bioRxiv preprint doi: https://doi.org/10.1101/538371; this version posted February 1, 2019. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission. Impaired M-current in KCNQ2 Encephalopathy Evokes Dyshomeostatic Modulation of 1 2 Excitability 3 4 Dina Simkin^{1,2}. Timothy J. Searl². Brandon N. Pivevskv¹. Marc Forrest^{3,4}. Luis A. Williams⁵. 5 Vaibhav Joshi⁵, Hongkang Zhang⁵, Steven J. Ryan⁵, Michael Schwake¹, Gabriella L. Robertson¹, 6 Peter Penzes^{3,4}, Linda C. Laux⁶, Owen B. McManus⁵, Graham T. Dempsey⁵, John J. Millichap⁶, 7 Alfred. L. George, Jr.^{2,*} and Evangelos Kiskinis^{1,3,*} 8 9 10 11 ¹The Ken & Ruth Davee Department of Neurology, Feinberg School of Medicine, Northwestern 12 University, Chicago, IL, USA 13 ²Department of Pharmacology, Feinberg School of Medicine, Northwestern University, Chicago, 14 15 IL, USA ³Department of Physiology, Feinberg School of Medicine, Northwestern University, Chicago, IL, 16 USA 17 18 ⁴Center for Autism and Neurodevelopment, Feinberg School of Medicine, Northwestern 19 University, Chicago, IL, USA

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31 **Conflicts of interest:** L.A.W., V.J., H.Z., S.J.R., O.B.M. and G.T.D. are employees and 32 shareholders at Q-State Biosciences. E.K. owns stock and is a consultant for Q-State 33 Biosciences.

34

35 ABSTRACT

Mutations in KCNQ2, which encodes a pore-forming K⁺ channel subunit responsible for 36 37 neuronal M-current, cause neonatal epileptic encephalopathy, a complex disorder presenting 38 with severe early-onset seizures and impaired neurodevelopment. The condition is exceptionally difficult to treat, partially because the effects of KCNQ2 mutations on the development and 39 40 function of human neurons are unknown. Here, we used induced pluripotent stem cells and gene editing to establish a disease model, and measured the functional properties of patient-derived 41 neurons using electrophysiological and optical approaches. We find that while patient-derived 42 excitatory neurons exhibit reduced M-current early, they develop intrinsic and network 43 hyperexcitability progressively. This hyperexcitability is associated with faster action potential 44 repolarization, larger afterhyperpolarization, and a functional enhancement of large conductance 45 Ca²⁺-activated K⁺ (BK) channels. These properties facilitate a burst-suppression firing pattern 46 that is reminiscent of the interictal electroencephalography pattern in patients. Importantly, we 47 were able to phenocopy these excitability features in control neurons only by chronic but not 48 acute pharmacological inhibition of M-current. Our findings suggest that dyshomeostatic 49 KCNQ2 loss-of-function compound and 50 mechanisms lead to alterations in the neurodevelopmental trajectory of patient-derived neurons. Our work has therapeutic implications 51 52 in explaining why KCNQ2 agonists are not beneficial unless started at an early disease stage.

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54 KEYWORDS

55 KCNQ2, Kv7.2, epileptic encephalopathy, human induced pluripotent stem cells, excitatory 56 neurons, M-current, epilepsy, dyshomeostatic and homeostatic plasticity, burst firing, disease 57 modeling

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67 INTRODUCTION

The KCNQ2 gene encodes Kv7.2 (referred to here as KCNQ2), a voltage-dependent 68 69 potassium (K⁺) channel widely distributed in central and peripheral neurons. In most mature 70 neurons, KCNQ2 and the paralogous KCNQ3 protein form heterotetramers (KCNQ2/3) (1). 71 Together these channels mediate the M-current, a slowly activating and non-inactivating voltage-72 dependent K⁺ conductance suppressed by Gq protein-coupled muscarinic acetylcholine receptor activation (2). The M-current activates as neurons approach action potential (AP) threshold and 73 acts to dampen neuronal excitability (2,3). Therefore, KCNQ2/3 channels help set the AP 74 threshold, and also contribute to the post-burst afterhyperpolarization (AHP), which limits 75 repetitive firing following bursts of action potentials (4,5). These channels are enriched at the 76 77 axon initial segment (AIS) and nodes of Ranvier of central and peripheral neurons (6-9), and are also expressed at lower densities at the soma, dendrites and synaptic terminals (8,10,11). 78

79 The importance of KCNQ2 in normal brain development and function is underscored by genetic epilepsies associated with this channel. Disorders caused by KCNQ2 mutation include 80 benion familial neonatal seizures (BFNS) characterized by seizures that spontaneously remit 81 within the first year of life (12,13), and the more severe neonatal epileptic encephalopathy (NEE), 82 which may present as Ohtahara syndrome or infantile spasms (14-17). A ClinVar search for 83 84 KCNQ2 variants with pathogenic, likely pathogenic, conflicting interpretations and uncertain significance results in 552 different variants some of which are recurrent in multiple patients, 85 86 accounting for approximately 5% of all mutations identified in genetic epilepsy (18,19) and 10% of those associated with early-onset forms of NEE (20). The main features of NEE are 87 developmental and cognitive disabilities, and early onset of severe seizures, occurring within a 88 few days after birth, that are refractory to antiepileptic drugs (21). 89

The earliest hypothesis to explain epilepsy associated with *KCNQ2* mutations posited that loss of KCNQ2 channel function allows for sustained membrane depolarization after a single action potential leading to increased repetitive firing within bursts in excitatory neurons (22). However, some variants associated with severe clinical phenotypes produce gain-of-function effects (23,24). Enhanced K⁺ conductance, specifically in the AIS, could hyperpolarize the AIS membrane, decreasing steady state inactivation for sodium channels. This would increase the rate of action potential activation and action potential repolarization (25).

97 The mechanisms by which developmental expression of KCNQ2 channels impact 98 neuronal excitability are not clear. What remains elusive is how the defects in M-current affect 99 the electrophysiological properties of human neurons leading to impaired neurodevelopment. 100 The use of patient-specific induced pluripotent stem cell (iPSC) technology has enabled a new approach for elucidating pathogenic mechanisms of genetic disorders such as the epileptic
channelopathies as it allows for the generation of otherwise inaccessible human neurons (2628). Here, we use KCNQ2-NEE patient-specific and isogenic control iPSC-derived excitatory
neurons to elucidate the dynamic functional effects of a prototypical *KCNQ2* mutation during
differentiation and maturation in culture.

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132 **RESULTS**

133 Establishing a Human Neuron Model of KCNQ2 Epileptic Encephalopathy

To investigate the effects of a KCNQ2 genetic variant in human neurons, we generated 134 135 iPSC lines from a 7-year old male clinically diagnosed with KCNQ2 NEE. The subject exhibited 136 seizures on the first day of his life (29), and had treatment-resistant epileptic spasms, myoclonic-137 tonic seizures and severe developmental delay. Genetic testing identified a de novo KCNQ2 138 variant (c.821C>T) resulting in a threonine to methionine missense mutation at position 274 139 (p.Thr274Met; T274M). The threonine residue is highly conserved and located in the pore domain 140 of the protein (Figure 1A). This prototypical mutation has been identified in at least 5 individuals 141 diagnosed with NEE (ClinVar).

We generated iPSCs from peripheral blood mononuclear cells (PBMCs) using integration-142 free, Sendai virus-mediated reprogramming. The resulting iPSCs exhibited a normal male 143 144 karyotype, typical stem cell morphology, and expressed pluripotency markers, including nuclear 145 NANOG and the cell surface antigen SSEA4 (Supplementary Figure 1, A and B). We validated the presence of the heterozygous mutation in the patient iPSCs by targeted PCR and Sanger 146 147 sequencing (Q2-01^{T274M/+}; Figure 1B). To create a model that would allow us to attribute any phenotypic differences to the disease-associated genetic variant, we generated an isogenic 148 149 control iPSC line from the patient-derived line. We specifically corrected the mutant allele using CRISPR/Cas9 genome editing and simultaneously introduced a silent mutation in the PAM site 150 151 to prevent re-cleavage (Supplementary Figure 1C). We identified a corrected isogenic clonal cell 152 line $(Q2-01^{+/+})$ that exhibited a normal karyotype (Figure 1B and Supplementary Figure 1B), and 153 found no evidence for off-target edits in any of the top 8 genomic regions with homology to the 154 targeted KCNQ2 exon (Supplementary Figure 1D-E and Supplementary Table 1).

155 Given the clinical presentation of *KCNQ2*-associated NEE and the focal source of seizures that reside in the cortex (30,31), we chose to study cortical excitatory neurons, differentiated 156 157 through a modified Nng2 overexpression protocol (Supplementary Figure 2A) (32,33). We simultaneously differentiated the Q2-01^{T274M/+} patient line, the engineered isogenic control line, 158 and two iPSC lines generated from unrelated, healthy, sex-matched controls (Supplementary 159 160 Figure 2B) (34). Co-expression of GFP marked lentiviral-transduced cells, which were co-161 cultured with primary mouse glia to facilitate in vitro neuronal maturation. To determine the 162 efficiency of differentiation, we used immunocytochemistry (ICC) to quantify the percentage of 163 MAP2 and GFP positive cells (Figure 1C). Over 85% of GFP-positive cells were also MAP2-positive 164 neurons for all iPSC lines (Figure 1C and Supplementary Figure 2C). As previously described (32) we found that these cultures expressed high levels of vGLUT2, FOXG1 and BRN2, that are 165

166 characteristic of excitatory layer 2/3 cortical neurons (Supplementary Figure 2D and
 167 Supplementary Table 2).

Using isoform-specific primers, we detected several KCNQ2 splice variants in the 168 169 differentiated neuronal cultures by RT-gPCR (Supplementary Figure 2E). To confirm the 170 presence of KCNQ2 protein, we performed Western blot analysis on proteins isolated from 171 neurons cultured for 4 weeks in vitro. We validated that we could detect both the wild type and 172 the T274M mutant KCNQ2 proteins expressed in transfected CHO cells (Supplementary Figure 2F). Unexpectedly, we found that Q2-01^{T274M/+} patient neurons exhibited significantly higher 173 174 KCNQ2 protein levels relative to their isogenic controls, suggesting a potential compensatory 175 mechanism (Figure 1D).

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177 KCNQ2-NEE Patient Neurons Exhibit Lower M-Current

Experimental evidence from Xenopus laevis oocytes indicated that pore mutations such 178 179 as T274M can produce a dominant negative loss of channel function (35). To determine the effect of the T274M KCNQ2 mutation in the context of human neurons, we measured the M-current in 180 181 week 4 excitatory patient-derived neurons. We recorded the total steady state current in wholecell voltage-clamp mode, and then blocked M-current by applying the selective KCNQ2/3 blocker 182 183 XE991. The remaining current was then subtracted from the baseline total current to determine 184 M-current density during the last 100 ms of 1 second voltage steps from -60 to +30 mV from a holding potential of -70 mV (Figure 1E). KCNQ2-NEE patient-derived neurons exhibited a 32% 185 186 reduction in total current density (p = 0.0005) and a 58% reduction in M-current density (p < 0.0005) 187 0.0001), as compared to neurons derived from two healthy control iPSC lines (Figure 1F-G). The 188 difference in total current density between controls and patient-derived neurons was equal to the 189 difference in M-current density, indicating the absence of compensatory outward conductances that would be active at the end of the 1-second voltage steps. Importantly, correcting the T274M 190 191 variant resulted in complete restoration of both total current density (p = 0.7282), as well as M-192 current density (p = 0.9446) to levels that were not significantly different from healthy control 193 neurons (Figure 1F-G).

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195 KCNQ2-NEE Neurons Exhibit Enhanced Spontaneous Neuronal Network Activity

The *KCNQ2* iPSC-based platform we developed presents an opportunity to dissect the functional consequences of chronic M-current impairment during neuronal development in culture. To assess spontaneous neuronal network activity, we used a multi-electrode array (MEA) platform and performed continuous recordings over a 3-week period (days 16-31, N = 4 independent differentiations, N = 40 well wide averages from 64 electrodes per well, per genotype). We plated an equal number of Q2-01^{T274M/+} and isogenic control neurons and carefully monitored neuronal attachment throughout the time course of experiments (Figure 2A and Supplementary Figure 3A). Neuronal cultures acquired significant spontaneous activity (>10% of the electrodes active) on or after day 19.

205 As illustrated by representative spike raster plots, patient-derived neurons exhibited an 206 enhanced spontaneous firing frequency relative to isogenic controls over time in culture (Figure 207 2B-C). As neurons matured, the spontaneous firing frequency increased in both cell lines, but the rate of increase was more rapid in Q2-01^{T274M/+} neurons (Figure 2C). Q2-01^{T274M/+} neuronal 208 209 cultures also had a significantly higher number of bursts, higher burst frequency, and greater 210 number of bursting electrodes (Supplementary Figures 2D and 3B). Furthermore, their spikes 211 were more restricted to bursts, as demonstrated by significantly higher number of spikes per 212 burst, percentage of all spikes occurring within bursts, and interspike interval covariance as 213 compared to isogenic controls (Figures 2C-D and Supplementary Figure 3B). There was no difference in number of active electrodes over time, although the rate at which more electrodes 214 became active over days in culture was higher in patient-derived neurons (Figure 2C). These 215 results indicate that Q2-01^{T274W/+} patient-derived neurons developed greater levels of 216 217 spontaneous activity and were more prone to fire in phasic bursts rather than single irregular 218 tonic spikes compared to isogenic control neurons (Figure 2E).

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220 KCNQ2-NEE Neurons Exhibit Progressive Enhanced Intrinsic Excitability

221 To determine if differences in firing behaviors exhibited by populations of Q2-01^{T274M/+} 222 neurons were driven by altered intrinsic excitability, we examined the firing frequency of large 223 numbers of neurons using the Optopatch, a recently developed system for high-throughput, all-224 optical electrophysiology with single cell resolution (36-38). The combined expression of CheRiff. 225 a blue light-activated channelrhodopsin, and QuasAr, a fluorescent voltage indicator allows for 226 the simultaneous stimulation and recording from multiple neurons within an elaborate network (Figure 3A). Using Synapsin 1 (SYN1)-driven expression constructs for CheRiff and QuasAr3 227 228 (Figure 3A), we imaged and analyzed the firing frequency of ~3000 neurons per genotype under 229 a blue-light illumination protocol (Figure 3B-D and Supplementary Figure 4). Day 35 neuronal 230 cultures were monitored for 2 sec without stimulation followed by five 500 ms pulses of blue light 231 of increasing intensity similar to a current injection step-protocol (Figure 3B). As shown in the 232 raster plot of spike timing for each neuron (Figure 3C, top), and the average firing rate of all 233 neurons (Figure 3C, bottom), KCNQ2-NEE patient-derived neurons exhibited a significantly higher firing frequency relative to their isogenic controls across the stimulation protocol (p < 0.01; Figure 3D). Within these populations of spiking neurons a significantly lower proportion of KCNQ2-NEE neurons responded to stimulations with a single spike (phasic spiking: p < 0.01; Figure 3E).

