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Citation for published version:

Digital Object Identifier (DOI):
10.1523/ENEURO.0426-17.2018

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Peer reviewed version

Published In:
eNeuro

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Research Article: Confirmation | Disorders of the Nervous System

Circadian and Brain State Modulation of Network Hyperexcitability in Alzheimer’s Disease

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DOI: 10.1523/ENEURO.0426-17.2018

Received: 5 December 2017
Revised: 8 March 2018
Accepted: 6 April 2018
Published: 27 April 2018

Author Contributions: RB, RRR and IO designed research; RB, ADL, AG-S, MJ, RC-CC, I-C, A-LH, TS and TS performed research; ADL, AY, RC-CC, MT, CYJ, A-LH and IO analysed the data; MAJ, A JC, SSC and KH contributed unpublished analytical tools; RB, ADL, KH and IO wrote the paper.

Funding: http://doi.org/10.13039/501100000320Alzheimer’s Society: PG-2012-208. RS Macdonald Charitable Trust; Muir Maxwell Epilepsy Centre; Euan MacDonald Centre; Patrick Wild Centre; http://doi.org/10.13039/100005339American Academy of Neurology (AAN); http://doi.org/10.13039/501100000295Epilepsy Research UK (ERUK); http://doi.org/10.13039/501100000265Medical Research Council (MRC): MR/M024075/1.

Conflict of Interest: Authors declare no conflict of interest.

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Cite as: eNeuro 2018; 10.1523/ENEURO.0426-17.2018

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Accepted manuscripts are peer-reviewed but have not been through the copyediting, formatting, or proofreading process.

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1. **Manuscript Title:** Circadian and brain state modulation of network hyperexcitability in Alzheimer’s disease

2. **Abbreviated title:** Hyperexcitability modulation in AD

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MAJ, AJC, SSC and KH contributed unpublished analytical tools. RB, ADL, KH and IO wrote the paper.

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6. **Number of Figures:** 7

7. **Number of Tables:** 2

8. **Number of Multimedia:** 0

9. **Number of words for Abstract:** 250

10. **Number of words for Significance Statement:** 111

11. **Number of words for Introduction:** 656

12. **Number of words for Discussion:** 2069

13. **Acknowledgements:** We thank the Gladstone Institute for providing J20 mice. We thank Dominic Walsh for supplying APPNL/F mice.

14. **Conflict of interest:** The authors declare no conflict of interest.

15. **Funding sources:** This work was supported by the grants awarded to IO from the following funders: the Alzheimer’s Society (PG-2012-208), the RS Macdonald Charitable Trust, The Muir Maxwell Epilepsy Centre, The Euan MacDonald Centre and The Patrick Wild Centre. ADL was supported
by the American Academy of Neurology Institute. AGS was supported by Epilepsy Research UK. RC-CC was supported by an MRC grant (MR/M024075/1) awarded to RRR.
Abstract

Network hyperexcitability is a feature of Alzheimer’s disease (AD) as well as numerous transgenic mouse models of AD. While hyperexcitability in AD patients and AD animal models share certain features, the mechanistic overlap remains to be established. We aimed to identify features of network hyperexcitability in AD models that can be related to epileptiform activity signatures in AD patients. We studied network hyperexcitability in mice expressing amyloid precursor protein (APP) with mutations that cause familial AD, and compared a transgenic model that overexpresses human APP (J20), to a knock-in model expressing APP at physiological levels (APPNL/F). We recorded continuous long-term electrocorticogram activity from mice, and studied modulation by circadian cycle, behavioural, and brain state. We report that while J20s exhibit frequent inter-ictal spikes (IIS), APPNL/F mice do not. In J20 mice, IIS were most prevalent during daylight hours and the circadian modulation was associated with sleep. Further analysis of brain state revealed that IIS in J20s are associated with features of rapid-eye movement (REM) sleep. We found no evidence of cholinergic changes that may contribute to IIS-circadian coupling in J20s. In contrast to J20s, intracranial recordings capturing IIS in AD patients demonstrated frequent IIS in non-REM sleep. The salient differences in sleep-stage coupling of IIS in APP overexpressing mice and AD patients suggests that different mechanisms may underlie network hyperexcitability in mice and humans. We posit that sleep-stage coupling of IIS should be an important consideration in identifying mouse AD models that most closely recapitulate
network hyperexcitability in human AD.

**Significance statement**

It is increasingly recognized that Alzheimer’s disease (AD) is associated with hyperexcitability in brain networks. Brain network hyperexcitability is also reported in several rodent models of AD. We studied the signatures of this hyperexcitability in two rodent models of AD as well as AD patients. Network hyperexcitability was prevalent in a transgenic model of AD, but was absent in a rodent model that is considered to be more physiological. Moreover, while network hyperexcitability was coupled to rapid-eye movement (REM) sleep in transgenic mice, hyperexcitability occurred in non-REM sleep in AD patients. We suggest that brain state coupling of hyperexcitability can be used as a method for screening animal models of AD.
Introduction

An increased incidence of seizures in Alzheimer’s disease (AD) is indicative of an underlying network hyperexcitability (Hesdorffer et al., 1996; Amatniek et al., 2006; Lozsadi and Larner, 2006; Vossel et al., 2013; Cretin et al., 2016). Interictal spikes (IIS) are also seen in a high proportion of AD patients without a history of clinical seizures (Vossel et al., 2016). Non-ictal network hyperactivity has also been detected by means of fMRI in individuals at risk of developing dementia, for example in people carrying the APOE4 allele (Bookheimer et al., 2000; Filippini et al., 2009), with other genetic predictors of AD (Quiroz et al., 2010) and also in patients with mild cognitive impairment – a diagnosis which is considered to be a prodromal stage of AD (Dickerson et al., 2005). Network hyperexcitability and seizure activity appear at early stages of the disease and have been suggested to be predictors of accelerated cognitive decline (Amatniek et al., 2006; Vossel et al., 2013; Cretin et al., 2016; Vossel et al., 2016).

Network hyperexcitability has also been reported in numerous mouse models of AD pathology (Palop et al., 2007; Minkeviciene et al., 2009; Busche et al., 2012; Šišková et al., 2014; Kazim et al., 2017), with the aberrant activity being a feature that occurs in advance of plaque deposition (Busche et al., 2012; Bezzina et al., 2015). These phenomenological similarities have led to the suggestion that these animal models can provide a tool by which to study network hyperexcitability in human AD (Palop and Mucke, 2016).
Aberrant network activity could in itself contribute to neurodegeneration and cognitive dysfunction in AD pathology (Cirrito et al., 2005; Bero et al., 2011; Busche and Konnerth, 2015; Wu et al., 2016). Reducing network hyperexcitability has been shown to ameliorate cognitive dysfunction in both patients and animal models (Bakker et al., 2012; Sanchez et al., 2012; Haberman et al., 2017), and to attenuate Aβ pathology (Yuan and Grutzendler, 2016). Hence, targeting network hyperexcitability has been suggested as a novel therapeutic avenue for AD. However, studying this therapeutic avenue by means of animal models (Sanchez et al., 2012) requires a deeper understanding of the shared features of network hyperexcitability between AD patients and animal models.

