were solubilized in sterile Ca\textsuperscript{2+} and Mg\textsuperscript{2+} free Dulbecco’s Phosphate Buffered Saline (PBS) (Sigma, St Louis, MO, USA), centrifuged, and filtered through a 0.22 μm filter (Sigma, St Louis, MO, USA). A stock solution with a concentration of 50 mg/ml was stored at -20ºC. Then, ASOs were further diluted to the desired concentration in sterile Ca\textsuperscript{2+} and Mg\textsuperscript{2+} free PBS (Sigma, St
Louis, MO, USA) immediately before injection.

**Quantification of mRNA levels**

Tissue harvest: Two weeks after ASO administration, mice were deeply anesthetized with 4-5% Isoflurane (IsoFlo; Abbott Laboratories, Abbott Park, IL, USA), before being decapitated. After the brain was removed from the skull, the cerebellum, right and left cortex were dissected. The spinal cord was harvested using hydraulic extrusion as previously described (71) and a 1 cm piece from the low thoracic/lumbar cord was extracted. All dissected tissue was snap frozen in liquid nitrogen, and stored at −80°C until RNA preparation.

Quantitative gene expression analysis: Total RNA was isolated from the mouse right cortex and spinal cord using TRIzol reagent (Cat #15596026, Thermofisher, Waltham, MA, USA) according to the manufacturer’s protocols. Contaminating genomic DNA was removed with rDNase I treatment (DNA-free Reagents; Ambion/Life Technologies, Carlsband, CA, USA). RNA was assayed for quality and quantity using a NanoDrop 2000c Spectrophotometer (Thermo Scientific).

For quantitative reverse transcription and polymerase chain reaction (RT-qPCR) presented in Figure 3 A-D, oligo deoxythymidylic acid (oligo-dT) primed cDNA was synthesized from 500 ng of total RNA using Murine Moloney Leukaemia Virus Reverse Transcriptase (Promega, Madison, WI, USA). RT-qPCR was performed on the ViiA 7 Real-Time PCR System (Applied Biosystems/Thermofisher, Foster City, CA, USA) using SYBR green technology and GoTaq qPCR master mix (Promega) according to the manufacturer’s protocols. The primers used for detection were:

\[ Kcnt1 \]

Forward: 5’-TCTTCCCTTTCTCAGGTCCAGG-3’
Reverse: 5’-AGGGAGAAGTTGAACAGCCG-3’

*Kcnt2*

Forward: 5’-AAGGCTGAGCAGAAAAGGGC-3’

Reverse: 5’-TCATTTCATCGTAGCCACCG-3’

*Rpl32*

Forward: 5’-GAGGTGCTGCTGATGTGC-3’

Reverse: 5’-GGCGTTGGGATTGGTGACT-3’

Relative gene expression values were obtained by normalization to the reference gene Rpl32 using the $-2\Delta\Delta Ct$ method, where $-2\Delta\Delta Ct = \Delta Ct$ sample−$\Delta Ct$ calibrator as previously described (72).

For Figures 6A, 6H and supplemental figure 3B, RT-qPCR, 1µg of rDNase I treated purified RNA was analysed using a Taqman™ RNA to Ct kit process (Applied Biosystems) with a Taqman probe to *Kcnt1* [probe ID # Mm00558471_m1 (FAM-MGB)] in duplex with the mouse endogenous gene Glucuronidase Beta (*Gusb*) [probe ID # Mm01197698_m1 (VIC-MGB)]. RT-qPCR was performed on the ViiA 7 Real-Time PCR System (Applied Biosystems) using cycling conditions specified by the manufacturer. All RT-qPCR reactions were performed in triplicate. Relative gene abundance values were calculated by normalization to *Gusb* and referenced to the control groups using the $-2\Delta\Delta Ct$ method.

**Western Blot**

The quantification of Kcnt1 protein levels was performed by Western blotting. Total protein lysate was prepared from the mouse left hemisphere. The Poly (vinylidene fluoride) (PVDF) membrane was incubated with an antibody against Kcnt1 protein (Anti KCNT1 mse monoclonal [N3/26]
8ug/mL, NeuroMab CAT # 75-051), Anti β-actin (Mse IgG1 monoclonal dilute 1:10,000, SIGMA CAT # A5441) as loading control and IRDye infrared secondary antibodies (IRDye 800CW [LI-COR #926-32210] and IRDye 680RD [LI-COR # 926-68070]).