We next validated the intrinsic hyperexcitability phenotype in KCNQ2-NEE neurons by 238 239 performing whole-cell current-clamp measurements. We systematically recorded from single Q2-01^{T274M/+} patient-derived and isogenic control GFP-positive cortical neurons at three time points 240 241 in culture defined as week 3 (days 14-16), week 4 (days 22-26), and week 5 (days 32-35; Figure 242 4A-B). As neurons matured over time in culture, the neuronal resting membrane potentials (RMP) 243 became similarly more hyperpolarized for both genotypes, but input resistance was significantly higher (p = 0.0054) in Q2-01^{T274M/+} neurons only during week 3 (Supplementary Figure 5A and 244 245 Table 1). The increasing action potential amplitudes for both genotypes over time indicated neuronal maturation (Supplementary Figure 5B and Table 1). 246

To assess neuronal membrane excitability, we measured the frequency of action 247 potentials (APs) evoked by ascending somatic current injection steps (1 sec, 10 - 80pA) from a 248 holding potential of -65mV (Figure 4C). More than 80% of both Q2-01^{T274M/+} and isogenic control 249 neurons produced at least one AP using this protocol, and >50% exhibited repetitive trains of 250 251 APs by week 3 (Supplementary Figure 5C). Analysis of the number of APs evoked from iPSC-252 derived neurons during weeks 3, 4 and 5 revealed significant differences between genotypes during week 4 (p < 0.0001) and week 5 (p = 0.02; Figure 4D-E). Although Q2-01^{T274M/+} neurons 253 254 were able to fire significantly more APs than isogenic controls during week 4 and 5, there were 255 no differences between the genotypes during week 3 (p = 0.46; Figure 4E). Collectively these results suggest that Q2-01^{T274M/+} patient-derived excitatory neurons develop robust intrinsic 256 257 hyperexcitability progressively as they mature in culture.

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259 Lower M-current is Not Sufficient to Induce Hyperexcitability

The lack of intrinsic hyperexcitability of Q2-01^{T274M/+} neurons during week 3 prompted us 260 to investigate whether M-current was abnormal at this early time point in culture. We 261 262 hypothesized that KCNQ2 may not be expressed at this early time point, or that neurons were 263 able to compensate for lower KCNQ2 channel function with other KCNQ channels (i.e. KCNQ3 or KCNQ5) earlier on. To test this, we recorded total and M-current density in Q2-01^{T274M/+} and 264 265 isogenic control neurons during week 3. Similar to what we observed during week 4, both total current and M-current density in Q2-01^{T274M/+} neurons were significantly lower (31%, p = 0.009; 266 267 and 57%, p < 0.001, respectively) than isogenic neurons during week 3 (Supplementary Figure

5D). This suggests that lower M-current amplitude is not sufficient for hyperexcitability at this early time point. Interestingly, we also found that the total current and M-current density in both Q2-01^{T274M/+} and isogenic control neurons were significantly higher during week 3 than week 4 (p= 0.007, p = 0.02; Supplementary Figure 5E).

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273 KCNQ2-NEE Neurons Exhibit a Distal Shift of the AIS

274 Recent studies have shown that pharmacological inhibition of M-current induces 275 remarkable structural plasticity affecting neurite outgrowth and the dynamics of the axon initial 276 segment (AIS) (39-41). To determine whether such changes occur in the context of Q2-01^{T274M/+} 277 patient-derived neurons, we assessed morphology in week 4 GFP+ cortical neurons (Figure 5A). As KCNQ2 channels are known to be localized in the AIS, we also characterized the AIS structure 278 279 and position by immunolabelling neuronal cultures with Ankyrin G (ANK3), a marker of the AIS, together with the dendritic marker MAP2 (Figure 5A-B). To obtain a single measure for AIS 280 281 location, we determined the AIS distance as the linear path length from the base of the soma to the start of the dense expression of ANK3 and lack of MAP2 signal. While there was no 282 differences in the soma size, number of primary branches or the total length of the AIS, we found 283 284 that patient-derived neurons exhibited a distal shift of the AIS as compared to isogenic controls 285 (p = 0.0242; N = 41, 30 for control and patient neurons respectively; Figure 5B).

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287 KCNQ2-NEE Neurons Exhibit Enhanced AP Repolarization and Post-Burst AHP

The lower M-current in Q2-01^{T274M/+} neurons during weeks 3 and 4 (Figure 1 and 288 289 Supplementary Figure 5D) does not correlate with enhanced excitability, which occurs only during 290 and after week 4 (Figures 2 and 4). Furthermore, altered dendritic arborization and distally shifted 291 AIS suggests the activation of homeostatic plasticity mechanisms utilized by patient-derived neurons (Figure 5). Together these results suggest that an alternative intrinsic neuronal 292 293 mechanism may be responsible for patient-derived neuron hyperexcitability. To determine the source of the progressive increase in excitability in Q2-01^{T274M/+} neurons, we examined the AP 294 properties and post-burst AHPs at 3, 4 and 5 weeks of differentiation. We observed that Q2-295 $01^{T274M/+}$ neurons had significantly lower AP thresholds (p = 0.0001) and faster AP repolarization 296 297 with shorter AP half-widths (p < 0.0001), and larger fAHPs (p < 0.0001) (Supplementary Figure 298 6A-B; Table 1). Consistent with the excitability measurements (Figure 4), the significant 299 differences in AP thresholds and repolarization were limited to weeks 4 and 5 with no difference 300 during week 3. Lower AP thresholds and faster repolarization would allow neurons to fire more APs with less synaptic input, which may explain the enhanced excitability in Q2-01^{T274M/+} neurons. 301

302 These differences in AP properties may also explain the higher tendency of patient-derived 303 neurons to fire within bursts (Figure 2).

The ability of neurons to fire in bursts is limited by several conductances that turn on 304 following a burst of APs acting to hyperpolarize the membrane potential and prevent neurons 305 306 from firing more bursts. Previous studies have shown the involvement of KCNQ2 channels in the 307 medium afterhyperpolarization (mAHP) in cortical neurons (7,42). We examined the post-burst 308 AHP using a 50 Hz train of 25 APs evoked by 2 ms/1.4 nA current pulses. We found that Q2-01^{T274M/+} neurons exhibited a significantly enhanced mAHP (peak of AHP) and slow post-burst 309 AHP (sAHP; 1 second after last stimulus; p = 0.003 and p = 0.01, respectively; Figure 6C-D; 310 311 Table 1). These differences were only significant during weeks 4-5 but not earlier (Figure 6C-D; 312 Table 1). Although an enhanced AHP increases the refractory latency of neurons to repolarize 313 and be able to fire again after a burst of APs, lower AP thresholds and faster AP repolarization in Q2-01^{T274M/+} neurons may overcome this hurdle. These findings support our MEA data showing 314 315 enhanced burst firing in patient-derived neurons because the changes in AP properties would account for more APs within bursts while enhanced post-burst AHPs would dampen spontaneous 316 iee manuscr 317 firing between bursts (Figure 2).

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319 Acute Inhibition of M-current Reduces AP Repolarization and Post-Burst AHP

320 Previous studies have shown that blocking M-current in rodent cortical excitatory neurons acutely with XE991 enhances neuronal excitability by lowering AP threshold, impairing AP 321 repolarization and attenuating the post-burst AHP (39,43). However, our analysis of Q2-01^{T274M/+} 322 323 neurons showed that while they have lower M-current, they develop hyperexcitability through 324 progressive enhancement of repolarization with reduced AP half-width and enhanced fAHP and 325 post-burst AHP (Figures 1-4 and 6). We propose two potential explanations for these divergent 326 mechanisms for hyperexcitability: either M-current inhibition in human cortical glutamatergic 327 neurons has different effects than what was previously reported, or chronic M-current 328 suppression leads to hyperexcitability and altered AP properties by indirect mechanisms that are 329 different from those related to acute inhibition.

330 To test the first possibility, we interrogated the intrinsic membrane properties of week 4 331 control excitatory neurons before and after acute treatment with 20 µM XE991. Consistent with 332 previous reports (39,43), XE991 significantly reduced AP threshold and AP repolarization 333 (Supplementary Figure 6A-B and F). Furthermore, XE991 application blunted the mAHP and 334 sAHP amplitudes in these neurons (Supplementary Figure 6C-D and F).

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335 To determine the effect of acute M-current inhibition on neuronal excitability, we quantified 336 the number of evoked APs before and after XE991 application. Acute block of M-current enhanced the ability of isogenic control neurons to fire APs only during 20, 30 and 40 pA current 337 338 steps (p-value = 0.003, 0.048, 0.021 respectively; N = 25 cells; Supplementary Figure 6E). This did not resemble the behavior of Q2-01^{T274M/+} neurons, suggesting that loss of M-current alone is 339 340 not sufficient for hyperexcitability. Collectively, these experiments suggest that KCNQ2-NEE 341 patient neurons likely develop hyperexcitability progressively as a result of chronic M-current 342 reduction and dyshomeostatic adaptation of AP properties.

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344 Chronic Inhibition of M-current in Control Neurons Phenocopies KCNQ2-NEE Neurons

While changes in the AP properties of Q2-01^{T274M/+} patient-derived neurons may explain 345 346 why they are hyperexcitable during weeks 4 and 5, hyperexcitability does not correlate with loss of M-current during week 3. Furthermore, the intrinsic AP properties of Q2-01^{T274M/+} neurons are 347 348 not consistent with a pure loss of M-current but rather with a gain of another fast or Ca²⁺dependent voltage-gated K⁺ conductance that increases over time in culture. Therefore, we 349 hypothesized that early and chronic suppression of M-current leads to adaptive enhancement of 350 repolarization and post-burst AHP. To test this, we chronically treated isogenic control neurons 351 352 with XE991 (starting on day 12 in culture; Figure 7A) and measured excitability during week 4.

Chronically XE991-treated control neurons exhibited lower AP threshold (p = 0.003), enhanced repolarization (HW: p = 0.0002; fAHP: p = 0.01) and larger post-burst AHPs (mAHP: p = 0.0001; sAHP: p = 0.0008) relative to untreated controls (Figure 7B-E, Supplementary Figure 7A-D and Table 2). The treatment also resulted in a significant increase in the number of APs evoked by current injection steps as compared to untreated isogenic control neurons (p < 0.0001; Figure 7F). Importantly, these effects were identical to the properties of Q2-01^{T274M/+} neurons.

We next monitored spontaneous neuronal network activity using MEAs and found that chronic M-current inhibition was associated with enhanced burst firing parameters such as burst duration, number of spikes/burst and percentage of all spikes fired that occurred within bursts (p< 0.0001; Figure 7G and Supplementary Figure 7E-F). This experimental paradigm effectively phenocopied the electrophysiological behavior of Q2-01^{T274M/+} neurons in control cells and suggests that the AP repolarization and post-burst AHP alterations that we identified in patientderived neurons occur as a result of long term reduction of M-current.

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367 KCNQ2-NEE Neurons Exhibit a Dyshomeostatic Increase in BK Channel Function

The development of a shorter AP half-width and larger fAHP, mAHP and sAHP by patient neurons on and after week 4, suggest an adaptive upregulation of K⁺ conductance. While a number of different channels could be contributing to this dyshomeostatic mechanism, large conductance, Ca²⁺- and voltage-gated, fast-activating BK channels can participate in both the fast AP repolarization and slow Ca²⁺-dependent post-burst AHP (44). We therefore investigated the contribution of BK channels to the AP properties and post-burst AHP in mature patient and isogenic control iPSC-derived neurons. Addition of paxilline (20 µm), a BK channel antagonist, increased the AP threshold (Figure 8A) and reduced the fAHP after a single AP (Figure 8B) in patient neurons but not in controls. The magnitude of change of the AP threshold (Δ Threshold: p = 0.0049) and fAHP (Δ fAHP: p < 0.0001) was significantly larger for patient neurons compared with controls. Furthermore, paxilline treatment reduced the mAHP and sAHP of patient neurons to the level of controls with no effect in control neurons, making the magnitude of change larger for patient neurons (Δ mAHP: p = 0.0017; Δ sAHP: p = 0.002; Figure 8C). These data suggest that functional enhancement of BK channels in KCNQ2-NEE patient neurons contributes to the increase in fast-repolarizing currents and slowly-deactivating Ca²⁺-dependent currents. see manusci

402 **DISCUSSION**

403 We developed and validated a patient-derived iPSC-based model of KCNQ2-associated epileptic encephalopathy that provides novel insight into the mechanisms by which the 404 405 dysfunction of this channel leads to progressive neuronal abnormalities. Consistent with previous reports using heterologous expression systems (35), we found that the KCNQ2-encoded M-406 407 current density was significantly lower in patient-derived cortical excitatory neurons. This early 408 defect led to the progressive development of intrinsic and network hyperexcitability, as neurons 409 matured over time in culture. Unexpectedly, this hyperexcitability was associated with faster AP repolarization and enhanced AHP, properties not previously associated with loss of M-current. 410 411 Our findings suggest that KCNQ2 dysfunction induces dyshomeostatic plasticity and alters the 412 neurodevelopmental trajectory of patient neurons.

413 Neurons dynamically adjust the expression and functionality of ion channels as well as the structure of their processes to regulate their intrinsic excitability in response to cell autonomous 414 415 defects or their external environment. For example, exposed to chronic hyperexcitability, neurons 416 homeostatically downscale their intrinsic excitability and alter the size and location of the AIS (45-417 47). However, homeostatic plasticity can become maladaptive and even pathogenic in epilepsy when these processes become dysfunctional (46-52). The "acquired channelopathy" hypothesis 418 419 suggests that proepileptic channel characteristics develop during or after the onset of epilepsy or 420 excitotoxicity (53,54). However, these ideas are not mutually exclusive and likely chronic 421 hyperexcitability or epileptic activity leads to protective homeostasis as well as epileptogenic 422 dyshomeostatic neuronal adaptation.