Expression of epileptiform activity frequently exhibits a circadian pattern and shows preferential activation with specific brain states in a range of epilepsies (Quigg, 2000; Ng and Pavlova, 2013; Sedigh-Sarvestani et al., 2015). Circadian dysfunction and sleep disruption are common features of AD, and are also considered as early features of disease pathogenesis (Musiek et al., 2015; Mander et al., 2016; Musiek et al., 2018). Two recent papers have reported modulation of epileptiform activity by circadian cycle and brain state in transgenic AD models. Epileptiform activity was more prevalent in daylight conditions, and was suggested to occur primarily during REM sleep (Born et al., 2014; Kam et al., 2016). If epileptiform activity is modulated by circadian-cycles and/or brain state in AD patients, it is possible that this might contribute to the reported circadian alterations and sleep dysfunction. In line with this, it has recently been shown that inter-ictal activity in AD patients is highly prevalent during sleep (Vossel et al., 2016; Horváth et al., 2017; Lam et al., 2017). The
modulation of ictal related activity by brain state points to a distinguishing
feature that could be used to (a) uncover distinct mechanisms underlying
hyperexcitability, and (b) ascertain the translational utility of specific animal
models in studying network hyperexcitability. To this end, the present study
aimed to investigate circadian and brain state modulation of network
hyperexcitability in two rodent models of AD of differing aetiology: one in which
hAPP is overexpressed and one in which APP is expressed at endogenous levels.
In order to shed light on the translational utility of rodent AD models for
studying network hyperexcitability in human AD, we further examined sleep-
stage modulation of epileptiform activity in two patients with AD, using
recordings from intracranial electrodes placed directly adjacent to the
hippocampus.

Methods

Animals and animal maintenance

All animal procedures were performed in accordance with the [Author
University] animal welfare committee regulations and were performed under a
UK Home Office project license.

Heterozygous mice expressing hAPP with the KM670/671NL (Swedish) and
V717F (Indiana) mutations on a PDGFβ promoter (J20; Mucke et al., 2000) were
bred by crossing J20 +/- males with C57Bl6J females. Experiments used J20+/-
(n=21) and J20 -/- (n=8) wild-type (WT) littermate controls. The mean age of J20
animals was 5 months (range: 3.3 – 6.5 months).
Homozygous knock-in mice expressing APP KM670/671NL (Swedish) and APP I716F (Iberian) mutations (APP<sup>NL/F</sup>; Saito et al., 2014) were back-crossed onto C57Bl6J strain for at least 3 generations and were >99.8% cogenic with C57Bl6J. Experiments used APP<sup>NL/F</sup> +/+ (n=20) and age-matched non-littermate C57Bl6J WT controls (n=15). Animals were either 8 months or 12 months of age.

Both male and female mice were used. Mice were kept on a 7h-19h light-dark cycle in standard, open cages. Mice were group-housed prior to surgery and were housed individually post-surgery and during telemetry data acquisition.

**Surgery and data acquisition**

A subdural intracranial electrocorticogram (ECoG) recording electrode was positioned in the cortex overlying the hippocampus (co-ordinates x: -2.25 y: -2.46). A reference electrode was implanted either in the skull of the contralateral hemisphere, or above the cerebellum. Electrodes were either bare wire, or skull screws. An EEG transmitter (A3028B, Open Source Instruments) was implanted on the back of the animal subcutaneously. Animals were left to recover for at least 24 hours after surgery before the commencement of telemetry data acquisition. Telemetric ECoG data was acquired for approximately 3 days from each animal. Recording was either carried out continuously between days 1 to day 3 after surgery, or day 1, followed by day 5 to day 6.

ECoG data was acquired using an OpenSource Instruments data acquisition system at 512sps as previously described (Chang et al., 2011).
Video data was acquired using a Basler aca1300-60gm gigE camera sampling at 10fps, or a Logitech C270 HD webcam sampling at 5fps. Video was acquired during the daylight hours.

**ECoG Data processing**

The raw ECoG data was analysed using custom written Tcl and C processors. ECoG data was analysed in 8s intervals. For each interval we extracted measures of data loss, spike count, delta power (0.1-3.9 Hz) and theta power (4-12 Hz). We defined intervals in which data loss exceeded 20% of samples as “lossy” intervals. Intervals in which delta power exceeded 0.16mV$^2$ were classified as artifacts. Lossy and artifact intervals were excluded.

Inter-ictal spikes (IIS) in rodent ECoG were detected as follows. Each 8-s interval of EEG was treated as a two-dimensional path. One dimension is voltage, which was normalized by dividing by the mean absolute step size of the voltage in the 8-s interval. The mean absolute step size is the sum of the absolute changes in voltage from one sample to the next, divided by the number of samples. For an 8-s interval, the number of samples would be 4096 and a typical mean absolute step size for mouse EEG is around 12 μV. The other dimension is time, which was normalized by dividing by the sample period. The spike-finder proceeds along this EEG path in steps. With each step, it moves to the nearest sample on the path ahead. Whenever the spike-finder steps past one or more samples, it classifies these samples as an aberration in the path. Solitary aberrations larger than 20 mean absolute step sizes are classified as IIS. A series of IIS in which single
spikes were separated by <78ms (40 samples) were treated as a burst event and counted as one IIS event within the 8s interval.

For each J20 animal, the false positive rate of IIS detection was determined by randomly hopping through 100 8s intervals identified by the processor as containing IIS, and scoring them as true or false positives. The animal was excluded from analysis if the false positive rate exceeded 10%. One animal was excluded from analysis on this basis. In the remaining animals, the false positive rate ranged from 0 – 6% (mean false positive rate: 1.9%).

We observed that lossy and artifactual intervals resulted from movement and external sources of interference. We could not exclude the possibility that these events are non-randomly distributed across the 24hr cycle. Non-random exclusion of intervals would impact the evaluation of coupling of IIS. We thus set a criterion: if >5% of all 8s intervals were excluded due loss or artifact, the animal was excluded from calculations of coupling of IIS to circadian cycles, sleep-wake, and θ/δ. Two J20 animals were excluded from data reported in Fig 2-4 on this basis (25% and 16% of 8s intervals excluded in these animals).

**Video analysis**

Video data was manually scored to classify periods as ‘sleep’ or ‘wake’. Based on previous reports, sustained inactivity ≥40s was classified as ‘sleep’, while stationary periods <40s and periods of movement were classified as ‘wake’ (Pack et al, 2007). Postural shifts during sleep epochs did not break sleep epochs.
Immunohistochemistry and imaging

Animals were killed by transcardial perfusion with N-methyl-D-glucamine (NMDG)-based saline solution (in mM: 92 NMDG, 2.5 KCl, 1.25 NaH$_2$PO$_4$, 20 HEPES, 30 NaHCO$_3$, 25 glucose, 10 MgCl$_2$, 0.5 CaCl$_2$, sucrose to adjust osmolarity to 315-330mOsm). Brains were post-fixed with 4% paraformaldehyde for 24h then washed and stored in PBS. Samples were put in 50% or 30% sucrose, PBS solution and 50% OCT solution for 24h before cutting, then placed in the same solution and cut using a freezing microtome.