**Immunohistochemistry**

Tissue preparation: Two weeks after ASO administration, mice were anesthetized with sodium pentobarbitone (80 mg/kg, lethal dose, i.p. injection) (Ilium, Troy Laboratories, Smithfield, Australia) and transcardially perfused with 0.1 M phosphate buffer (PB) followed by a 4% paraformaldehyde solution (PFA; pH7.4). Brains were removed from the skull and stored in 20% sucrose solution overnight at 4°C, and then frozen in isopentane cooled in liquid nitrogen before being sectioned coronally at 20 μm thickness with a cryostat (Leica Microsystems, Wetzlar, Germany). Sections were mounted on Superfrost plus slides (Thermofisher, Waltham, MA, USA) and stored at -20 ºC until used for immunostaining.

Immunostaining: Sections were air-dried for 1 hour at room temperature and then blocked for 1 hour in a humidified chamber, with a mixture of 10% normal goat serum, 0.3% Triton X-100 (Sigma, St Louis, MO, USA) in PB. The blocked sections were incubated overnight with primary antibodies: polyclonal rabbit anti-ASO (against the phosphorothioate [PS] backbone) 1:7500 (Ionis Pharmaceuticals); polyclonal guinea-pig anti-NeuN 1:500 (Millipore; Cat #ABN90-clone A60 [MAB377]); After washing in PB, the sections were incubated for 2 h with secondary antibodies: goat-anti-rabbit Alexa-647 1:500 (Thermo-fisher; Cat#A31573) and donkey-anti-guinea pig Alexa 488 1:500 (Thermo-fisher; Cat# A-11073). To identify the nuclei, sections were then stained with DAPI (Sigma-Aldrich). All incubations were conducted at room temperature. Slides were covered with Prolong Gold Anti-fade (Invitrogen) and stored at -30 ºC. Confocal image stacks were acquired on a Zeiss LSM 780 microscope equipped with a 20 x/0.8 NA lens. Z-
stack images were acquired within Nyquist sampling parameters and de-convolved using Huygens 4.4.0 software (Scientific Software Imaging).

**Statistical analysis**

The latencies for the seizure susceptibility tests were plotted as Kaplan Meier survival curves and tested for significance using the Log rank (Mantel Cox) test. In all the survival plots, animals that did not reach the end point by 40 minutes (20 minutes for the thermogenic assay) were marked as censored. Dose-response curves were fitted with the Motulsky regression. For all other measures, Fischer test, Mann–Whitney U test, nonparametric one-way and two-way ANOVA with post hoc analysis were used accordingly. Statistics were computed using GraphPad Prism, San Diego, CA, USA. For all tests, statistical significance was set at a p < 0.05.

**Study approval**

All animal experiments were approved by the animal ethics committee of the Florey Institute of Neuroscience and Mental Health (protocols #14-026, 16-061, 16-062 and 17-014) and were performed in accordance with the guidelines of the National Health and Medical Research Council Code of Practice for the Care and Use of Animals for Experimental Purposes in Australia.

**Author contributions**

LEB, ML, NJ, PJ-N and S Pe designed the research. LEB, ML, NJ, PJ-N, BR, AS, ASo, KR, EG and LJ performed the experiments. SP, FA, PQT designed and generated the Kcnt1 mouse model. PJ-N, ASo and FR contributed ASOs and provided advice on designing experiments. CAR, SM, and S Pe supervised experimental design and provided advise on data interpretation.

LEB, CAR and SM wrote the manuscript. All authors read and contributed to the revision of manuscript.
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Conflict of interest