423 We identified adaptive features of Q2-01^{T274M/+} excitatory neurons that may be protective 424 such as: a) increased KCNQ2 protein expression to potentially compensate for loss of KCNQ2 425 channel function, b) an enhanced post-burst AHP associated with longer inter-burst latency (55) 426 and c) a shift in the AIS away from the soma. A distal shift in the AIS has been suggested to lead 427 to an increase in the current thresholds for AP firing (45). At the same time we identified 428 progressively acquired intrinsic AP properties that lead to a hyperexcitable phenotype in Q2-01^{T274M/+} patient neurons despite the protective homeostatic changes. These dyshomeostatic 429 430 changes include a reduction in AP threshold and faster AP repolarization. The combination of the 431 hyperexcitable AP properties and the protectively increased post-burst AHP in single neurons 432 result in a neuronal network burst firing pattern that has been suggested to contribute to abnormal 433 interictal and ictal phenotypes on patient EEGs (29,56,57).

434 While it is not clear how reduced M-current in Q2-01^{T274M/+} patient-derived neurons leads 435 to the adaptive changes described here, we were able to recapitulate these defects by chronically blocking the M-channel in control neurons. Indeed, while acute suppression of M-current evokes
hyperexcitability by slowing AP repolarization and reducing the post-burst AHP, KCNQ2-NEE
patient neurons and chronically treated control neurons exhibit hyperexcitability that is associated
with faster repolarization and enhanced AHPs. These findings support a model where early
KCNQ2 channel dysfunction drives dyshomeostatic neuronal adaptation and hyperexcitability
through enhanced speed of AP repolarization and size of post-burst AHP.

AP repolarization and post-burst AHP are modulated by various K⁺ channel types, and it 442 443 is therefore likely that multiple conductances compound these alterations in KCNQ2-NEE patient neurons. Fast-activating and fast-inactivating K⁺ conductances are implicated in AP 444 445 repolarization while more slowly activating, slow-deactivating Ca²⁺-dependent K⁺ conductances are responsible for the medium and slow AHP (58). Our voltage-clamp recordings measuring M-446 447 currents, revealed that the difference in total current density between control and patient-derived neurons was equal to the difference in M-current density. This indicated the absence of 448 compensation for the loss of M-current by several slowly inactivating K⁺ conductances, allowing 449 us to eliminate these from the large list of potential culprits in the dyshomeostatically altered 450 451 intrinsic properties. r

The kinetics of fast activating BK (K_{ca}1.1; KCNMA1) channels are modulated by several 452 453 β subunits enabling them to be quickly inactivated if associated with $\beta 2$ (KCNMB2) or slowly 454 deactivated if associated with $\beta 4$ (*KCNMB4*) subunits (44,59-61). This property allows BK 455 channels to participate in both the fast AP repolarization and slow Ca²⁺-dependent post-burst 456 AHP. Our experiments with paxilline suggest that BK channels (K_{Ca}1.1; KCNMA1) contribute to the increased K⁺ conductance that enhances AP repolarization and AHPs, and leads to 457 458 hyperexcitability in KCNQ2-NEE patient-derived neurons. It is also possible that other K⁺ 459 channels implicated in AP repolarization and post-burst AHPs, such as A-type (Kv4.x/1.4; 460 KCND/KCNA4) and IK/SK (K_{ca}3.1/2.x; KCNN) channels, respectively, contribute to the changes 461 in intrinsic excitability of KCNQ2-NEE patient-derived neurons. Interestingly, several gain-of-462 function mutations in K⁺ channels have been reported in other genetic epilepsy disorders such as Kv7.5 (KCNQ5), BK (KCNMA1), Kv4.2 (KCND2), Kv2.1 (KCNB1), Kv1.2 (KCNA2) and Kca4.1 463 464 (KCNT1) channels (62).

Analysis of firing patterns from populations of cells using MEAs indicated that mutant *KCNQ2* patient-derived neurons developed greater levels of spontaneous activity and are more prone to fire in phasic bursts rather than single tonic spikes compared to controls. This is supported by the combination of the hyperexcitable AP properties and the protectively increased post-burst AHP in single neurons, which result in increased number of spikes per burst intermixed with longer refractory periods. This burst-suppression firing pattern is reminiscent of the interictal EEG pattern on KCNQ2-NEE patients (16,29,56,57). Repeated firing of discrete groups of neurons or bursts of high-frequency action potentials is typically observed in chronic epileptic conditions (63-66). Importantly, this type of activity does not necessarily reflect more neuronal APs, but rather an alteration in the neuronal discharge pattern (i.e. bursts rather than irregular single spikes). This can lead to hypersynchronous activity during seizure episodes and a burst-suppression pattern that is characteristic on KCNQ2-NEE patient EEGs during persisting periods without seizures (16,29,56,57).

Treatment with an M-current activator such as retigabine (ezogabine) has been shown to be most effective when given to patients early in development (29). Our findings support this clinical observation as early intervention may interrupt the dyshomeostatic adaptation observed in patient-derived neurons. Unfortunately, because KCNQ2-NEE presents in the first days of life, targeting the underlying cause (i.e. KCNQ2 loss-of-function) might not be effective if started late in the disease course. Perhaps, targeting these dyshomeostatically altered functional features might offer an alternative therapeutic strategy for this vulnerable patient population. The iPSC-based platform that we developed here may be used to identify effective therapeutics and address further questions regarding the spatiotemporal mechanisms of NEE due to KCNQ2 mutations.

508 METHODS

509

510 Cell Lines

511 Control 1 hiPSC line (11a; RRID:CVCL_8987) was derived previously (34). Control 2 512 hiPSC line (NCRM-5; NHCDR Cat# ND50031, RRID:CVCL_1E75) was obtained from the NIH 513 Center for Regenerative Medicine (NIH CRM). KCNQ2-NEE patient and isogenic control iPSC 514 lines were derived as described below. Further information on all iPSC lines can be found in 515 Supplementary Figure 2B.

516

517 Cell Culture

518 All iPSCs were grown on Matrigel (BD Biosciences) with mTeSR1 media (Stem Cell 519 Technologies) and passaged weekly using 1mM EDTA or Accutase (Sigma). All cell cultures 520 were maintained at 37°C and 5% CO₂. All lines were determined to be mycoplasma-free.

521 Primary glial cell cultures were derived from brain cortex of postnatal day 0-2, CD-1 mice (Charles River). Briefly, brain cortices were dissected free of meninges in dissection buffer HBSS 522 (Thermo Fisher), then digested with trypsin (Thermo Fisher) and DNAse I (Worthington) for 10 523 524 min at 37°C. The tissue was dissociated in glia medium: MEM (Life Technologies) supplemented 525 with Glutamax (0.6%), D-glucose, 10% normal horse serum (Life Technologies), and penicillin-526 streptomycin (Thermo Fisher). After centrifugation and resuspension, cells were filtered through 527 a 0.45 micron cell strainer and plated on poly-D-lysine coated plates with glia media at 37°C, 5% 528 CO₂ for 2 weeks. Afterwards, glial cultures were tested for mycoplasma, dissociated for 529 expansion, and frozen in 10% DMSO/horse serum. All animal experiments were approved and 530 conducted in accordance with the policies and guidelines set forth by the Northwestern University 531 Institutional Animal Care and Use Committee (IACUC).

532

533 Generation of iPSCs

Peripheral blood mononuclear cells (PBMCs) were isolated from a whole cell blood draw 534 following informed consent under protocols approved both by Ann & Robert H. Lurie Children's 535 536 Hospital of Chicago and Northwestern University. Reprogramming of PBMCs into iPSCs was 537 performed at the Northwestern Stem Cell Core Facility using Invitrogen's CytoTune®-iPS 2.0 Sendai Reprogramming system (A16517, Thermofisher), following the manufacturer's 538 539 instructions. This reprogramming system uses four transcription factors (Oct4, Sox2, Klf4, c-Myc). Briefly, PBMCs (5 × 10⁵) were seeded into one well of a 24-well plate and cultured for four 540 541 days in StemSpan[™] SFEM II PBMC complete medium (STEMCELL Technology, 09655) 542 supplemented with 100 ng/ml SCF (PeproTech, 300–07), 100 ng/ml FLT3 (PeproTech, 300–19),

20 ng/ml IL-3 (PeproTech, 200–03) and 20 ng/ml IL-6 (PeproTech, 200–06). Immediately after 543 544 plating, the cells were infected with Sendai virus for 48 h at 37 °C. The infected cells were transferred onto MEF feeders and cultured in StemSpan[™] SFEM II. Following 21–28 days of 545 546 culture, individual iPS colonies were picked and transferred to matrigel coated 6 well plate for 547 expansion and were maintained in mTeSR1.

548

549 **CRISPR/Cas9** Gene-Editing

Isogenic control iPSCs were generated using CRISPR/Cas9 from the Q2-01^{T274M/+} patient-550 551 derived iPSC line in collaboration with Applied StemCell (Milpitas, CA). Briefly, iPSCs were co-552 transfected with a plasmid encoding Cas9 nuclease along with the single guide RNA (sgRNA) KCNQ2.g2 (Supplementary Table 1), a puromycin-resistance plasmid, and a donor single 553 554 stranded repair oligonucleotide (Supplementary Table 1). Cells were grown in the presence of 555 puromycin for two days then colonies were isolated and expanded two weeks later. Clones with 556 the desired genetic modification were identified by PCR genotyping and confirmed by DNA sequencing (Supplementary Figure 1C). All primer sequences can be found in Supplementary 557 see manusci Table 1. 558

559

Analysis of Off-Target Cas9 Sites 560

561 Potential off-target sites were identified with the online tool: http://crispr.mit.edu. We selected the top 8 genomic regions of homology and thus most likely off-target sites, and 562 563 amplified each one by targeted PCR of genomic DNA from the corrected iPSC clone, for further analysis either by Sanger Sequencing or by a T7 Endonuclease assay (Supplementary Figure 564 565 1D-E; Supplementary Table 1). The same PCR conditions were used to amplify the positive 566 control DNA template and primer mix, included in the Genecopoeia TM T7 Endonuclease I Assay Kit. The amplified DNA from each potential off-target site was purified using the Wizard ® SV Gel 567 568 and PCR Clean-Up System (Promega). The concentration of the purified DNA from the potential off-target sites and the template DNA from the positive control was assessed by using a 569 Nanodrop 2000 Spectrophotometer (Thermo-Fisher). 500 ng of DNA product from each potential 570 571 off-target region of the isogenic and parental cell lines and 500 ng of the positive control DNA 572 template were heated to 95°C for five minutes and subsequently allowed to cool to room 573 temperature to denature and re-anneal the PCR products. 2U of T7 Endonuclease I 574 (Genecopoeia) was added to the re-annealed PCR products and incubated at 37°C for 60 575 minutes. The PCR products from the potential off-target sites and the positive control template were then run on a gel with 6x loading buffer, alongside a 2-log DNA ladder (New England
BioLabs Inc.). All primers sequence can be in Supplementary Table 1.

578

579 Cortical Excitatory Neuron Differentiation

iPSCs were differentiated into glutamatergic neurons using a modified version of a 580 581 protocol based on Ngn2 overexpression (32). Stem cells were dissociated as single cells using 582 Accutase (Sigma), re-suspended in mTeSR1 with 10µM ROCK inhibitor (Y-27632, DNSK 583 International), then incubated with lentiviruses (pLV-hPGK-M2rtTA, TetO-Ngn2-Puro, TetO-584 FUW-EGFP) in suspension for 5-10 min before plating (95,000 cells/cm²; Supplementary Figure 585 2A). After 24h (day 1), lentivirus was removed and replaced with knockout serum replacement medium (KOSR) consisting of KnockOut DMEM supplemented with Knockout replacement 586 serum KSR, nonessential amino acids (NEAA), Glutamax (Life Technologies), 55μM β-587 mercaptoethanol (Gibco, Cat# 21985023), 10µM SB431542 (DNSK International), 100nM LDN-588 589 193189 (DNSK International), 2µM XAV939 (DNSK International) and 2 µg/ml of doxycycline (Sigma). On the following day (Day 2), media was replaced with a 1:1 ratio of KOSR to neural 590 591 induction media (NIM) composed of DMEM:F12 supplemented with NEAA, Glutamax, N2 (Gibco, Life Technologies), 0.16% D-glucose (Sigma) and 2µg/ml heparin sulfate (Sigma). Doxycycline 592 593 (2µg/ml) and puromycin (2µg/ml; Sigma) were added to this NIM media. On Day 3, the media 594 was replaced with NIM containing doxycycline (2µg/ml) and puromycin (2µg/ml). All neurons were 595 frozen in 10% DMSO/FBS on Day 4 (Supplementary Figure 2). For all experimental analysis, 596 iPSC-derived neurons were plated on primary CD1 mouse cortical glia, derived as previously 597 described (67). Glial cells were first plated on PDL/laminin-coated plates or coverslips in glia 598 media composed of MEM (Life Technologies) supplemented with Glutamax (0.6%), D-glucose, 599 and 10% Horse serum (Life Technologies). After 5-7 days, neurons were thawed (Day 5 post-600 induction) and plated, at a density of 20.000/cm², directly onto the monolayer of mouse glia in Neurobasal medium (NBM), supplemented with NEAA, Glutamax, N2 and B27 (Life 601 602 Technologies) containing BDNF (10ng/mL, R&D systems), 2% FBS (VWR), doxycycline and 603 ROCK inhibitor. Half of the media was replaced every other day.

604

605 **Preparation of Plasmids and Lentivirus**

606 Overexpression of KCNQ2 in CHO cells for Western blotting was achieved using a plasmid 607 containing wild type KCNQ2-IRES2-EGFP that was a gift from David McKinnon (68). Two base 608 pairs were modified through mutagenesis, in this plasmid (c.2002G>A, p.K668E; c.2467T>C, p.R823C), to match NM_172108.4 sequence. Mutagenesis was performed to insert the T274M
 patient mutation into the wildtype construct.

TetO-Ngn2-puro (Addgene plasmid #52047) and TetO-FUW-EGFP (Addgene plasmid #30130) plasmids were gifts from Marius Wernig (32,69). FUW-M2rtTA was a gift from Rudolf Jaenisch (Addgene plasmid # 20342) (70). Lentiviruses were generated in HEK293T cells using the second generation packaging vectors, psPAX2 and pMD2.G, as described previously (71) by the Northwestern University DNA/RNA Delivery Core.