50μm sections were stored in PBS at 4°C. Slices were presoaked with 5% rabbit normal serum (RNS, Vector S-5000), 0.2% Triton X-100, PBS solution for 30min at room temperature (RT), followed by incubation with 3% RNS, 0.2% Triton X-100, anti-Choline Acetyltransferase (ChAT; 1:500, Millipore #AB 144P, RRID: AB_2079751), PBS solution for 48h at 4°C. The sections were washed 3 times with PBS 0.2% Triton X-100 for 5min each and then incubated in 3% RNS, anti-Goat biotinylated (1:200), Dapi (1:5000, Sigma D9542-1MG), PBS solution for 1h at RT. After 3 PBS 0.2% Triton X-100 washings of 5min each, the sections were incubated with ABC reagent (Vectastain PK-6105 kit) prepared half an hour before using and stored in foil at 4°C containing 0.1% of A, 0.1% of B, 0.01% Triton X-100, PBS for 1h at RT. After 6 PBS washings of 10min each, the sections were put in 3 3'-diaminobenzidine (Sigma D5905-50TAB), 0.02% CoCl$_2$ (1% wt/vol), 0.04% (NH$_4$)$_2$Ni(SO$_4$)$_2$ (1% wt/vol) dH$_2$O solution for 30min at 4°C over agitation. Then stained by adding 1.2% of fresh 1% H$_2$O$_2$ per slice for 10 to 20s until the slice darkened. The slices were then transferred and washed in PBS, 6 times for 10min each, mounted on a slide and dried for 30min at 50°C then
finally covered with Mowiol Embedding Medium and coverslips. Slides were stored at RT.

Imaging was performed on a Zeiss AX10 microscope using StereoInvestigator Software with a 5x/0.16 (420630-9900) apochromat air objective. Quantification was performed using StereoInvestigator Software "Optical Fractioner Workflow" probe with the following settings. Thickness of 50μm was manually defined and regions were selected using a 1.25x/0.03 (420310-9900) apochromat air objective for low magnification and then counted with a 10x/0.45 (420640-9900) apochromat air objective for high magnification. The border between medial septum (MS) and diagonal band of Broca (DB) was defined as a line between the two major island of Caleja. The regions were separated using different lines. The counting frame used was a square of 75μm size and the grid was a square of 150μm size. The counter was blind to genotype.

Oral administration of Donepezil

Donepezil hydrochloride (Sigma Aldrich, D6821) was orally administered in a jelly. Mice were trained to voluntarily consume jelly following the protocol described by Zhang (2011). Mice were given placebo jelly or a jelly containing a Donepezil dose of 1.8mg/kg. For experiments studying the effects of Donepezil on acetylcholinesterase (AChE) activity, jelly was given at 8am daily. For experiments studying the effects of Donepezil on IIS, jelly was given daily at either 8am, or 8pm to assess interactions of AChE modulation and circadian cycle. Since there was no effect of AChE on IIS, results were pooled.
Acetylcholinestrase (AChE) assay

Quantitative measurements of AChE enzymic activity were made using a modified Ellman method (Ellman et al., 1961; Rosenfeld et al., 2001). Stock solutions were acetylthiocholine iodide, used as the enzymic substrate (ATH; 1.7 mg/ml in PBS, Sigma-Aldrich), 5,5'-dithio-bis(2-nitrobenzoic acid) (DTNB, 0.8 mg/ml in PBS, Sigma-Aldrich). Briefly, brains were rapidly dissected from either WT or J20 mice. Neocortex was isolated, weighed and then homogenised using a Pellet Pestle (Sigma, Z 359971) in 9 volumes of 0.1M sodium phosphate buffer (pH 7.4) (Patel et al., 2014). 5 μl of brain homogenate was aliquoted into each well of a 96–well plate, volume made up to 200 μl with PBS. DTNB (50 μl from stock) was added, followed by 50 μl of ATH substrate from stock. Measurement of absorption at 450 nm began immediately (<2 hours from dissection) and was measured every 5 mins for up to 30 mins using a MRX microplate reader (Dynex Technologies, Chantilly, USA). Thiocholine production in the test wells was expressed in units of nmoles/minute, calibrated with reference to the absorbance change over a range of concentrations giving a linear response using glutathione as the DTNB reactant (Eyer et al., 2003). Neostigmine (10 μM, Sigma-Aldrich) was used to completely inhibit AChE activity and establish there was no baseline drift during the measurements.

Human scalp EEG and foramen ovale (FO) electrode recordings

Human scalp EEG and FO electrode recordings were performed at the Massachusetts General Hospital, as described in detail previously (Lam et al., 2017). Scalp EEG electrodes were placed using the International 10-20 system, with additional T1 and T2 electrodes.
Sleep staging in patient data was performed by a board-certified clinical neurophysiologist (ADL) based on visual analysis of the full scalp EEG data. While dedicated electrooculogram and electromyogram channels were not recorded for these studies, the frontopolar scalp EEG electrodes allowed assessment of eye movements, while the frontopolar, frontal, and temporal electrodes allowed assessment of myogenic activity. Scalp EEG data was reviewed in 30 second epochs in the longitudinal anterior-posterior bipolar montage, using the Python module wonambi (https://github.com/wonambi-python/wonambi). Each 30 second epoch was classified as awake, NREM1, NREM2, NREM3, or REM, based on the American Academy for Sleep Medicine's manual for sleep scoring.

Spike quantification in patient data was performed by a board-certified clinical neurophysiologist (ADL), using a custom-made GUI in Matlab (Mathworks, Natick, MA). The GUI displayed 15 second epochs of left and right sided FO data, in both bipolar and common reference montages (common reference = C2), along with the EKG trace to allow exclusion of EKG artifact. The reviewer could adjust amplitudes for each trace as needed. For the MCI patient analyzed, contact #3 from the left FO electrode did not record properly and was excluded from analysis. The reviewer marked all spikes in each epoch. Epochs were presented in consecutive order, but the reviewer was otherwise blinded to the sleep stage for each epoch during the review. Instantaneous spike rates were calculated by determining the total number of left FO and right FO spikes detected within all 30 second epochs of the recording (which corresponded to
the sleep staging epochs above), and converting these rates to spikes per hour. Average spike rates within each sleep stage were calculated by summing the total number of spikes that occurred during each sleep stage and dividing by the total number of hours the patient spent in each respective sleep stage in the recording.

Spectral analysis of the FO electrodes was performed in Matlab, using the freely available Chronux toolbox (Mitra and Bokil, 2007). Analysis was performed on the LF01, LF02, RF01, and RF02 channels, as these were the deepest contacts and thus least prone to noise or artifact. Channels were each normalized to zero-mean, unit-variance. Multi-taper spectrograms were calculated for each normalized channel, using the Chronux script `mtspecgramc` with the following parameters: frequency range: 1-20Hz, window: 30 seconds; step size: 30 seconds; time-bandwidth product: 3, tapers: 5. This provided a spectral resolution of 0.2Hz. An average spectrogram across all FO channels was then generated, and the average spectral powers within the δ band (0-4Hz) and θ band (4-12Hz) were then calculated.

Statistics

Statistical data analysis was performed using R (version 3.2.0) including the 'dplyr' (Wickham et al., n.d.) and ggplot2 (Wickham, 2009) packages. Assumptions for parametric tests were tested using Q-Q plots and residual plots. Data transformations or non-parametric tests were used for two-group comparisons in which test assumptions were violated.
For evaluating the effects of the fixed effects of age and genotype on the proportion of intervals containing more than one spike in APPNL/F animals, the data first underwent a square-root transformation and then fit using a linear model:

$$\sqrt{\text{IntervalProportion}} \sim \text{Age} + \text{Genotype} + \xi$$

where $\xi$ is the error term.