Steven Petrou has received payments from Praxis Precision Medicines. Paymaan Jafar-Nejad, Armand Soriano and Frank Rigo are paid employees of Ionis Pharmaceuticals.
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Figure 1. Phenotype of the Kcnt1 L/L mouse model. A. Difference in size at P21 of L/L mice (red arrow) compared to their L/+ and +/+ littermates. B. Body weight is significantly reduced in L/L mice compared to their +/+ littermates at weaning age (Kruskal-Wallis test, followed by Dunn’s post hoc analysis; +/+ n=15, L/+ n=21, L/L n=18), data is presented in a box and whiskers plot with maximal and minimal data points. C. Life span is shortened in L/L mice with a median survival of 43 days (Kaplan-Meier curve, Log-rank test p<0.0001; +/+ n=15, L/+ n=15 and L/L n=35). D. Representative ECoG trace of ictal activity and interictal spikes in the L/L mice. Top: Acute interictal spikes. Bottom: Seizures can be preceded by an increase in frequency of acute interictal spikes. Spontaneous tonic-clonic seizures correlated with fast, high amplitude signal, followed by electric suppression (black arrow). E. Acute high amplitude (>500 µV) spikes are present in the L/L mice, with a median of 1470 spikes in 24 hours (p=0.03, Kruskal-Wallis test, followed by Dunn’s post hoc analysis, n=3 for each genotype). F. Seizure frequency over 72 hrs (+/+ n=3, L/+ n=3, L/L n=6; median seizure frequency for L/L of 23 events. Kruskal-Wallis test, followed by Dunn’s post hoc analysis p=0.024).
Figure 2. Behavioral profile of the Kcnt1 L/L mouse model. A. Representative images of minimal and maximal scores of nesting behavior. The left image corresponds to a score of 5 (+/+ mouse), while the right image exemplifies a score of 1 (L/L mouse) B. Nesting behavior is impaired in L/L mice (+/+ n=7, L/+ n=7, L/L n=8, Kruskal Wallis test with Dunn’s post hoc analysis). C. Total ambulatory distance explored in the locomotor cells test. L/L mice are more active compared to L/+ and +/+ (+/+ n=12, L/+ n=20, L/L n=25, Kruskal-Wallis test with Dunn’s post hoc analysis). D. L/L mice spend less time in the light compartment during the light/dark box test (+/+ n=12, L/+ n=20, L/L n=17, Kruskal-Wallis test with Dunn’s post hoc analysis) E. L/L mice display a preference for the open arms of the elevated plus maze (+/+ n=10, L/+ n=11, L/L n=15, Kruskal-Wallis test with Dunn’s post hoc analysis).
Figure 3. *Kcnt1* ASO produces a dose dependent knockdown of *Kcnt1* mRNA in the mouse CNS. Dose response curves for *Kcnt1* ASO in the brain cortex (A) and thoracic spinal cord (B) for +/+ mice injected at P40. The tissue was collected 2 weeks after injection and processed for mRNA quantification (n=3 for each dose and PBS control, curves were fitted with the Motulsky regression). C. Mouse cortex was collected 2 weeks after i.c.v. injection for mRNA quantification. The i.c.v. administration of *Kcnt1* ASO reduced the levels of *Kcnt1* mRNA (Kruskal-Wallis test p=0.0027), without affecting the paralogous gene *Kcnt2* (Kruskal-Wallis test p=0.4794); Data is presented as bar plots with mean and SEM, untreated n=4, *Kcnt1* ASO n=5, Control ASO n=3. (D). Western blot showing a reduction of Kcnt1 protein in WT mice left hemisphere 2 weeks after i.c.v injection E. Average band signal for Kcnt1 protein. Data is presented as bar plots with mean and SD (n= 3 mice for each treatment condition, ns for all comparisons except PBS vs 500 ug, One way ANOVA with Dunnett’s multiple comparisons test). F. Distribution of the *Kcnt1* ASO in the mouse brain. Coronal brain sections of +/+ mice treated with *Kcnt1* ASO 75 μg. Tissue was collected 2 weeks after i.c.v. injection and stained with an ASO antibody (red), neuronal marker (NeuN; green) and counterstained with nuclear stain DAPI (blue). *Kcnt1* ASO was found throughout the meninges, hippocampus and cortical layers (n=3 experiments). Scale bars represent 500 μm.
Figure 4. ASO mediated knockdown of Kcnt1 at P40 markedly improves the disease phenotype of adult L/L mice. 

A. Experimental timeline for behavioral studies. 

B. Kaplan-Meier curves shows a dose dependent improvement in survival of adult L/L mice treated with Kcnt1 ASO (p<0.0001 for Kcnt1 ASO ED50, ED80 and 500 μg, log-rank test), while mice treated with control ASO showed a survival similar to that of untreated animals (p=0.237, log-rank test, untreated n=16, control ASO n=16, Kcnt1 ASO ED50 n=13, ED80 n=11, 500 μg n=11). 

C. Acute spike frequency over 24 hrs (untreated n=4, Kcnt1 ASO ED50 n=6, ED80 n=7, 500 μg n=8; ANOVA F [3, 21] = 7.978, p=0.001). 

D. Seizure frequency was significantly reduced after treatment with Kcnt1 ASO ED80 and 500 μg. Although ED50 did not reach statistical significance, a trend towards reduction was observed. Treatment with control ASO did not reduce the occurrence of seizures (control ASO n=9; ED50 n=8; ED80 n=10, 500 μg n=9; seizure frequency was compared using the non-parametric Wilcoxon matched-pairs signed rank test, with Pratt’s method for identical rows). 