616

617 Immunocytochemistry

618 iPSCs and neurons were plated on Matrigel or PDL/laminin-coated glass coverslips, respectively. Cells were fixed with 4% formaldehyde (Sigma) in 4% sucrose/PBS for 15 min at 619 620 room temperature. Cells were permeabilized and blocked simultaneously in PBS containing 0.1% Triton and 5% normal goat serum (Jackson Immunoresearch) for 2 hr at 4°C followed by 621 incubation with primary antibodies overnight at 4°C. The following primary antibodies were used: 622 GFP (Abcam ab13970, RRID: AB 300798, 1:10,000), ankyrin-G (Neuromab 75-146, 623 AB 10673030, 1:200), synapsin (Cell signaling #5297, AB 2616578, 1:200), Map2 (Millipore 624 MAB3418, AB 94856, 1:1000), and Map2 (AB15452, AB 11211337, 1:1000). Cells were 625 626 washed three times in PBS and incubated with secondary antibodies at room temperature for 1 627 hour followed by another three washes in PBS. The following secondary antibodies were used: Alexa 488 goat anti-chicken, Alexa 568 goat anti-mouse, Alexa 568 goat anti-chicken, Alexa 647 628 629 goat anti-rabbit, and Alexa 647 goat anti-mouse (Thermo Fisher Scientific, 1:1000). Primary and 630 secondary antibodies were diluted in PBS containing 5% normal goat serum. Coverslips with 631 immunostained cells were washed briefly in distilled water and mounted onto microscope slides 632 using Prolong anti-fade reagent (Life Technologies). Neurons from the same differentiation 633 experiment were fixed and stained at the same time with identical antibody dilutions. Images 634 were acquired at 10x for MAP2/GFP+ neuron counting using a Leica inverted Ti microscope.

635

636 Confocal Imaging and Analysis of Neuronal Morphology

For morphometric analysis images of immunostained neurons were acquired using a ×63 oil immersion objective lens on a Nikon C2+ confocal microscope with NIS-Elements software. Each channel was acquired sequentially to prevent fluorescence bleed-through with 2x line averaging for each channel. Neurons were imaged on multiple z-planes to capture full dendritic and axonal projections, and maximum intensity projections (max projections) of z-stacks created in Image J (Fiji) were used for downstream analyses. Images were acquired using identical
 settings for each experiment to allow comparison between experimental groups.

For analysis of the AIS and soma area, thirty to forty neurons were imaged for each 644 645 isogenic line (10-15 neurons from each of three independent differentiations). Regions of interest (ROI) defining the outline soma were drawn in Image J from max projections of the GFP channel. 646 647 Z-stacks were consulted for cells with ambiguous somas shapes, to avoid the inclusion of any protrusions. To identify the AIS, a set threshold (>1000 mean pixel intensity) was applied to 648 649 confocal images containing Ankyrin G (ANK3) immunostaining. Any signal above this threshold and devoid of MAP2 immunostaining was defined as the AIS. The AIS was traced from start to 650 651 finish using the freehand line tool in Image J to determine the total AIS length. If a neuron contained multiple regions of ANK3 staining (i.e. multiple axons) the length of all ANK3+ 652 653 segments was summed for that cell. To determine the distance of the AIS from the soma, the shortest possible line was traced from the edge of the soma ROI to the start of the AIS, following 654 655 neuronal processes. Morphometric measurements [soma area (µm²), AIS length (µm), the distance of the AIS to the soma (µm)] were all performed in Image J and recorded in Microsoft 656 Excel. Statistical significance between two experimental groups was determined using a Student 657 t-test (for parametric data) or Mann-Whitney (for non-parametric data). All graphs and statistics 658 659 for morphometric analyses were performed in Statview.

660

661 **RNA Isolation and qRT-PCR**

Cells were harvested by scraping from 6-well plates at the indicated time points after 662 induction of neuronal differentiation. Cells were resuspended in TRIzol Reagent (Life 663 664 Technologies), and RNA was isolated following manufacturer's protocol. RNA (0.5 to 1 µg) was 665 treated with DNase I (Invitrogen) and subsequently used for the generation of cDNA using iSCRIPT Reverse Transcription Supermix (Bio-Rad) following manufacturer's instructions. RT-666 667 PCR was performed using SYBR green on the CFX system (Bio-Rad). All assays were performed in triplicate. The averaged cycle of threshold (Ct) value of two housekeeping genes (GPI/GAPDH) 668 was subtracted from the Ct value of the gene of interest to obtain the Δ Ct. Relative gene 669 expression was determined as $2^{-\Delta Ct}$ ($\Delta \Delta Ct$) and expressed relative to the control sample or the 670 671 highest expressed sample in the experiment. All primer sequences are listed in Supplementary 672 Table 2.

- 673
- 674 Western Blot

675 Cells were harvested from 6-well plates at the relevant times of neuronal differentiation. Briefly, cells were twice washed with ice-cold PBS, and scraped from the wells in PBS with mini 676 EDTA-free protease inhibitor cocktail table (Roche). Cell pellets were lysed in EBC Lysis Buffer 677 678 (Boston Bio Products) with protease inhibitor cocktail on ice for one hour and spun at 14,000 679 RPM in an Eppendorf 5417R centrifuge at 4°C for 10 minutes. The supernatant was used to 680 perform a BCA assay. Protein samples were electrophoresed on NovexTM 4-20% Tris-Glycine 681 gels (Thermo-Fisher) and transferred onto nitrocellulose membrane using the CriterionTM Blotter 682 transfer system (Bio-Rad). Membranes were blocked in 5% milk in .1% Tween/Tris-buffered saline (TBST) for 30 min and probed with primary antibodies in 5% BSA-TBST with .05% sodium 683 684 azide against: KCNQ2 (1:1000; custom made), GAPDH (Millipore MAB374; 1:5000) and βTubulin-III (BioLegend #801201, previously Covance: #MRB-435P; 1:5000) overnight at 4°C, 685 686 followed by Peroxidase-AffiniPure Goat Anti-Mouse or Anti-Rabbit IgG (H+L) secondary antibody (Jackson ImmunoResearch 115-035-003 or 111-035-144, respectively; 1:10000) in 5% milk-687 688 TBST for 1hr. To visualize protein, blots were incubated with secondary antibody in 5% milk-TBST in the dark for 1 hr. Finally, blots were visualized using SuperSignal West Pico PLUS 689 chemiluminescence (ThermoScientific) on a Bio-Rad Chemidoc using ImageLab software. 690

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PP 692 **Multi-Electrode Array Recordings**

693 For multielectrode array (MEA) studies, 12 well MEA plates with 64 electrodes per well 694 were coated with PDL and laminin according to Axion Biosystems protocols. Mouse glial cells 695 were seeded at a density of 40.000 cells/well then 30.000 neurons/well were added one week 696 later. Every other day, half of the media was removed from each well and replaced with fresh 697 media. Spontaneous activity was recorded using Axion Biosystems Maestro 768 channel 698 amplifier and Axion Integrated Studios (AxIS) v2.4 software. The amplifier recorded from all channels simultaneously using a gain of 1200× and a sampling rate of 12.5 kHz/channel. After 699 700 passing the signal through a Butterworth band-pass filter (300–5000 Hz), on-line spike detection 701 (threshold = 6× the root-mean-square of noise on each channel) was performed with the AxIS adaptive spike detector. All recordings were conducted at 37°C in 5% CO₂ / 95% O₂. 702 703 Spontaneous network activity was recorded for 5 min each day starting on day 15 of 704 differentiation. Active electrodes were defined as having ≥ 1 spikes/min. The mean firing rate (Hz). 705 burst number, and number of spikes per burst were used as a measure of neuronal activity as 706 this demonstrates maturity of neuronal functional properties. MEA recordings were done with 4 707 independent differentiations. All data reflects well-wide averages, with the number of wells per 708 condition represented by N values.

709

710 **Optopatch Measurements**

711 Cryo-stocks of Ngn2 neurons (preserved on day 4) differentiated from KCNQ2-NEE 712 patient and isogenic control iPSC lines were thawed and plated at a density of 100,000 cells/cm² 713 onto Poly-D-Lysine/Laminin pre-coated 96-well custom-made Ibidi® tissue culture dishes. Cells 714 were co-plated with P1 primary C57BL/6 mouse cortical glial cells, at a density of 30,000 715 cells/cm². Primary glial cells were prepared as previously described (67). These co-cultures were 716 maintained in complete NBM (NBM medium supplemented with 1x Gibco® N2, 1x Gibco® B27, 717 10ng/mL BDNF (R&D), 2ug/mL Doxycycline (Sigma) and 2% Hyclone[™] Fetal Bovine Serum). Cells were cultured for 30 days with complete medium exchanges every 3 days. Two weeks prior 718 719 to all-optical (Optopatch) electrophysiology measurements (at Day 21), cells were transduced with lentiviral particles encoding Optopatch components CheRiff-BFP and QuasAr3-Citrine 720 (36,72), driven by the human Synapsin I promoter for neuronal-specific expression. Lentiviral 721 preparation and transduction were carried out as previously reported (38). Cells were recorded 722 using Optopatch imaging (36-38) on day 35 (Figure 3A). Cells were imaged in Tyrodes Buffer 723 (10mM HEPES, 125mM NaCl, 2mM KCl, 3mM CaCl₂, 1mM MgSO₄, 30mM Glucose, pH 7.40) in 724 725 the presence of synaptic blockers (10µM NBQX (Sigma), 25µM D-AP5 (Tocris) and 20µM 726 Gabazine (Sigma) to block AMPA, NMDA and GABA currents, respectively) to allow for 727 measurements of intrinsic spontaneous and evoked neuronal activity. Optopatch imaging was 728 performed on a custom built, ultra-wide field fluorescence microscope described previously 729 (38.73). Briefly, samples were illuminated with ~100 W/cm² 635 nm laser excitation to monitor 730 changes in membrane potential through changes in QuasAr3 fluorescence. In order to evoke 731 neuronal activity, a custom blue light (470nm LED) stimulus protocol was used to depolarize the 732 cell membrane through excitation of CheRiff. The stimulus protocol consisted of 2 seconds of 733 spontaneous activity (blue light off) followed by five 500 msec steps of increasing blue light 734 intensity (2.45, 5.5, 14.67, 33, and 88 mW/cm²), with 500 msec of rest in between each stimulus 735 step. Imaging data were recorded on a Hamamatsu ORCA-Flash 4.0 sCMOS camera across a 736 4 mm x 0.5 mm field of view at a 1 kHz frame rate. Data acquisition was performed using custom 737 control software written in MATLAB. Analysis of Optopatch data, using a custom analytics 738 pipeline written primarily in MATLAB, was carried out as previously described⁴. Statistical 739 significance/p-values were determined using the Kolmogorov-Smirnov statistic with a custom MATLAB routine (** or *** indicates p<0.01 and p<0.005, respectively). All reported error bars 740 741 are standard error of the mean.

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743 Patch Clamp Electrophysiology

744 Whole-cell voltage and current clamp recordings were made from visually identified GFP-745 expressing neurons using an inverted Olympus IX51 microscope equipped with a 40X objective. Recording pipettes were made of glass capillaries using a horizontal Sutter P-1000 puller yielding 746 a 3-5M Ω resistance pipette when filled with standard intracellular solution containing (in mM): 747 748 120 K-MeSO₄, 10 KCl, 10 HEPES, 10 Na₂-phosphocreatine, 4 Mg-ATP, 0.4 Na₃-GTP, pH 7.3 749 adjusted with KOH; osmolality 285-290 mOsm/Kg. Neurons were continuously perfused with oxygenated aCSF bath solution (in mM): 125 NaCl, 26 NaHCO₃, 2.5 KCl, 1.25 NaH₂PO₄, 1 750 MgSO₄, 25 glucose, 2 CaCl₂, pH 7.4 at 33-35°C. Whole-cell voltage clamp recordings were 751 752 acquired using an Axopatch 200B amplifier (Molecular Devices, USA). An M-current blocker, XE991 (20µM, Abcam), was applied 2-3 minutes after establishing the whole cell configuration, 753 754 and XE991-insensitive currents recorded 10 minutes after application were subtracted offline. Mcurrents were measured in the final 100ms of the voltage steps (Figure 1F-G). 755

756 Current-clamp recordings were acquired using a Multiclamp 700B amplifier (Molecular Devices, USA) and digitized at 10kHz (filtered at 3kHz) with the neurons held at -65mV (V_h). All 757 758 reported potential values were corrected for the liquid junction potential, calculated to be -8.4 mV. Resting membrane potential was measured immediately after establishing the whole-cell 759 760 patch clamp configuration. Input resistance (R_N) was calculated as the slope of the voltagecurrent curve determined using 500 ms current steps from -50 pA to 30 pA in 10 pA steps. 761 762 Medium (mAHP) and slow (sAHP) afterhyperpolarizations (AHPs) were measured as the 763 difference between V_h and the negative going peak and 1 sec after the offset of the last current 764 step, respectively, induced by a 50 Hz train of 25 APs evoked by 2 ms/1.2 nA current injection 765 pulses. Single AP properties, including fast afterhyperpolarization (fAHP), were measured using 766 direct somatic current injection ramps (10-80 pA, 500 ms). AP amplitude was calculated as the difference between V_h to the peak of the first AP of the ramp protocol. AP threshold was 767 768 calculated where the first derivative of the up phase of the trace equaled 5mV/ms. Using a 1 ms 769 sliding average, the fAHP measurement was taken when the mean first derivative of the trace reached 0.0 ± 0.5 after initial spike in each sweep. AP width measurements were taken at half 770 771 the AP peak amplitude relative to V_h . Accommodation was measured during 1000 ms current 772 steps from 10 to 80 pA; the total number of APs during the entire steps were counted. Neurons 773 meeting the following criteria were used: series resistance (Rs) < 30 M Ω , membrane resistance (R_N) > 200 M Ω , resting potential (V_{rest}) < -45 mV, and AP amplitude > 80 mV from V_h. Data were 774 775 analyzed using custom MATLAB protocols. Statistical analyses were conducted using Statview.

We also tested the action of chronic and acute application of 1 and 20 µM XE991 (expected to block 50% and 100% of M-current, respectively) (Wang et al., 1998). 1 µM XE991 was chronically applied to neuronal culture media starting day 12 in differentiation (right before beginning of week 3 time point) and AP properties were measured on week 4. Acute application of 20 µM XE991 or 20 µM paxilline (Tocris) was done during week 4 and AP properties were measured before and 10 min after continuous perfusion of aCSF with XE991 or paxilline.