The time of IIS was treated as circular variable. Each interval in which 1 or more IIS were detected was considered an event. The time of each event was evaluated as a phase of a circadian cycle. Circular data was analysed using circular statistics by means of the ‘circular’ package (Agostinelli and Lund, 2013). Circular outliers were identified using ‘CircOutlier’ package (Rambli et al., 2016).

For tests entailing random variables, linear models were fit using ‘lme4’(Bates et al., 2015). Significance was tested using a log-likelihood test comparing the full model to a null model without the factor of interest.

For evaluation of the relationship between spike count and $\theta/\delta$, we described each $\theta/\delta$ value as a member of one of three levels: i) $\theta/\delta \leq 1$; ii) $1 < \theta/\delta \leq 2$, and iii) $\theta/\delta > 2$. We then modelled spike count (Poisson-distributed) as a function of levels of $\theta/\delta$, using the R package ‘MCMCglmm’ (Hadfield, 2010). It should be noted that due to poor properties of a single model fitted across all animals
(fitting animal as a random effect and $\theta/\delta$ factor as a fixed effect), separate models were fitted to individual animals without including a random effect. Thus, the data do not allow for inference about the population.

Event-triggered averages of IIS were evaluated by considering each interval in which an IIS was detected as an event. If no intervals within $\pm 80s$ around the event were excluded, then the 160s window was included in the calculation of the event-triggered averages, else the event was excluded from the averaging. An event-triggered average was also evaluated around 2000 randomly sampled points.

For comparing $\theta/\delta$ in intervals with IIS to $\theta/\delta$ in intervals preceding IIS, we considered only interval pairs where the preceding interval did not contain IIS and fit the model:

$$\left(\frac{\theta}{\delta}\right)^{1/4} \sim \text{Index} + \text{Subject} + \xi$$

where Index was a factor labelling whether the interval contained IIS or the preceding interval and modelled as a fixed effect, and Subject was a random effect with a random intercept.

For comparison of ChAT+ cells between genotypes, the model used was:

$$\text{EstimatedCount} \sim \text{Genotype} + \text{Region} + \text{Subject} + \xi$$

where genotype and region were fixed effects and subject was a random effect with a random intercept.
To study the effect of genotype and treatment of the Thiocholine production rate, the data of Thiocholine production was log-transformed. The model used was

$$\log(\text{ThiocholineRate}) \sim \text{GenotypeTreatment} + \text{RepeatID} + \xi$$

where GenotypeTreatment was a fixed effect and RepeatID was a random effect with a random intercept. Post-hoc tests for the linear model were performed using package ‘multcomp’ with the Holm correction method (Hothorn et al., 2008). It should be noted that while the treatment levels of control and donepezil were independent, the neostigmine treatment was applied to a sample of wild-type control tissue and thus was not independent. This repeated factor was not accounted for in the model.

Significance was tested using $\alpha=0.05$. Two-sided hypothesis testing was used.

Superscripts following statistical reporting in the results section refer to the statistical table (Table 1).

Code and data accessibility

The processor script used for quantification of IIS, $\theta$ and $\delta$ power in rodent ECoG data is available from

http://www.opensourceinstruments.com/Electronics/A3018/HTML/SCPP4V1.tcl

Code used for quantifying IIS in human data is available from

https://github.com/mauriceaj/GUI-EEG_Spike.Annotation
The datasets used for figures 1 – 6 (rodent data) are available from http://dx.doi.org/10.7488/ds/2319

Results

Network hyperexcitability in mouse models of AD pathology

To establish circadian patterns of network hyperexcitability in J20 mice, we recorded ECoG activity from freely-moving J20 and littermate wild-type (WT) mice using wireless telemetry over a period of three days. As network excitability has been suggested to be an early event in AD pathogenesis (Vossel et al., 2013; Sarkis et al., 2015), we focused our study on ages which precede overt plaque pathology in J20s (Mucke et al., 2000).

As previously reported (Palop et al., 2007), non-seizure, inter-ictal spikes (IIS; Fig 1A) were detected in J20 ECoG (note that while ictal activity was not assessed, we refer to these as inter-ictal events due to the similarity with IIS that have been reported in the literature). We applied automated event detection (see methods), on 8s intervals of continuous ECoG. The percentage of intervals in which 1 or more spikes were detected was negligible in WTs (mean percentage: 0.8%, sd=0.7%, n=8). In contrast, the percentage of intervals with 1 or more spikes was greater in J20s (mean percentage: 11.6%, sd=5.1%, n=18; t=10.6, df=23.98, p<0.0001, t-test on square root transformed data with Welch correction, Figure 1B,C)².

Seizures and IIS have been reported in numerous strains of transgenic mice that express hAPP and that exhibit Aβ pathology (Del Vecchio et al., 2004; Palop et al.,...
2007; Minkeviciene et al., 2009; Rasch and Born, 2013). However, it has been suggested that such network hyperexcitability is the result of overexpression of hAPP (Born et al., 2014). To determine whether network hyperexcitability is associated with Aβ pathology in the absence of hAPP overexpression, we performed telemetric ECoG recordings as above, in mice expressing the humanized Aβ sequence of APP (APPNL/F; Saito et al., 2014) and age-matched controls. We recorded from mice at ages preceding overt plaque pathology (8 months) and at ages where plaques begin to appear (12 months)(Saito et al., 2014; Masuda et al., 2016). We found no significant effect of genotype in the proportion of intervals containing spikes between WT and APPNL/F (Fig 1D; F(2, 32)=3.1, R²=0.11, p=0.06) with a negligible proportion of intervals with one or more spikes detected (mean percentage of intervals with one or more spikes, pooled across genotype and age = 1.2%, 95%CI (0.8%, 1.6%)). A post-hoc power calculation based on the effect size from the J20 group (Cohen's d=2.5) and the sample sizes of the APPNL/F and WT groups yielded a power of >0.99 at α=0.05 for an effect of genotype. Hence, we conclude that APPNL/F mice show no evidence of network hyperexcitability compared to control animals.

Circadian coupling of IIS

It has been suggested that seizure-related activity shows circadian fluctuations in epilepsies (Quigg, 2000). Hence we next asked whether the likelihood of IIS in J20s varies across the day/night cycle. Quantifying the number of IIS per hour revealed that IIS are more frequent during daylight hours (inactive phase; Fig 2A). We used circular statistics to extract measures of the phase coupling of IIS to the circadian cycle within individual J20 animals (Fig 2A, see methods). To
evaluate the degree of phase coupling of IIS in each animal, we evaluated the mean angular vector length ($\rho$) from the time of IIS. $\rho$ can vary between 0 (no phase coupling) to 1 (perfect phase coupling). To evaluate the time to which IIS were coupled, we extracted the mean coupling phase off IIS, expressed as a time on a 24-hour cycle ($\theta_{IIS}$).

The distribution of IIS phases differed significantly from a random distribution in all animals (Rayleigh Test of Uniformity: $p<10^{-11}$). The extent of phase coupling was variable across the sample of J20s (Fig 2B; mean $\rho_{IIS}=0.24$, sd=0.13, n=16).

Evaluating the coupling phase revealed that IIS occurred predominantly in the light condition (Fig 2A). Across the sample of J20s, the mean $\theta_{IIS}$ confirmed this (Fig 2B; $\bar{\theta}_{IIS} = 15h05$, $\rho =0.38$, n=16, $p < 0.0001$, Rayleigh's test).