E. Representative images of nesting behavior of Kcnt1 ASO ED80 (top) and control ASO treated (bottom) L/L mice. 

F. Nesting score of animals treated with Kcnt1 ASO showed a significant improvement compared to control ASO treated animals (ED50 vs control p=0.0002; ED80 vs control p=0.0015; 500 μg vs control p<0.0001; untreated vs control p>0.9. Kruskal-Wallis test with Dunn’s post hoc analysis. Untreated n=8, control ASO n=12; ED50 n=13; ED80 n=10, 500 μg n=12).
Figure 5. ASO mediated knockdown of Kcnt1 at P40 improves the behavioral phenotype of adult L/L mice. A. Total ambulatory distance was reduced in mice treated with Kcnt1 ASO 500 μg compared to control treated mice (p=0.0002), but not for mice treated with ED50 or ED80 (p=0.775 and 0.839, respectively) (Kruskal-Wallis test, followed by Dunn’s multiple comparisons, +/+ n=12, untreated n=25, control ASO n=13; ED50 n=13; ED80 n=10, 500 μg n=11). B. Time spent in the light compartment during the light/dark box test (Kruskal-Wallis test, followed by Dunn’s multiple comparisons p=0.588; +/+ n=12, L/L n=17, control ASO n=11, ED50 n=13, ED80 n=10, 500 μg n=12). C. Time spent in the open arms of the elevated plus maze (Kruskal-Wallis test, followed by Dunn’s post hoc analysis; +/+ n=10; untreated n=15; control ASO n=9; ED50 n=13; ED80 n=10, 500 μg n=12). D. Time spent in the novel arm of the Y-maze (Kruskal-Wallis test, followed by Dunn’s post hoc analysis p=0.551; +/+ n=8; untreated n=10; ED50 n=12; ED80 n=10, 500 μg n=10).
Figure 6. Neonatal administration of Kcnt1 ASO is safe and effective in the L/L mouse model.
A. Dose response curve for +/+ mice injected at P2 and tested 2 weeks after injection (n=3-11 per dose, the curve fitted with the Motulsky regression). B. Kaplan-Meier curve for neonatal (P2-P3) administration of Kcnt1 ASO in L/L mice showing prolonged lifespan of Kcnt1 ASO compared to Control ASO treated (p<0.0001). 50 μg of control ASO produced a smaller improvement in survival (p=0.003, log-rank test) (untreated n=36; control ASO n=23; Kcnt1 ASO 3.4 μg n=17). C. Weight at P40 of treated L/L mice improved (p=0.016, +/+ n=28, untreated n=17, control ASO 50 μg n=14, Kcnt1 ASO 3.4 μg n=17). D. Nesting behavior improved in L/L mice treated with Kcnt1 ASO (p =0.0001, untreated n=8; control ASO 50 μg n=9; Kcnt1 ASO 3.4 μg n=15). E. Seizure frequency was reduced after neonatal administration of Kcnt1 ASO when compared to control treated (50 μg) mice (p<0.0001, control ASO 50 μg n=15, Kcnt1 ASO 3.4 μg n=17). F. Total ambulatory distance was reduced in mice treated with Kcnt1 ASO 3.4 μg compared to untreated mice (+/+ n=12, untreated n=25, Kcnt1 ASO 3.4 μg n=17). G. Time spent in the open arms of the elevated plus maze (+/+ n=10, untreated n=15, Kcnt1 ASO 3.4 μg n=17). Mann-Whitney test (C-G). H. A second injection of Kcnt1 ASO at P30 was tolerated in +/+ mice with a further reduction of Kcnt1 mRNA compared to a single injection at P2. Cortical tissue was
collected at P42 [Kcntl ASO 3.4 μg n=3; Kcntl ASO 3.4 μg + Control ASO 500ug n=3; Kcntl ASO 3.4 μg + Kcntl ASO 35 μg n=2; Kcntl ASO 3.4 μg + Kcntl ASO 75 μg n=3]. I. Survival curve for L/L mice treated with a second dose of ED80 ASO at P167 (dotted line) (Untreated n=36; ED80 reinjected n=4). J. Nesting score of L/L mice reinjected with ED80 ASO. After the injection at P2, nesting behavior was present at P40 but had declined by P165. A second injection rescued the nesting behavior (n=4); Significant difference was found for P40 vs P165 (p<0.0001, One-way ANOVA with Dunnett’s multiple comparisons test).