Drugs

Drugs were prepared as a stock solutions using distilled water or DMSO, then diluted to the required concentration in aCSF or culture media immediately before use. Bath-applied drugs were perfused for at least 10 min to ensure complete equilibration within the recording chamber OI for details before recording.

Statistical Analysis

Differences were evaluated using t-test, one-way or two-way ANOVA, repeated-measures ANOVA, and Fisher's protected least significant difference post hoc tests where appropriate. All data are reported as means ± SEM.

Study Approval

Written informed consent was received from participants prior to inclusion in the study under protocols approved both by Ann & Robert H. Lurie Children's Hospital of Chicago and Northwestern University IRB (#2015-738).

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820 AUTHOR CONTRIBUTIONS

D.S., A.L.G., and E.K. conceived the study. D.S. designed and performed experiments, 821 analyzed and interpreted all the data presented, assembled figures and wrote the manuscript. 822 B.N.P. and D.S. generated and maintained all iPSC and neuronal cultures. B.N.P. and G.L.R. 823 performed quality control and qPCR experiments. T.J.S. and D.S. performed patch-clamp 824 825 electrophysiological experiments. D.S. performed MEA experiments. M.F. and B.N.P. performed and analyzed immunocytochemistry and confocal imaging experiments under the supervision of 826 827 P.P. M.S. performed Western blot experiments. J.J.M. and L.C.L. recruited patients for the study. 828 L.A.W., V.J., H.Z., S.J.R., O.B.M. and G.T.D. performed and analyzed data from Optopatch experiments. A.L.G. and E.K. designed experiments, interpreted the data, provided guidance and 829 supervision and wrote the manuscript. All authors helped to edit the manuscript. 830

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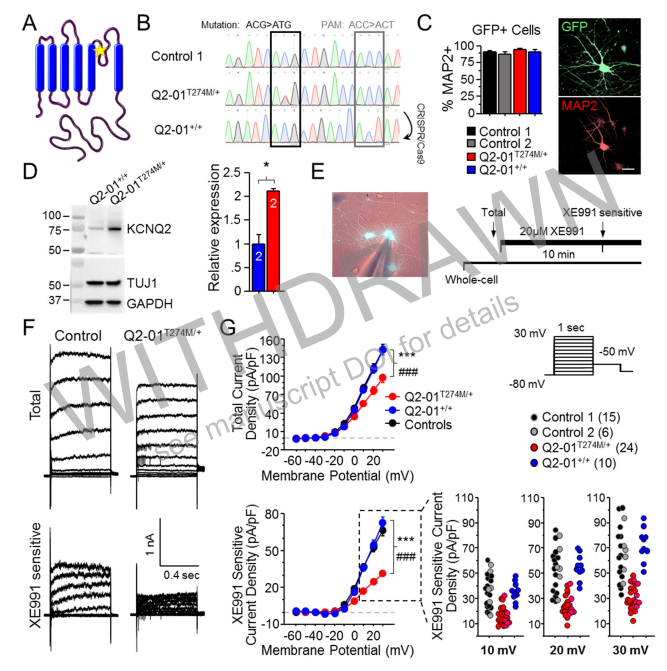
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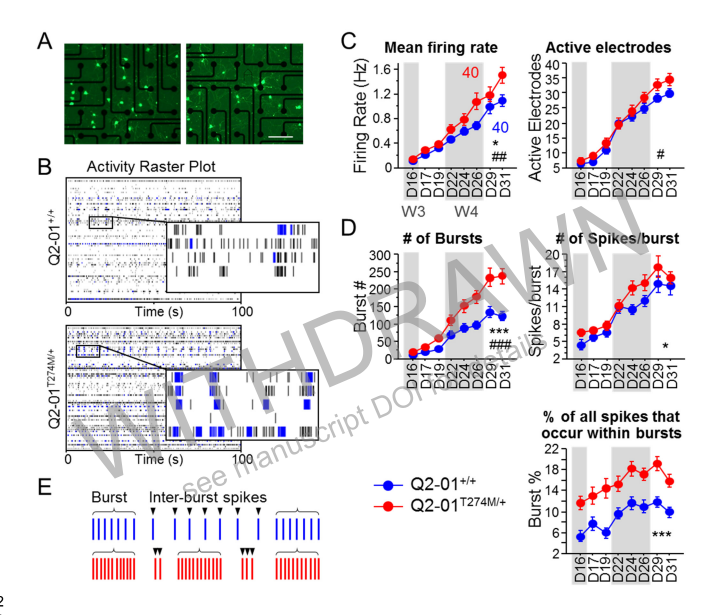




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1004 Figure 1. KCNQ2-NEE patient-specific iPSC-derived neurons exhibit reduced M-currents. (A) Proposed structure of KCNQ2 channel subunit containing the patient mutation T274M (yellow 1005 1006 star) near the ion selectivity filter within the pore-loop domain. (B) DNA sequence electropherograms of KCNQ2 in control and patient iPSCs before and after gene editing, 1007 demonstrate the correction of the heterozygous (T274M; c.821C>T) mutation. A silent mutation 1008 1009 (T276T: ACC>ACT) was concurrently introduced in the PAM site (See Supplementary Figure 1 and Supplementary Tables 1-2). (C) Quantification of GFP fluorescence coincident with MAP2 1010 1011 immunopositive staining in two unrelated healthy controls and patient and isogenic control iPSC-1012 derived neurons (See Supplementary Figure 2). Scale bar: 20µm. (D) Western blot analysis of KCNQ2 protein in patient-derived neurons relative to isogenic controls (t-test: *p=0.02; N=2 1013 independent differentiations). TUJ1 quantification indicates that cell lysates contained similar 1014 1015 levels of mature neurons. GAPDH was used as a loading control. (E) Left: Representative image of GFP-fluorescing neuron during patch-clamp recording. Right: Experimental protocol. Total 1016

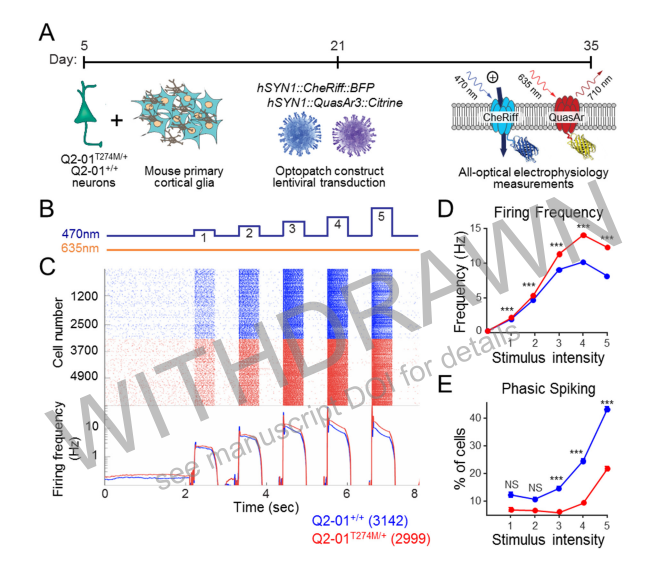
steady-state current density and M-current density were measured in voltage-clamp mode after establishing whole-cell configuration. (F) Representative traces recorded from control and patient-derived neurons before XE991 application (top; total current) and after subtraction of XE991-insensitive currents (bottom; XE991 sensitive/M-current). (G) Total and XE991-sensitive current density plotted against test potential. Top, repeated measures ANOVA for total current density during week 4 by cell line: F_(2,468)=8.93, ****p*=0.0005; cell line/voltage step interaction: F(18.468)=11.11, ###p<0.0001; Fisher's PLSD posthoc comparing healthy controls (black) to isogenic control (blue) neurons (p=0.7282) and Q2-01^{T274M/+} (red) to either isogenic control (p=0.0071) or healthy control neurons (p=0.0002). Bottom, M-current density repeated measures ANOVA: F_(2,468)=29.83, ***p<0.0001; cell line/voltage step interaction: F_(18,468)=33.36, ###p<0.0001; Fisher's PLSD posthoc test comparing healthy controls to isogenic controls (p=0.9446) and Q2-01^{T274M/+} neurons to either isogenic controls or healthy controls (p<0.0001; N=21 healthy controls combined). Bottom right: plot of M-current densities from all neurons recorded during 10, 20 and 30mV steps. Number of neurons analyzed is displayed within the figure. Values displayed as mean±SEM. anuscript DOI for details



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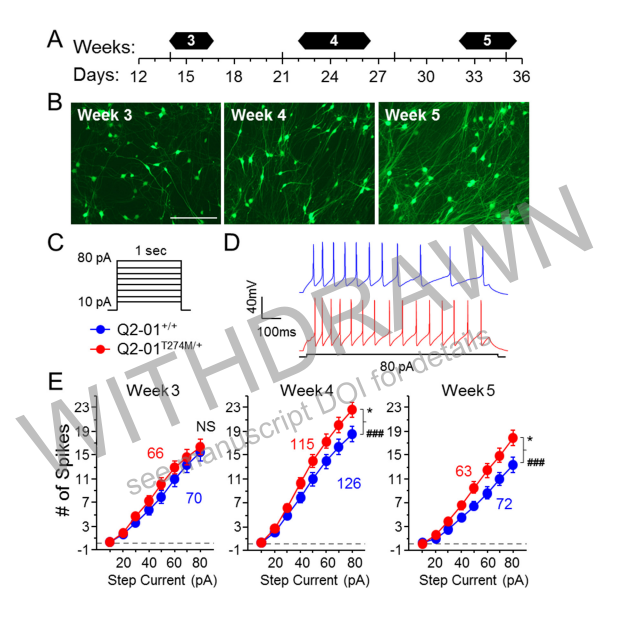
Figure 2. KCNQ2-NEE neurons exhibit enhanced spontaneous neuronal network activity. 1064 (A) Representative images of GFP-fluorescing patient-derived (Right) and isogenic control 1065 neurons (Left) plated on MEA wells at 31 days in culture. Scale bar: 200µm (B) Representative 1066 1067 raster plots from a single MEA well of patient-derived and isogenic control neurons on day 31. Each row represents the signal detected by a single electrode, black ticks indicate single spikes 1068 and blue ticks spikes that occur within bursts. (C) Longitudinal analysis of the mean firing 1069 1070 frequency (repeated measures ANOVA for genotype: $F_{(1,546)} = 5.63$, *p = 0.02; genotype/day interaction: $F_{(7,546)} = 3.17$, ^{##}p = 0.0027); number of active electrodes (repeated measures) 1071 ANOVA for genotype: p = 0.198; genotype/day interaction: $F_{(7,546)} = 2.49$, # p = 0.0159). (D) 1072 1073 Number of bursts detected (repeated measures ANOVA for genotype: $F_{(1,546)}$ = 18.82, ***p < 0.001; genotype/day interaction: $F_{(7,546)} = 6.71$, $^{\#\#}p < 0.0001$); number of spikes within bursts 1074 (repeated measures ANOVA for genotype: $F_{(1,546)} = 4.17$, *p = 0.0445); and percentage of spikes 1075 1076 which were found to occur within bursts (repeated measures ANOVA for genotype: $F_{(1.546)}$ = 31.62, ***p < 0.0001; See Supplementary Figure 3). (E) Illustration of firing pattern showing 1077 increased phasic firing in bursts of patient neurons (red) as compared to isogenic controls (blue). 1078 Number of wells analyzed per cell line is displayed within the figure from 4 independent 1079 1080 differentiations. Values displayed are mean ± SEM. 1081



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1084 Figure 3. KCNQ2-NEE neurons are hyperexcitable relative to their isogenic controls as 1085 measured by Optopatch recordings. (A) Experimental outline: neurons were co-cultured with primary glial cells and on day 21 were infected with lentiviruses encoding the Optopatch 1086 constructs. CheRiff was coupled to a BFP fluorescence reporter and QuasAr3 to Citrine. 1087 1088 Optopatch imaging was performed on day 35. (B) Illumination protocol: following 2 sec of 1089 spontaneous activity recording, neurons were subjected to five, 500 ms stimulation pulses of increasing blue light intensity (1: 2.45, 2: 5.5, 3: 14.67, 4: 33, and 5: 88 mW/cm²) at 1Hz, to 1090 activate CheRiff at 470 nm. Changes in membrane potential were monitored through changes in 1091 QuasAr3 fluorescence excited by red light at 635 nm. (C) Top: raster plot showing spike timing 1092 1093 for individual neurons across the stimulation protocol. Each row represents an individual neuron 1094 and each point an AP. Bottom: average firing frequency of mutant and control neurons across the stimulation protocol. (D-E) Average firing frequency and percentage of cells that exhibit 1095 1096 phasic spiking for each stimulus intensity step. Number of neurons analyzed is displayed within 1097 the figure. Values displayed are mean ± SEM. Statistical significance determined by using the Kolmogorov-Smirnov statistic, **p<0.01 and ***p<0.005, NS: not significant. 1098 1099



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1101 Figure 4. KCNQ2-NEE neurons exhibit progressive enhancement of intrinsic excitability. 1102 1103 (A) Experimental time line. (B) Representative images of GFP-fluorescing isogenic control neurons during week 3, 4 and 5. Scale bare: 500µm. (C) Current-clamp protocol. Intrinsic 1104 1105 excitability was measured in current-clamp mode using 1 sec somatic current injection steps (10-1106 80pA) from a holding potential of -65mV. (D) Representative traces of patient-derived (red) and 1107 isogenic control (blue) neurons firing APs during an 80 pA current stimulus. (E) Numbers of APs that patient-derived and isogenic control neurons fired during week 3 (repeated measures 1108 1109 ANOVA for genotype: p = 0.21; genotype/current injection amplitude interaction for week 3: p =0.38), week 4 (repeated measures ANOVA for genotype: $F_{(1,1673)} = 5.14$, *p = 0.015; 1110 genotype/current injection amplitude interaction: $F_{(7,1673)} = 4.79$, ###p < 0.0001) and week 5 1111 1112 (repeated measures ANOVA for genotype: $F_{(1,931)} = 5.68$, **p* = 0.018; genotype/current injection amplitude interaction: $F_{(7,931)} = 5.27$, $^{\#\#}p < 0.0001$). NS: not significant. Number of neurons 1113 analyzed is displayed within the figure and in Table 1 (See Supplementary Figure 5). Values 1114 1115 displayed are mean ± SEM.