Inspection of the $\theta_{IIS}$ distribution revealed potential outliers. Testing for outliers on a circular distribution (Rambli et al., 2016) identified four outliers. These four animals were amongst the 5 that showed a cluster of weakest phase coupling as measured by $\rho_{IIS}$ (range: 0.06 – 0.11). We used the upper bound of the range of $\rho_{IIS}$ of the four outlier animals to classify phase-coupling as weak or strong.

Henceforth, we refer to the five animals with $\rho_{IIS} \leq 0.11$ as showing weak phase-coupling and the other 11 animals as showing strong phase-coupling ($\rho_{IIS}>0.17$).

Sleep/wake modulation of IIS
Since IIS predominantly occurred in the normal inactive phase of the circadian cycle, we next asked whether this circadian modulation of IIS could be accounted
for by the sleep/wake state of the animals. In a subset of J20s, we acquired simultaneous video recordings while recording ECoG data (n=4). We manually scored the video and classified periods as ‘sleep’ or ‘wake’ (see Methods). Two of these four J20 animals showed strong circadian phase-coupling of IIS, and two showed weak phase-coupling. For the two animals that showed strong phase-coupling of IIS, IIS occurred more frequently in sleep than during waking (Fig 3A, B). In contrast, the modulation of IIS probability did not show a consistent pattern in animals showing weak phase-coupling (Fig 3B). This suggests that the strong phase-coupling of IIS may be accounted for by differences in behavioural state across the circadian cycle.

Brain state modulation of IIS in J20 mice

Sleep-related ictal and inter-ictal activity is differentially modulated by REM and NREM sleep in different forms of epilepsy (Bazil and Walczak, 1997; Herman et al., 2001; Sedigh-Sarvestani et al., 2014; Ewell et al., 2015). REM and NREM can be distinguished by the relative power in the \( \delta \) (defined here as 0.1 – 3.9 Hz) and \( \theta \) (4-12 Hz) frequency bands, with high \( \theta/\delta \) associated with REM (Ewell et al., 2015) as well as waking exploration (Buzsáki, 2002). Thus, we next asked whether IIS are more likely to occur in particular brain states. To this end, we performed spectral analysis of the ECoG data from a subset of the mice (n=5 J20s) in which a reference electrode was implanted at cerebellar coordinates (a non-cortical reference for detection of cortical rhythms). ECoG recordings from J20 mice, exhibited periods showing a peak in \( \theta \)-band power when animals were either awake (i.e. moving) or asleep, while periods of elevated \( \delta \)-band power were seen during sleep (Fig 4A). We evaluated the \( \theta/\delta \) ratio for each 8s interval
and related it to the number of IIS in the interval. Transient increases in $\theta/\delta$ were observed during sleep and were associated with increased occurrences of IIS (Fig 4B).

To quantify whether IIS were more likely in particular brain states, we next investigated the relationship between $\theta/\delta$ and IIS count/8s interval. As we were interested in discriminating between REM and NREM sleep, we limited the analysis to daylight hours when animals are more likely to be asleep. We used a value of $\theta/\delta<1$ and $>2$ to classify periods as NREM-like and REM-like respectively (Ewell et al., 2015). This revealed significantly higher spike counts during REM-like vs NREM-like periods in all 5 animals (Fig 4B, p<0.0005 for all 5 animals, Markov Chain Monte Carlo generalised linear model). Interestingly, IIS were associated with increased $\theta/\delta$ in animals showing both weak and strong phase-coupling (Fig 4C). Since sleep and wake are not predictive of IIS in animals with weak phase-coupling, this suggests that there is a mismatch between $\theta/\delta$ and behavioural state in animals with weak phase-coupling. Moreover, high $\theta/\delta$ states are predictive of IIS, regardless of behavioural state.

To examine the temporal dynamics of $\theta/\delta$ around IIS, we evaluated the IIS-triggered average of $\theta/\delta$ (Sedigh-Sarvestani et al., 2014) for 160s window around each interval in which at least one IIS was identified. In all animals, $\theta/\delta$ was increased around the time of IIS relative to $\theta/\delta$ averaged around randomly sampled points (Fig 4D). In three strongly phase-coupled animals, $\theta/\delta$ returned to baseline levels within the 160s window around the event. However, in the
weakly phase coupled animals, \(\theta/\delta\) remained elevated above baseline levels in this window. The peak in the \(\theta/\delta\) IIS-triggered average did not occur at \(t=0\) in any of the animals. Since intervals neighbouring the IIS-containing interval show increased \(\theta/\delta\), this suggests that the IIS contribution to spectral power did not underlie the association between increases in \(\theta/\delta\) and IIS probability. To further examine whether IIS could directly contribute to the increased \(\theta/\delta\), we compared \(\theta/\delta\) in intervals with IIS to \(\theta/\delta\) in the preceding intervals only in cases where the preceding interval contained no IIS. We found no significant difference in \(\theta/\delta\) between intervals with IIS and the preceding interval \(\text{(Linear mixed model, } \chi^2(1) = 0.35, p=0.56, \text{ data not shown})\).

To determine whether the spectral ECoG patterns in J20 mice are a reflection of normal sleep or a result of pathology, we performed similar analysis of video-scored ECoG data from 3 wild-type mice. As in the J20, intervals of strong \(\delta\)-band activity were seen in sleep. Transient increases in \(\theta/\delta\) during sleep akin to those seen in J20s were also observed in all WT animals, suggesting that such increases are a feature of normal sleep, and not pathological \(\text{(Fig 5)}\). To compare the distribution of \(\theta/\delta\) during sleep between genotypes, we calculated the range and 90th percentile of \(\theta/\delta\) while animals were asleep \(\text{(using data for which we had video-scoring)}\). Group sizes were too small for statistical comparison but suggested that \(\theta/\delta\) values spanned a narrower range in J20 mice than in WT mice \(\text{(J20 mean range=(0.02, 10.0), 90}^{\text{th}} \text{ percentile=2.4, SD(1.1), n=4; WT mean range=(0.04, 19.3), 90}^{\text{th}} \text{ percentile =5.4, SD(1.4), n=3; data not shown).}\)
No evidence of cholinergic changes in J20 mice

Cholinergic levels exhibit a circadian modulation (Hut and Van der Zee, 2011), and high cholinergic tone is implicated in generating θ oscillatory states (Buzsáki, 2002). In addition, cholinergic dysfunction has been suggested to be a key feature of AD pathogenesis (Craig et al., 2011). Recently, it has been suggested that cholinergic alterations may contribute to network excitability in the Tg2576 model of AD (Kam et al., 2016). Hence, we hypothesized that cholinergic changes might underlie the brain-state dependent modulation of IIS in the J20 mice. We used immunohistochemistry to quantify the number of ChAT+ cells in the MS and DB, and asked whether the number of ChAT+ cells differs between J20 (n=7) and WT (n=5) mice. Fitting a linear mixed model to the data, we found no effect of genotype on the estimated number of ChAT+ cells in the MS or DB (Fig 6A; Linear mixed model, $\chi^2(1) = 0.0002$, $p = 0.99$).