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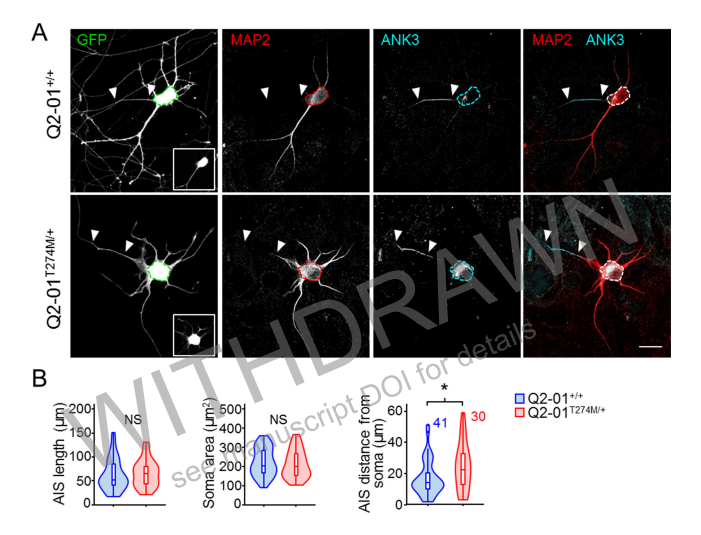
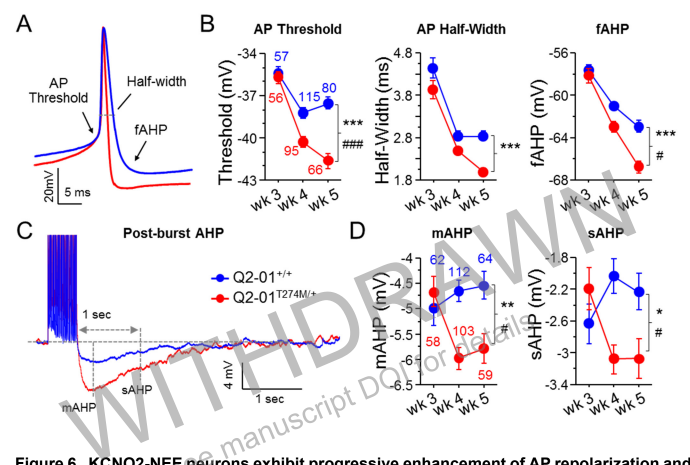


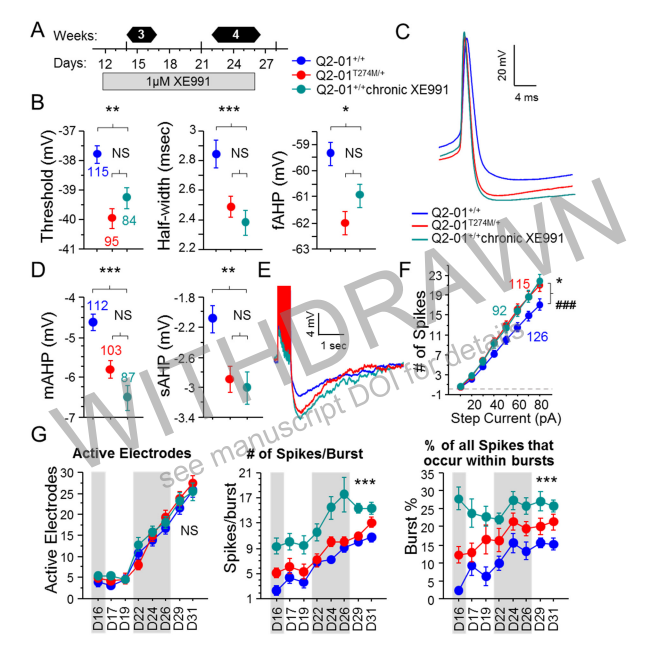
Figure 5. KCNQ2-NEE neurons exhibit a distal shift of the AIS. (A) Representative confocal images of 4-week old patient-derived and isogenic control neurons immunostained with GFP, MAP2 and ANK3. Dotted lines outline the soma. Insert at the bottom of GFP panel shows the specific image used to select the soma. White triangles indicate beginning and end of the axon initial segment (AIS). The end of the MAP2 signal that coincided with the beginning of the ANK3 signal was used at the start of the AIS. Scale bar: 20µm. (B) Morphometric analysis of patient-derived and isogenic control neurons. AIS longest path length, AIS total length and soma area were not significantly different. Patient-derived neurons exhibited a greater distance between the soma and the start of the AIS (24.9±2.7 µm) as compared to isogenic controls (17.5±1.7 µm; ttest: p = 0.0188). Number of neurons analyzed is displayed within the figure. NS: not significant. Values displayed are mean ± SEM.



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Figure 6. KCNQ2-NEE neurons exhibit progressive enhancement of AP repolarization and 1139 **post-burst AHP.** (A) Representative traces showing AP amplitude, threshold, half-width and 1140 fAHP. (B) Analysis of AP properties measured during week 3, 4 and 5. Patient-derived (red) 1141 neurons had significantly more hyperpolarized AP threshold (two-way ANOVA for genotype: 1142 1143 $F_{(1,463)} = 42.17$; ****p* < 0.0001; genotype/weeks interaction: $F_{(2,463)} = 9.16$; ###*p* = 0.0001) and 1144 enhanced repolarization with shorter AP half-widths (two-way ANOVA for genotype: $F_{(1,463)}$ = 1145 27.88; ***p < 0.0001) and larger fAHPs (two-way ANOVA for genotype: $F_{(1,463)} = 27.18$; ***p < 0.00010.0001; genotype/weeks interaction: $F_{(2, 463)} = 4.17$; #p = 0.02). Post hoc analysis using t-tests to 1146 1147 compare patient-derived and isogenic control neurons at each time point revealed differences in these properties only on week 4 and 5, with no significant differences on week 3. (C) 1148 1149 Representative whole-cell current-clamp traces showing post-burst AHPs after 50 Hz train of 25 APs evoked by 2 ms/1.2 nA suprathreshold current stimuli. (D) Patient-derived neurons had 1150 1151 enhanced mAHP (two-way ANOVA for genotype: $F_{(1,452)} = 9.83$; **p = 0.0018; genotype/weeks interaction: $F_{(2,452)} = 4.93$; #p = 0.0076) and sAHP (two-way ANOVA for genotype: $F_{(1,452)} = 4.86$; 1152 1153 *p = 0.0281; genotype/weeks interaction: $F_{(2, 452)} = 5.14$; #p = 0.0062). Post hoc analysis using ttests to compare patient and isogenic control neurons at each time point revealed differences in 1154 mAHP and sAHP only on week 4 and 5, with no significant differences on week 3. Number of 1155 neurons analyzed is displayed within the figure and in Table 1 (Also see Supplementary Figure 1156 1157 5). Values displayed are mean \pm SEM.

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Figure 7. Chronic Inhibition of M-current in Control Neurons Phenocopies KCNQ2-NEE AP 1163 **Properties.** (A) Experimental time line. (B) AP threshold, half-width and fAHP of isogenic control 1164 neurons chronically treated with 1 µM XE991 (Q2-01^{+/+}chronic XE991) were significantly different 1165 from untreated isogenic control neurons (ANOVA, Fisher's PLSD posthoc test for AP threshold: 1166 **p=0.003; AP half-width: ***p=0.0002; fAHP: *p=0.0107), but were not significantly different from 1167 patient-derived neurons (ANOVA, Fisher's PLSD posthoc test for AP threshold: p=0.137; AP half-1168 width: *p*=0.421; fAHP: *p*=0.112). (**C**) Representative AP traces from Q2-01^{T274M/+}. Q2-01^{+/+} and 1169 Q2-01^{+/+}chronic XE991 neurons. (D) The post-burst AHP is enhanced after chronic XE991 1170 treatment in isogenic control neurons (One-way ANOVA and Fisher's PLSD posthoc test for 1171 1172 mAHP: ***p < 0.0001; sAHP: **p=0.0008), to levels similar to patient-derived neurons (mAHP: p=0.0532; sAHP: p=0.69). (E) Representative traces showing post-burst AHPs of Q2-01^{T274M/+}, 1173 Q2-01^{+/+} and Q2-01^{+/+}chronic XE991 neurons. (F) Q2-01^{+/+}chronic XE991 were able to fire 1174 significantly more APs per current stimulus than untreated control neurons (repeated measures 1175 ANOVA: F_(2,2310)=3.88, *p=0.0216; interaction: F_(14,2310)=3.97, ###p<0.0001), and were not 1176 significantly different from patient-derived neurons (repeated measures ANOVA: ρ =0.9822). 1177 1178 Number of neurons analyzed is displayed within the figure and in Table 1 and 2 (also see

Supplementary Figure 6). (G) MEA recordings from day 16 to 31 in culture recorded from Q2-1179 01^{1274M/+}, Q2-01^{+/+} and Q2-01^{+/+} chronic XE991 neurons. There was no difference in the number 1180 of active electrodes between the three groups of neurons (repeated measures ANOVA: 1181 p=0.5603). Within the detected bursts Q2-01^{+/+}chronic XE991 neurons exhibited a significantly 1182 greater number of spikes as compared to both Q2-01^{T274M/+} and Q2-01^{+/+} neurons (repeated 1183 measures ANOVA: F_(2,315)=27.63, ***p<0.0001). The percentage of spikes that occurred within 1184 bursts was significantly greater in Q2-01+/+ chronic XE991 neurons as compared to both Q2-1185 1186 01^{T274M/+} and Q2-01^{+/+} neurons (repeated measures ANOVA: F_(2,315)=28.02, ****p*<0.0001). 16 wells were analyzed per group from 2 independent differentiations (See Supplementary Figure 1187 7). NS: not significant. Values displayed are mean ± SEM. 1188

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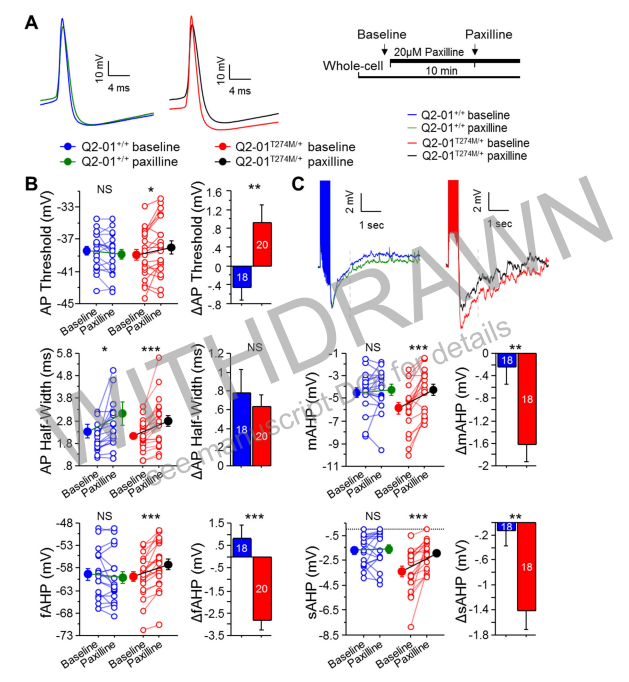


Figure 8. 1192 KCNQ2-NEE Neurons Exhibit a Dyshomeostatic Increase in BK Channel 1193 **Function.** (A) Right: Experimental protocol. Baseline measures were made after establishing the whole-cell configuration in current-clamp mode. After exactly 10 min of continuous perfusion of 1194 20 µM paxilline in aCSF, the AP properties and post-burst AHP were remeasured. Left: 1195 Representative trace of action potential before and after application of 20 µM paxilline. (B) Acute 1196 application of paxilline significantly decreased AP threshold in patient neurons (*p = 0.0193) but 1197 not in controls (p = 0.111). Paxilline reduced AP repolarization by increasing AP half-width equally 1198 1199 for patient and control neurons (ΔAP half-width: p = 0.589) but fAHP was only reduced in patient neurons (***p < 0.0001) and not affected in controls (p = 0.1882). (**C**) Top: Representative traces 1200 of post-burst AHPs. Acute application of paxilline significantly reduced mAHP (**p < 0.0001) and 1201 1202 sAHP (**p = 0.0002) in patient neurons but not in controls (mAHP: p = 0.393; sAHP p = 0.556). Repeated measures ANOVA was used to compare paxilline effects. Number of neurons analyzed 1203 1204 is displayed within the bar graphs, values displayed are mean \pm SEM.

1205	TABLES:
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1207 Table 1: Developmental Timeline of Intrinsic Membrane Properties of Patient-Derived and

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Weeks:	Week 3		We	ek 4	Week 5		
Genotype:	Q2-01+/+	Q2-01 ^{T274M/+}	Q2-01 ^{+/+} Q2-01 ^{T274M/+}		Q2-01+/+	Q2-01 ^{T274M/+}	
Resting Potential (mV)	-54.6 ± 0.6 (84)	-54.2 ± 0.6 (91)	-56.5 ± 0.5 (145)	-57.1 ± 0.5 (136)	-60.2 ± 0.6 (86)	-60.3 ± 0.6 (78)	
Input Resistance R _N at RMP (MΩ)	860.6 ± 35.4	998.3 ± 33.8 *	875.5 ± 25.2	932.9 ± 28.7	746.2 ± 28	732.6 ± 34.7	
Series Resistance Rs RMP (MΩ)	14.6 ± 0.3	13.9 ± 0.3	14.2 ± 0.3	13.6 ± 0.3	13.2 ± 0.4	13.9 ± 0.4	
AP Amplitude from baseline (mV)	87.6 ± 0.7 (57)	87.2 ± 0.8 (56)	91.9 ± 0.6 (115)	90.8 ± 0.6 (95)	93.2 ± 0.7 (80)	94.9 ± 0.8 (66)	
AP Threshold (mV)	-35.4 ± 0.5	-35.7 ± 0.4	-37.8 ± 0.3	-40 ± 0.3 ***	-37.5 ± 0.4	-41.6 ± 0.5	
AP Half-Width (ms)	4.4 ± 0.2	3.9 ± 0.2	2.9±0.1	2.5 ± 0.1 **	2.8 ± 0.1	2 ± 0.1 ***	
fAHP (mV)	-57.7 ± 0.6	-58.1±0.7	-59.4 ± 0.4	-62 ± 0.4 ***	-63 ± 0.6	-66.8 ± 0.6	
mAHP (mV)	-5 ±0.3 (62)	-4.7 ± 0.3 (58)	-4.7 ± 0.2 (112)	-5.9 ± 0.2 *** (103)	-4.5 ± 0.3 (64)	-5.7 ± 0.3 ** (59)	
sAHP (mV)	-2.7 ± 0.3	-2.2 ± 0.3	-2.2 ± 0.2	-3 ± 0.2 ***	-2.3 ± 0.2	-3.1 ± 0.3 *	

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1210 *p < 0.05, **p < 0.005, ***p < 0.0005: Fisher PLSD test comparing patient-derived neurons to 1211 isogenic controls during each week. Values displayed are Mean ± SEM. Number of neurons is 1212 indicated in ().

1213 TABLE ABBREVIATIONS: RMP - resting membrane potential; AP - Action potential; AHP -1214 afterhyperpolarization; fAHP - fast AHP; mAHP - medium AHP; sAHP - slow AHP

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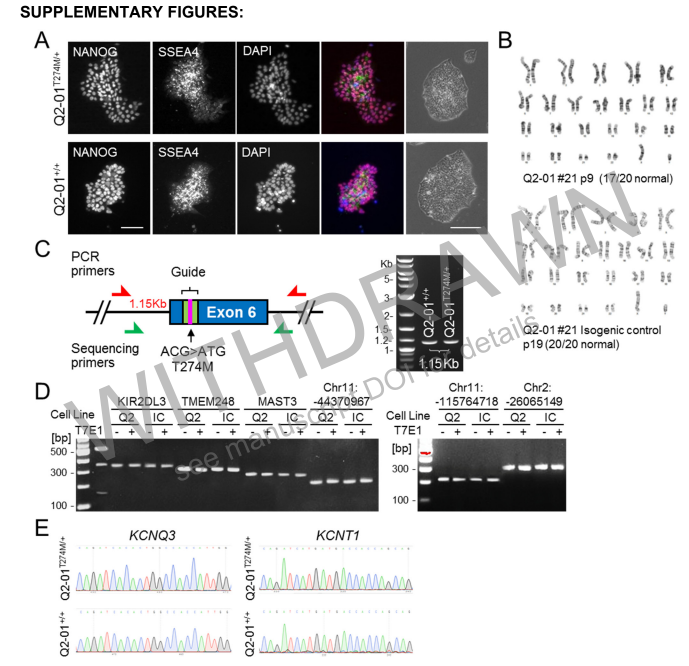
1223 Table 2: Week 4 intrinsic membrane properties of isogenic control neurons chronically

1224 treated with XE991

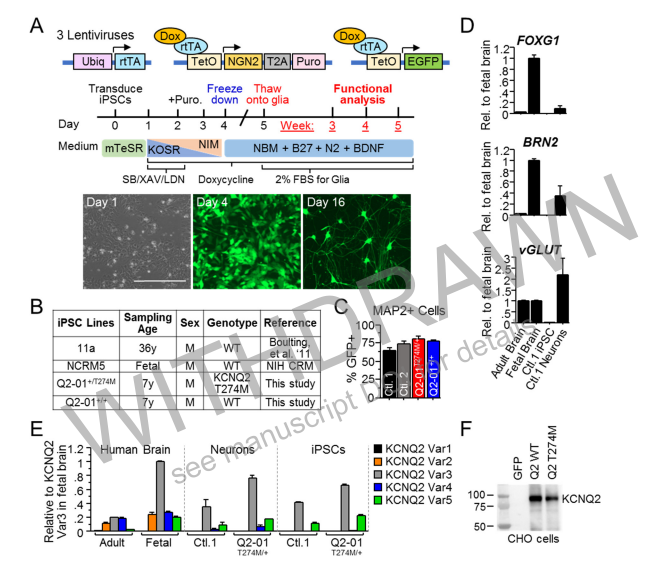
	Q2-01 ^{+/+} Chronic XE991 week 4	
Resting Potential (mV)	-60 ± 0.7 (106) ***	
Input Resistance R_N at RMP (M Ω)	984.1 ± 36 *	
Series Resistance Rs RMP (MΩ)	14 ± 0.3	
AP Amplitude from baseline (mV)	92.6 ± 0.7 (84)	
AP Threshold (mV)	-39.2 ± 0.4 **	
AP Half-Width (ms)	2.5 ± 0.1 ***	< r
fAHP (mV)	-61 ± 0.5 *	details
mAHP (mV)	-6.5 ± 0.3 (87) ***	of for details
sAHP (mV)	-3 ± 0,2 ** 0	99
	manus	

1226 *p < 0.05, **p < 0.005, **p < 0.0005: Fisher PLSD test comparing Q2-01^{+/+} neurons during week 1227 4 that were untreated (See Table 1) and chronically treated with XE991. Number of neurons is 1228 indicated in ().

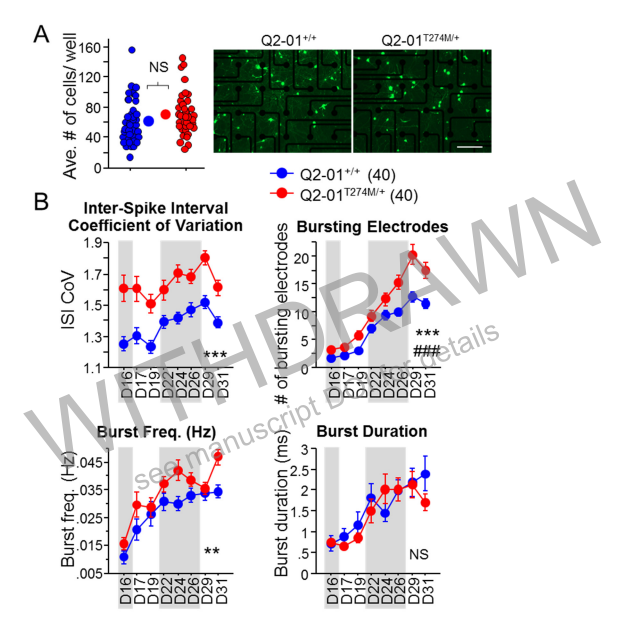
1229 TABLE ABBREVIATIONS: RMP - resting membrane potential; AP - Action potential; AHP -1230 afterhyperpolarization; fAHP - fast AHP; mAHP - medium AHP; sAHP - slow AHP



Supplementary Figure 1. Quality control studies of iPSC lines, related to Figure 1. (A) Left: 1242 Immunocytochemical labeling of KCNQ2-NEE patient-derived (Q2-01^{T274M/+}) and isogenic control 1243 1244 (Q2-01^{+/+}) iPSC lines with the pluripotency markers NANOG, SSEA4 and DAPI. Scale bar: 75µm. 1245 Right: Phase contrast images of iPSC colonies. Scale bar: 200 µm. (B) Karyotype analysis of patient and isogenic control iPSC lines. (C) Analysis of KCNQ2 targeted region after 1246 1247 CRISPR/Cas9 gene correction. To validate the absence of large indels or mosaicism, a 1.15Kb 1248 DNA fragment of genomic DNA around the targeted site was amplified and subjected to Sanger 1249 sequencing (Figure 1B). Products from both the parental and corrected iPSC lines exhibited one 1250 band of the expected size. (D-E) Analysis of potential CRISPR off-target sites. The top eight 1251 genomic regions of homology with the CRISPR/Cas9 targeted region (Supplementary Table 1) were analyzed by a T7 endonuclease assay or Sanger Sequencing. No mutations were identified. 1252 1253 (E) Two potential off-target sites were found in K⁺ channels (*KCNQ3* and *KCNT1*). PCR primers 1254 were designed to amplify ~850bp fragments around off-target sequence (Supplementary Table 1). DNA sequencing electropherograms showed no mutations in these sites. 1255

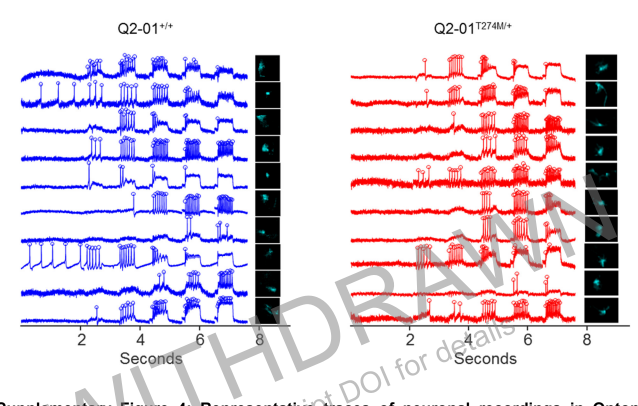


Supplementary Figure 2. Quality control studies of iPSC-derived neurons, related to 1256 1257 Figure 1. (A) Top: Design of lentiviral vectors for Nan2-mediated conversion of iPSCs to cortical excitatory neurons through tetracycline-inducible expression of exogenous proteins driven by a 1258 tetracycline-inducible promoter (TetO). Cells are transduced with (i) a virus expressing reverse 1259 tetracycline-controlled transactivator (M2rtTA) driven by the Ubiquitin promoter, (ii) a virus 1260 1261 expressing Ngn2 linked by T2A to a puromycin (puro) resistance gene driven by TetO, and (iii) EGFP driven by the TetO promoter. Middle: Protocol for generation of cortical excitatory neurons 1262 1263 (See methods). Bottom: Representative images illustrating the time course of the conversion of 1264 iPSCs into neurons. Corresponding bright field and GFP fluorescence images on day 1 before addition of doxycycline, day 4 before cryopreservation, and day 16 after thawing onto a mouse 1265 1266 glia monolayer (day 5). This protocol is modified from Zhang et al., 2013. Scale bar: 400µm. (B) 1267 Table of iPSC lines used in this study. (C) Quantification of MAP2 immunopositive staining with MAP2 antibody coincident with GFP fluorescence in iPSC-derived neurons. More than 65% of 1268 1269 the MAP2 positive neurons were also positive for GFP in all iPSC lines as quantified by 1270 fluorescent microscopy (Figure 1C). (D) RNA expression analysis using quantitative RT-PCR of vGLUT2, FOXG1 and BRN2 that are characteristic of excitatory layer 2/3 cortical neurons in 1271 healthy control iPSC-derived neurons. Human adult and fetal brain were used as controls. (E) 1272 1273 RT-qPCR expression analysis of KCNQ2 splice variants in the differentiated neuronal cultures 1274 using isoform-specific primers (Supplementary Table 2). Human adult and fetal brain were used as controls. (F) Western blot analysis of proteins isolated from CHO cells transfected with 1275 1276 plasmids encoding either wild type or T274M mutant KCNQ2 human variant 4.

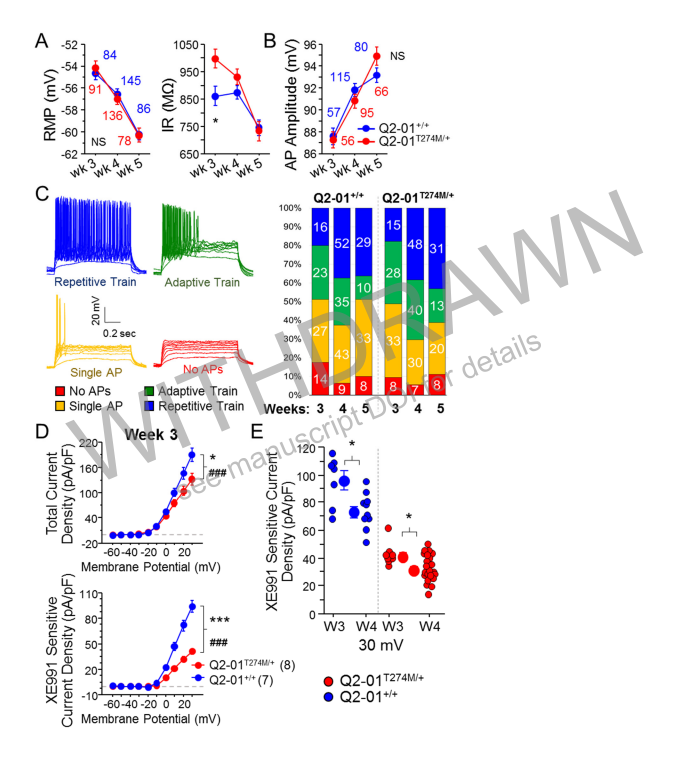


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1278 Supplementary Figure 3. MEA quality control and bursting measurements, related to Figure 2. (A) Each MEA plate well was imaged during week 4 in culture. GFP-fluorescing 1279 neurons on the electrode field area were counted for each well of every plate. The average 1280 number of cells per well was not different between Q2-01^{T274M/+} patient-derived (71.1 ± 4.6 1281 1282 neurons/well) and Q2-01^{+/+} isogenic control (61.4 \pm 4.3 neurons/well) wells (p = 0.1295, N = 40). 1283 Right: Representative images from which GFP-fluorescing cells were counted. Scale bar: 200 µm. (B) The interspike interval coefficient of variation (ISI CoV) was significantly greater in 1284 1285 patient-derived neurons (repeated measures ANOVA for genotype: $F_{(1,546)} = 47.94$, ***p < 0.0001). The number of bursting electrodes was significantly greater in patient-derived neurons 1286 (repeated measures ANOVA for genotype: $F_{(1,546)} = 18.82$, ***p < 0.0001; genotype/day 1287 interaction: $F_{(7,546)} = 6.71$, ###p < 0.0001). The burst frequency was higher in patient-derived 1288 neurons (repeated measures ANOVA for genotype: $F_{(1,546)} = 10.35$, **p = 0.0019). However, the 1289 1290 burst durations were not different between patient-derived and isogenic control neurons 1291 (repeated measures ANOVA for genotype: $F_{(1,546)} = .289$, p = 0.592; See Figure 2). NS: not significant. Number of neurons analyzed is displayed within the figure. Values displayed are 1292 1293 mean ± SEM.