AChE activity is reduced in AD (García-Ayllón et al., 2011). We assayed cholinergic function by measuring AChE activity. AChE activity was quantified by estimating the rate of thiocholine production in neocortical brain homogenates (see methods). There was no significant difference in the rate of thiocholine production in brain homogenates prepared from WT and J20 mice ($V = 15$, $p = 0.06$, $n = 5$ WT/J20, Wilcoxon signed rank test, matched by day of assay, Fig 6B). We also wanted to directly test the effect of modulation of ACh levels on IIS. However, using oral administration of Donepezil at a dose previously suggested to achieve clinically relevant drug plasma levels (Dong et al., 2009) was ineffective at altering AChE activity in brain homogenates. In contrast, a positive
control treatment of direct application of neostigmine to brain homogenate led to a significant reduction in AChE activity (Fig 6B; Linear mixed model: $\chi^2(4)=75.3, p<0.0001$. Post-hoc using Tukey paired comparisons: $p<0.0001$ for neostigmine vs. each of the treatment and genotypes. $p>0.05$ for all other group comparisons). Two days of oral Donepezil administration at this dose did not affect the IIS rate in J20 mice ($t(11)=0.8, p=0.43$, paired t-test, data not shown).

Sleep stage modulation of IIS in human AD

The first intracranial recordings in humans with AD were recently reported and demonstrated marked activation of mesial temporal lobe (mTL) IIS during sleep compared to the awake state (Lam et al., 2017). We further analyzed the combined scalp EEG and intracranial electrode recordings from these two patients to better understand the relationship between sleep stage and mTL IIS rate in AD patients. One patient with advanced AD did not achieve REM sleep but showed mTL IIS preferentially during NREM sleep as opposed to waking states (Table 2, Patient 1). The second patient was a 67-year-old woman with amnestic mild cognitive impairment (aMCI), an early stage of AD that is thought to correspond to the early stage of AD modelled in our young J20 mice. The data from this patient was used to compare the frequency of IIS in wake, NREM and REM states.

We analyzed 14.25 consecutive hours of combined scalp EEG and FO recordings from the aMCI patient, which spanned from ~7PM on the first day of FO recording (FOD1) to 9:15AM the following morning (FOD2). Further recordings were not analyzed, as the patient was initiated on treatment with the
anticonvulsant levetiracetam on the afternoon on FOD2. Of note, the patient underwent implantation with FO electrodes on FOD1 from ~ 12:40PM – 1:50 PM and received sevoflurane, propofol and midazolam during the procedure. She was awake and answering questions appropriately by 2:15PM on FOD1.

We performed sleep staging of the recording using the full scalp EEG data, and measured mTL spike rates using the bilateral FO electrode data (Figure 7A,B). As described previously, we found that mTL spiking in the aMCI patient was largely activated during sleep. In contrast to what we found in the young J20 mice, mTL spiking in the aMCI patient occurred with highest frequency during NREM sleep stages, particularly during NREM3, and were lowest during REM sleep (Figure 7 and Table 2). mTL IIS rates during REM sleep were also markedly lower than during wakefulness (Table 2). We also calculated spectral power in the $\theta$ and $\delta$ bands, as well as the $\theta/\delta$ ratio, in the FO electrodes across sleep states (Figure 7C-E). Increases in both $\theta$ and $\delta$ power were seen with deepening stages of NREM sleep, while a reduction was seen with REM sleep. In contrast to what we observed in the J20 mice, the $\theta/\delta$ ratio was reduced during periods of highest spike frequency (Figure 7E).

**Discussion**

Network hyperexcitability is a feature of AD. Here we compared patterns of network hyperexcitability in two rodent models of AD, as well as in two AD patients, in order to reveal shared phenomenological features with the disease. We show that while J20, (hAPP overexpressing) mice exhibit frequent IIS as
previously reported, APPNL/F mice (that express APP at physiological levels) do not show evidence of network hyperexcitability. Moreover, IIS in J20s occur primarily during daylight hours, and this circadian fluctuation is accounted for by an increased probability of IIS during sleep. Interestingly, we found that IIS in J20 mice are modulated by brain state, with increased likelihood of IIS in brain states with high \( \theta/\delta \) activity, a marker of REM sleep. In contrast, patients with AD showed prevalent IIS during NREM sleep. Moreover, in the one AD patient who exhibited REM sleep, IIS frequency was lowest in REM compared to other sleep states.

Circadian dysfunction and network hyperexcitability in AD

Brain network hyperexcitability in the form of IIS and seizures has now been reported in numerous models of AD pathology (reviewed in Scharfman, 2012; Born, 2015). Our data, along with those reported by others (Born et al., 2014; Kam et al., 2016) reveal that network hyperexcitability in animals models of AD can be modulated by the circadian cycle. Circadian disturbances in AD include sleep fragmentation, increased daytime somnolence, and sundowning, the phenomenon in which neuropsychiatric symptoms are heightened late in the day (Peter-Derex et al., 2015). Animal models of AD have also been reported to show disturbances in the circadian cycle, some of which overlap with patterns of circadian alterations seen in patients (Huitrón-Reséndiz et al., 2002; Vloeberghs et al., 2004; Wisor et al., 2005; Jyoti et al., 2010; Sterniczuk et al., 2010; Duncan et al., 2012; Roh et al., 2012). Our findings of circadian modulation of network hyperexcitability in AD raise the question of whether IIS might causally contribute to the alterations in circadian-coupled behaviour observed in AD.
Future work investigating the effects of anti-epileptic drugs on circadian alterations in AD would go towards answering this.

**Brain state modulation of network excitability**

Here we report that IIS in J20 animals are modulated by $\theta/\delta$, with higher IIS rates seen in states of high $\theta/\delta$ during sleep. The spectral patterns of EEG that we report here are in line with previous reports in WT mice, that have shown increases in cortical EEG $\theta$ power in REM sleep relative to wake and NREM (Brankačk et al., 2010). We also report transient increases in $\theta/\delta$ in sleep in both WT and J20 mice. Since these increases in $\theta/\delta$ occur in both WT and J20s, they are likely to be indicative of REM sleep periods (Ewell et al., 2015). Given that J20 animals with strong circadian phase-coupling show highest IIS rates during sleep this suggests that IIS in these animals are associated with REM sleep.

An alternative explanation for the association between IIS and high $\theta/\delta$ during sleep may be that IIS occur during ectopic $\theta$ in sleep, in the absence of a concomitant drop in muscle tonus. A phenomenon of ictal activity during ectopic $\theta$ has been reported in a mouse model of Huntington’s disease (Pignatelli et al., 2012). Without simultaneous EMG recordings, the present data cannot conclusively distinguish between REM states and ectopic $\theta$. In the human data, analysis of $\theta/\delta$ ratios showed that these were lowest during periods of highest IIS frequency. This argues against the idea of IIS coupled to ectopic $\theta$ in humans, though a more definitive assessment will require data from more AD subjects as well as healthy elderly controls.
Our finding of an association between IIS and high $\theta/\delta$ is in line with recent reports that young Tg2576 model of AD as well as mice overexpressing WT-hAPP also demonstrate IIS predominantly during states of high $\theta$ which the authors suggest is indicative of REM sleep (Kam et al., 2016).

The findings that IIS in multiple mouse models of AD are most likely to occur in REM-like states begs the question of what makes REM a pro-ictal state in these models. Both REM sleep and the awake state share common features of high $\theta/\delta$ activity and high cholinergic tone (Vazquez and Baghdoyan, 2001; Lee et al., 2005), yet IIS occur much less frequently in the awake state in these models. There are several potential explanations for this. Firing rates of hippocampal neurons increase during REM (Grosmark et al., 2012), which might contribute to the propensity to seize. In addition, systems that normally show distinct activity in REM sleep vs. waking and NREM sleep might contribute to the pro-ictal REM state in these models (Sedigh-Sarvestani et al., 2014; Ewell et al., 2015; Kam et al., 2016). Unlike cholinergic neurons, which increase their activity in both REM and waking, monoaminergic neurons in brainstem nuclei (including the locus coeruleus and the tuberomammillary nucleus) as well as the dorsal raphe nucleus of the hypothalamus, show differential activity between these brain states. These neurons are highly active in waking, exhibit low firing rates in NREM sleep, and are quiescent during REM sleep (Lee and Dan, 2012). It may be that brain state modulation of one or more of these systems is disrupted in these mouse AD models, and other forms of epilepsy which show REM-coupling (Sedigh-Sarvestani et al., 2014; Ewell et al., 2015).

The present study quantified cholinergic neurons in MS and DB. Cholinergic
neurons in laterodorsal tegmental and pedunculopontine tegmental nuclei of the pontomesencephalic tegmentum have been suggested to control REM onset (Van Dort et al., 2015). In the rat, these neurons have been shown to be active during both wake and REM, however, firing rates are higher in REM, and correlate with $\theta/\delta$ (Boucetta et al., 2014). Thus, changes to these neurons are also potential candidates for mediating the pro-ictal nature of REM sleep in J20 mice.

Kam et al.,(2016) reported that MS-DB cholinergic neuron number was unchanged in young Tg2576 mice. However, they found evidence to support the notion that overactivity of cholinergic neurons might contribute to IIS by showing that antagonism of muscarinic receptors reduced IIS in these animals. Hence they concluded that IIS during REM might be the result of cholinergic hyperfunction. We did not find evidence for cholinergic changes in J20 mice as quantified by the number of cholinergic neurons in MS-DB, or AChE activity. If cholinergic activity is indeed unaltered in J20 mice, future experiments using muscarinic antagonism in J20 mice could be used to investigate whether atropine can act to reduce IIS by reducing overall neuronal excitability, rather than by reversing cholinergic hyperfunction.

Our assay of cholinergic function was based on measurements of AChE enzymic activity in brain homogenate. There was no significant difference between AChE levels in WT and J20, or with donepezil treatment. While it is possible that post-mortem degradation of AChE could have masked differences in AChE levels, the robust effect of neostigmine supports the conclusion that the tissue contained functional AChEs.
In a subset of our animals, IIS were weakly-coupled to the circadian cycle and the sleep-wake pattern, but were still modulated by $\theta/\delta$. This suggests that the relationship between $\theta/\delta$ and behavioural state might be disturbed in these animals. It is possible that these animals also exhibited greater disturbances in other elements of the circadian-cycle, such as a circadian decoupling of sleep quantity/quality.

During both REM and NREM, hippocampal neurons have been shown to replay firing patterns that were experienced prior to sleep (Skaggs and McNaughton, 1996; Louie and Wilson, 2001), and such precisely timed sequences are likely to be involved in the memory facilitation role of sleep. IIS are thought to arise from depolarization and synchronous firing of neurons. This firing is followed by an inhibition and reduction of firing (Holmes and Lenck-Santini, 2006). Thus, IIS during sleep are likely to interfere with the coordinated replay of firing sequences, and consequently, would be expected to contribute to memory impairments. In support of this, it has recently been shown that reducing IIS by treatment with anti-epileptic drugs, rescues memory deficits in J20s (Sanchez et al., 2012).

Relationship between IIS and AD pathology in mouse models

Here we report that while IIS are prevalent in hAPP overexpressing mice, APPNL/F mice that exhibit A$\beta$ pathology without APP overexpression, do not exhibit IIS at two ages preceding widespread plaque deposition (8 and 12 months). This finding is in line with other reports that it is overexpression of hAPP that is causal in generating network hyperexcitability in these animal models.
models (Born et al., 2014; Xu et al., 2015; Kam et al., 2016). An alternative explanation of the presence of IIS in J20 but not APPNL/F mice may be differences in the levels of Aβ between the two models. However, levels of soluble Aβ in 6 month old J20 and 12 month old APPNL/F are comparable, and levels of total Aβ are higher in APPNL/F (Shankar et al., 2009; Saito et al., 2014). Thus it is unlikely that higher levels of Aβ in the J20s are a cause of IIS in this model.

Interestingly, APPNL/F mice begin to exhibit cognitive deficits at 8 months of age (Masuda et al., 2016), which suggests that cognitive deficits at these ages are not the result of IIS, as has been suggested for J20s (Sanchez et al., 2012). Moreover, differences in the types of memory affected in J20 and APPNL/F at ages preceding overt plaque deposition have been reported. Specifically, 4 – 6 month old J20s show impairments in hippocampal dependent spatial memory (Sanchez et al., 2012). In contrast, in 8-month old APPNL/F mice, spatial memory as assayed by a place preference task is intact. However, place-avoidance memory, which is also dependent on amygdala-circuits (Wilensky et al., 2000), is impaired (Masuda et al., 2016). It may be that hippocampus dependent processes are susceptible to interference by IIS while the disturbances in the non-hippocampal circuits result from processes independent of IIS.

Differential sleep-stage coupling between mouse models of AD and human AD

Lam et al., (2017) recently used intracranial electrode recordings to detect mTL IIS in two AD patients without a history of epilepsy. Here, we report that in these patients, IIS were predominantly associated with NREM sleep (ie. low θ/δ). In the patient with aMCI, IIS occurred most frequently in N3 sleep and were least
frequent in REM, with a greater than 4.5-fold difference in spike rates between N3 and REM. In the AD patient, frequent IIS were seen during NREM sleep, though REM sleep was absent from this patient’s brief recording, in line with previous reports of REM deficits in AD (Vitiello et al., 1984). Our findings from intracranial electrodes in AD patients are consistent with prior scalp EEG studies by Vossel et al., (2016), who reported that epileptiform discharges are highly prevalent in sleep stages >2 (although the authors did not differentiate between REM and NREM sleep). Although the means of characterising sleep differed between rodents and patients, combined, these results point to important differences in sleep stage coupling of epileptiform activity between rodent AD models and humans with AD and suggest that the specific mechanisms that underlie hyperexcitability in AD may differ between certain mouse models and humans.