- Supplementary Figure 4: Representative traces of neuronal recordings in Optopatch experiments. Fluorescence-time traces showing detected spikes (Δ F/F₀; circles above spikes) and corresponding images of neurons for KCNQ2-NEE patient neurons (right) and isogenic controls (left).
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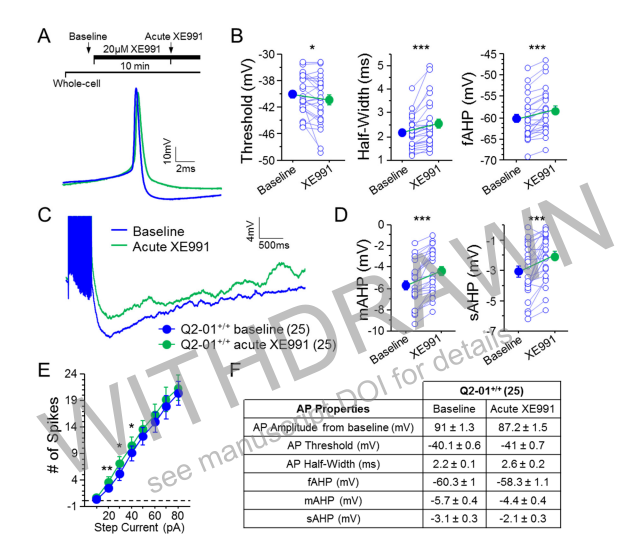
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1305 Supplementary Figure 5. Intrinsic excitability by week in culture and M-current during week 3, related to Figures 3 and 5. (A) Passive membrane properties. Left: Time course of 1306 1307 resting membrane potential (RMP) and input resistance (IR) for iPSC-derived neurons in culture. As neurons matured over the 3-week period, the RMP became more hyperpolarized with no 1308 differences between cell lines, whereas IR became smaller over time in culture but was higher 1309 1310 for patient neurons only during week 3 (Two-way ANOVA for input resistance measured at RMP: $F_{(1,614)} = 5.54$, *p = 0.019; t-test comparing isogenic control pair at week 3: *p = 0.0054; week 4: 1311 p = 0.132; week 5: p = 0.759). (B) Action potential amplitudes over time in culture. There were 1312 no differences between cell lines. (C) Qualitative analysis of neuronal firing patterns. At each time 1313 point in culture there was heterogeneity in the firing patterns exhibited by the neurons during a 1 1314 sec somatic current stimulus (10-80pA) protocol. Neurons were grouped based on specific 1315

defined criteria: No APs; Single APs; at least 1 AP fired during any of the 10-80pA steps but no more than 10 APs; Adaptive train: 11 or more APs; Repetitive train: 11 or more APs with at least 10 APs having amplitudes more than 85 mV from baseline. The percent of neurons that fell into specific group for each time point and cell line is indicated within columns. (D) Outward current measured during week 3 in culture recorded from patient-derived (red) or isogenic control (blue) neurons. Quantification of total current density (top: repeated measures ANOVA for total current density during week 3 by genotype: $F_{(1,117)} = 4.7$, **p* = 0.0493; genotype/voltage step interaction: $F_{(9,117)} = 6.57$, ###p < 0.0001), and XE991-sensitive current (bottom: repeated measures ANOVA) for M-current density during week 3 by genotype: $F_{(1,117)} = 27.64$, ***p = 0.0002; genotype/voltage step interaction: F(9,117) = 36.99, ###p < 0.0001). (E) Plot of M-current densities recorded at 30 mV from all neurons during week 3 or week 4 (Figure 1G) Comparison between weeks was significantly different for both groups of neurons (*t test* for isogenic controls neurons: **p* = 0.015, patient-derived neurons: p = 0.016). NS: not significant. Number of neurons analyzed is displayed within the figure and in Table 1. Values displayed are mean ± SEM.

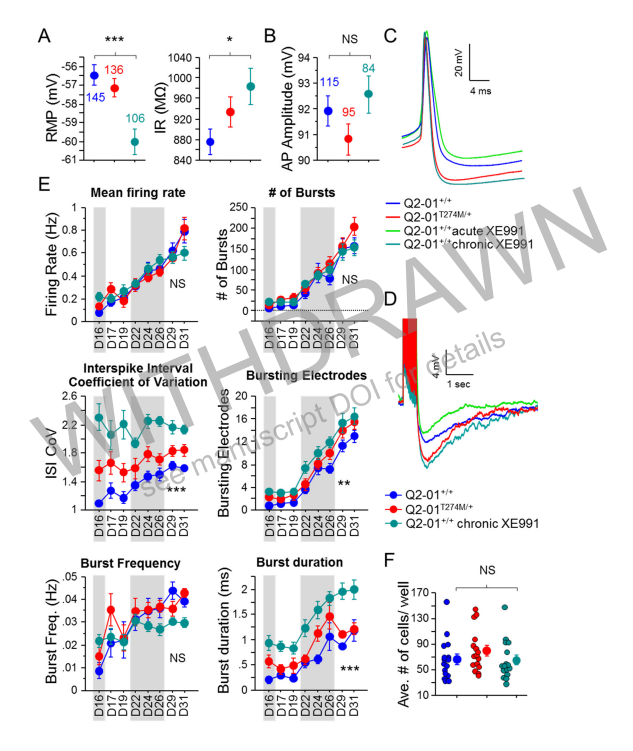
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Supplementary Figure 6. Acute inhibition of M-current in control neurons enhances 1360 excitability and decreases AP repolarization and post-burst AHP, related to Figures 6 and 1361 7. (A) Top: Experimental protocol. Baseline measures were made after establishing the whole-1362 cell configuration in current-clamp mode. After exactly 10 min of continuous perfusion of 20 µM 1363 XE991 in aCSF, the AP properties, post-burst AHP and number of spikes fired per current 1364 stimulus were remeasured. Bottom: Representative whole-cell current-clamp recordings showing 1365 AP properties of isogenic control neurons during week 4 before (blue) and after (green) acute 1366 1367 application of XE991. (B) Acute application of XE991 significantly decreased AP threshold (*p =0.033), and reduced AP repolarization with longer AP half-width (***p = 0.0003) and smaller fAHP 1368 (***p < 0.0001). (C) Representative whole-cell current-clamp recordings showing post-burst 1369 1370 AHPs before and after acute application of 20 µM XE991. (D) Acute application of XE991 significantly reduced mAHP (***p < 0.0001) and sAHP (***p = 0.0001). (E) Number of APs fired 1371 1372 by isogenic control neurons in response to 1 sec somatic current stimuli (10-80pA) from a holding 1373 potential of -65mV. Neurons were able to fire before and after acute application of 20µM XE991, but blocking M-currents resulted in an increase of APs fired only with 20 – 40pA current stimuli 1374 (p-value = 0.003, 0.048, 0.021; for 20, 30 and 40pA, respectively). (F) Summary of AP and AHP 1375 1376 properties before and after XE991 application in isogenic control neurons during week 4. Repeated measures ANOVA was used to compare XE991 effects. N = 25 neurons, values 1377 1378 displayed are mean ± SEM.



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1382 Supplementary Figure 6. Effects of chronic XE991 treatment on control neuron intrinsic membrane properties and MEA recordings, related to Figure 6. (A) Passive membrane 1383 1384 properties. Left: RMP became more hyperpolarized with chronic XE991 treatment (Q2-01^{+/+}chronic XE991) in isogenic control neurons (ANOVA, Fisher's PLSD post hoc test: 1385 ****p*<0.0001). Right: Input resistance was greater in Q2-01^{+/+}chronic XE991 neurons compared 1386 to Q2-01^{+/+} (ANOVA, Fishers PLSD post hoc test: *p=0.0114). Number of neurons analyzed is 1387 displayed within the figure and in Tables 1 and 2. (B) Action potential amplitudes were not 1388 different between cell lines and after chronic treatment with XE991 (See Figure 7). (C) 1389 Representative AP traces from patient-derived neurons, isogenic control neurons, and control 1390 neurons chronically or acutely treated with 1 or 20 µM XE991, respectively. (D) Representative 1391 1392 traces showing post-burst AHPs of patient-derived neurons, isogenic control neurons, and control

neurons chronically or acutely treated with 1 or 20 µM XE991, respectively. (E) Longitudinal analysis of MEA recordings from patient-derived neurons (Q2-01^{T274M/+}) and isogenic control neurons, which were either not treated (Q2-01^{+/+}) or treated chronically with 1µM XE991 starting on day 12. There was no difference in mean firing frequency or number of bursts fired by control neurons chronically treated or not treated with XE991. However, the interspike interval coefficient of variation (ISI CoV) was significantly higher in Q2-01^{+/+}chronic XE991 neurons as compared to both patient-derived neurons and isogenic control neurons (repeated measures ANOVA: F_(2.315)=50.15, ***p<0.0001). The number of bursting electrodes was significantly greater in Q2-01^{+/+}chronic XE991 neurons as compared to untreated isogenic controls (repeated measures ANOVA: F_(2,315)=6.08, **p=0.0046; Fisher's PLSD: p=0.0011). While the burst frequency was not different between groups (repeated measures ANOVA: $F_{(2,315)}=2.75$, p=0.0752), the burst duration was significantly longer in Q2-01^{+/+}chronic XE991 neurons (repeated measures ANOVA: $F_{(2,315)}$ =39.68, ***p<0.0001). 16 wells were analyzed per genotype treatment group from 2 independent differentiations (See Figure 7). (F) Each MEA plate well was imaged during week 4 in culture. GFP-fluorescing neurons on the electrode field area were counted for each well of every plate. The average number of cells per well counted was not different between Q2-01^{T274M/+} see manuscript Dol for det patient-derived neurons (80.3±8.2 neurons/well). Q2-01+++ isogenic control neurons (66.8±8 neurons/well) and Q2-01^{+/+}chronic XE991 neurons (65.3±7.6 neurons/well) wells (p=0.3469). NS: not significant. Values displayed are mean ± SEM.

1436 SUPPLEMENTARY TABLES:

Supplementary Table 1. Single guide RNA off targets, related to Figure 1 and Supplementary Figure 1.

Single Guide RNA (sgRNA) KCNQ2	5' CAGATCATGCTGACCACCATTGG 3'
Donor Single Stranded Repair Oligonucleotide	5'-AGTCGCCAGCGGGCGTCCAGCCTGCCCTCAGGGGTGTGAGCAGGCCCTTCGTGTGAC TAGAGCCTGCGGTCCCACAGATC <u>ACG</u> CTGACC <u>ACT</u> ATTGGCTACGGGGACAAGTACCCC CAGACCTGGAACGGCAGGCTCCTTGCGGCAACCTTCACCCTCATCGG -3'

Off Target Sequence CAGATC <u>ATG</u> CTGACC <u>ACC</u> ATTGG	Score	Gene Locus	PCR primer set		Size (bp)	s	equencing Primers
CAGATCACGCTGACCACCATTGG 100 NM 172107		F	AACTAAGCACAACC CCTGGG	1150	F	GCCATGGCTCGG TGGAGA	
CAGATOA <u>C</u> GOTGACCACCATTGG	100	chr20:-62071037		TCTGCAGGCCCATC TTGAAG	I I SU	R	TGAAGGGCACAC AGTGAAGG
CAGATCATGATGACCACCAGCAG	3.3	KCNT1 NM_020822	F	GCACACATATATTC ACAAACATGGC	842	F	ACACACACATGA ATGCTCGTG
	5.5	chr9:-138651650	R	GGACAAGAGAAGG GAACTCACA	042	R	AAATGAAGGACC TGCTCCGT
	0.6	KCNQ3 NM_004519 chr8:-133186572	F	CCTGGCTGTGGAT GGGAAAT	867	F	CTGTCCACCTTG ACCCTTCC
	mai nai		R	ATCCCCTGCTCCCA GAGAAT		R	CCAGGGACCTAG GAGAGAGC
CACACCATGCTGACGACCATGAG		KIR2DL3 NM 015868	F	GACGTCTTTTGAGT CTGGTCG	328		
		chr19:-55250012	R	ATCTCCATCCCCGC ACT	520		
	0.4	TMEM248 NM_017994 chr7:-66406902	F	CTGTTGGCAAGTCA GTCCTTG	297	207	
C <u>T</u> GATCATG <u>AA</u> GACCACCA <u>G</u> GGG			R	TATATGGCAAGCCA CAGGGTG	291		
0404404T00T040040444000	0.3	MAST3 NM_015016 chr19:-18241435	F	CAGCGCTTTGCCAT CAAGAAG	270		
CAGA <u>A</u> CATGCTGACCAC <u>A</u> A <u>A</u> GGG	0.5		R	ACAGAACAGACCTC AACTTGGT			
AAGGTCATCCTGACCACCATAAG	1.6	Intronic/ Intergenic chr11:-44370967	F	AGCTCTGGAAAGCT GCCTTATC	204		
			R	GAGACAGTCCAAGA CGTGGC	204		
CACATCCTGGTGACCACCATCAG	1.6	Intronic/ Intergenic chr11: 115764718	F	GGTCTTGGCATGCC TTTCTG	207		
			R	TCTGTCTTCCAACC CTGGCT			
CAGTACATGCTGACCACCCTTAG	1	Intronic/ Intergenic	F	GAACACAGAACACT ACAACAAACT	277		
		chr2:-26065149	R	AGGTAGTAAAGTGT GAATCATCAGA	211		

1443 Supplementary Table 2: RT-qPCR primers, related to Supplementary Figure 2

	NCBI Gene ID	Gene Target		Primer 5'->3' Sequence	Size (bp)	Source
Neuronal Markers	2290	FOXG1	F	ATACTGTGGTCATATGCCCGTG	165	E.K.
			R	AGCCTTTGAATTCCCTATAAGTTGA	100	
	5454	POU3F2/ BRN2	F	TGGATTCCCATCAGGAAAGAGG	170	E.K.
eui Aar	5454		R	AAAATCCCCCAAAACGGCAA	170	L.N.
Z 2	33427	vGLUT2	F	GGGAGACAATCGAGCTGACG	156	E.K.
	55427	VGLUTZ	R	TGCAGCGGATACCGAAGGA	150	L.K.
KCNQ2 Splice Variants	3785	KCNQ2 Var 1 NM_172107.3	F	TACGGGGCCTCCAGACTTAT	129	D.S.
	3705		R	GCACGGGCTGCCTTTACTTG	129	
	3785	KCNQ2 Var 2 NM_172106.2	F	CGCAAACTCAAACCTACGGG	139	D.S.
			R	CTGACCTTCTGGCTTGGAGA		
	3785	KCNQ2 Var 3 NM_004518.5	F	GCACTTTGAGAAGAGGCGGA	144	D.S.
			R	ATAAGTCTGTACATGGGCACG	144	
	3785	KCNQ2 Var 4 NM_172108.4	F	ACGGCAGAACTCAGAAGCAAG) 155	D.S.
×			R	TGAACTTCCGCTTGGACACC	100	
	3785	KCNQ2 Var 5 NM_172109.2	F	GCCTGGAGATTCTACGCCAC	123	D.S.
			R	TTGCTTGGTGGCAGGTG		
5	5080	GAPDH	F	ACAACTTTGGTATCGTGGAAGG	101	PrimerBank: 378404907c2
inç			R	GCCATCACGCCACAGTTTC	101	
House keeping	2821		F	GTGTACCTTCTAGTCCCGCC	101	
- <u>×</u>		GPI (F1/R1)		GGTCAAGCTGAAGTGGTTGAAGC	101	G.L.R.