Analysis of ictal and inter-ictal activity in epilepsy patients has led the view that NREM sleep is a generally pro-ictal state, whereas REM sleep is an anti-ictal state (Sammaritano et al., 1991; Herman et al., 2001; Minecan et al., 2002; Ng and Pavlova, 2013). Many animal models of epilepsy have also shown that seizures are more frequent in NREM and rarely occur in REM (Shouse et al., 2000). Interestingly, rodent models of the same type of epilepsy can still exhibit differences in the sleep-stage coupling of epileptiform activity. For example, in both the kindling as well as the pilocarpine models of temporal lobe epilepsy in rats, IIS are most common during NREM sleep (Colom et al., 2006; Gelinas et al., 2016). In contrast, rats with either the tetanus toxin or the low-dose kainate models of temporal lobe epilepsy have seizures that occur most commonly
during REM sleep (Sedigh-Sarvestani et al., 2014; Ewell et al., 2015). Based on this, we hypothesize that different mouse models of AD may have specific mechanisms underlying their network hyperexcitability, which could be differentially expressed through sleep-stage coupling of IIS. We propose that sleep-stage coupling of IIS should be an important factor for identifying mouse AD models that more closely resemble the EEG signature of network hyperexcitability in human AD.
**Figures**

**Figure 1:** Inter-ictal spikes (IIS) are prevalent in J20 mice, but not in APP knock-in mice. (A) ECoG trace recorded from a J20 mouse showing IIS. Inset is 250ms expansion around IIS event marked by *. (B) Empirical cumulative distribution frequency plots for individual animals quantifying the number of detected IIS in 8s intervals across 3 days of recording in WT and J20s. Colours represent distributions for individual animals. (C) Plot showing the proportion of intervals with one or more detected IIS in WT and J20. (D) Plot showing the proportion of intervals with one or more detected IIS in WT and APPNL/F at 8 months and 12 months. Bars represent medians. Whiskers extend to 1.5 IQR. *** p<0.001.

**Figure 2:** Circadian modulation of IIS. (A) Circular histogram of IIS counts over three days of recording in an individual J20 mouse plotted on 24hr cycle. Light condition indicated by shading. For the animal shown, $\psi_{IIS} = 14h38$ and $\rho=0.43$. (B) Summary data for $\psi_{IIS}$ vs $\rho$ for all animals, shown on circular plot. Solid symbols are strongly-coupled animals. Weakly coupled animals are shown with orange fill.

**Figure 3:** The probability of IIS is modulated by behavioural state in strongly phase-coupled animals. (A) IIS count/8s interval versus time over 2 hours of ECoG recording in a J20 mouse, with sleep and wake indicated by shading. (Bi) Mean spike rate in sleep and wake condition for strongly and weakly phase coupled animals. Error bars: 95% CI. (ii) Circular histograms for a strongly (left) and weakly (right) phase coupled animals using conventions as in Fig 2A.
Figure 4: IIS occur during high $\theta/\delta$ states. (A) 8s ECoG signals (left) and corresponding power spectra (right) during different behavioural states recorded from a J20 mouse. A single IIS is seen in the sleep high $\theta$ state (ii). (B) Time series of $\delta$ power, $\theta$ power, $\theta/\delta$ and spike count per 8s intervals across 2 hours of ECoG recorded from the same J20 mouse as shown in A. Black/grey symbols indicate sleep/wake as classified by simultaneous video data. Red symbols and vertical dotted lines indicate the 8s intervals for which the ECoG signal is shown in panel A. (C) Spike number per 8s interval as a function of $\theta/\delta$ in 5 animals (represented by different colours and connected by lines). The increase spike count in intervals with high $\theta/\delta$ was seen in animals with both strong (filled symbols) and weak (open symbols) circadian phase-coupling. *** $p<0.001$ (D) IIS-triggered averages of $\theta/\delta$ for 5 individual animals (black) and windowed averages triggered around 2000 randomly sampled points (grey) show an increased $\theta/\delta$ around IIS. Strong/weak coupling shown in filled/open symbols. Error bars in B and C represent 95% CI.

Figure 5. Transient increases in $\theta/\delta$ are non-pathological features of sleep. (A) 8s ECoG signals (left) and corresponding power spectra (right) during different behavioural states recorded from a WT mouse. (B) Time series of $\delta$ power, $\theta$ power and $\theta/\delta$ per 8s interval across 2 hours of ECoG recorded from same WT mouse as shown in A. Black/grey symbols indicate sleep/wake as classified by simultaneous video data. Red symbols and vertical dotted lines indicate the 8s intervals for which the ECoG signal is shown in panel A.
Figure 6: No evidence of cholinergic alterations in J20s. (A) Immunostained brain section showing ChAT+ cells in medial septum (MS) and diagonal band of Broca (DB). Lower panel shows zoomed in region of upper panel (left) and corresponding regions of a negative control stained section (right). Upper right: quantification of stereological estimates of ChAT+ cell count in MS and DB in WT and J20. (B) AChE activity was assayed by the rate of thiocholine production in brain homogenate from WT and J20 in control conditions and following oral administration of donepezil (DPZ). The AChE activity was compared to a positive control of direct application of neostigmine (10µM) to the brain homogenate. Experimental repeat groups are indicated by different colours and connected lines. *** p<0.001.

Figure 7: Sleep stage coupling of mesial temporal lobe (mTL) spiking in a human with aMCI, a suspected early stage of AD. (A) Hypnogram showing the patient's sleep architecture, spanning from ~ 7PM on FOD1 to 9:15AM on FOD2. (B) Bar plot showing instantaneous mTL lobe spike rates over the course of the recording. Bars are colored by sleep stage, with light green for Wake, light blue for NREM (includes NREM1, NREM2, and NREM3), and dark blue for REM. The patient had three brief subclinical seizures (SZ) from the left FO electrodes during this recording, the timing of which is depicted by red vertical bars. (C-E): Plots showing (C) δ power (0-4Hz), (D) θ power (4-12Hz), and (E) θ/δ ratio of bilateral mTL activity, based on FO electrodes recordings. Dots represent the spectral power for each non-overlapping 30 second window of the recording. Power is measured in arbitrary units.
Table 1: Statistical table

Table 2: Average mTL spike rates were evaluated from foramen ovale electrodes and related to sleep stage as assayed by scalp EEG in two patients with AD.
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A

Spike Count/8s

Time (s)

B

(i)

Mean spike counts

Strong  Weak

(ii)

Sleep  Wake  Sleep  Wake
Table 1: Statistical table

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<tr>
<td>IIS count data – (analysed with log-link function)</td>
<td>MCMC generalized model</td>
<td>Difference between estimates of θ/δ&lt;1 vs. θ/δ&gt;2. Provided for animals JF221, JF220, JF218, J0460, J0456, respectively</td>
<td>(1.619, 2.122)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>(0.261, 0.471)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>(0.254, 0.478)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>(1.166, 1.392)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>(2.128, 2.372)</td>
</tr>
<tr>
<td>Normal (fourth root transformed)</td>
<td>Linear mixed model</td>
<td>β-genotype</td>
<td>(-0.004, 0.008)</td>
</tr>
<tr>
<td>Normal</td>
<td>Linear mixed model</td>
<td>β-Genotype</td>
<td>(-1015.7, 1029.0)</td>
</tr>
<tr>
<td>Non-normal</td>
<td>Wilcoxon signed rank test</td>
<td>Difference of medians</td>
<td>(0.08, 0.65)</td>
</tr>
</tbody>
</table>
### g
**Normal (log transformed)**

<table>
<thead>
<tr>
<th>Tukey contrasts</th>
<th>Difference of mean IIS rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>J20_Ctrl - WT_Ctrl</td>
<td>(-0.24, 0.03)</td>
</tr>
<tr>
<td>WT_DPZ - WT_Ctrl</td>
<td>(-0.15, 0.12)</td>
</tr>
<tr>
<td>J20_DPZ - WT_Ctrl</td>
<td>(-0.08, 0.19)</td>
</tr>
<tr>
<td>WT_NSTG - WT_Ctrl</td>
<td>(-1.50, -1.23)</td>
</tr>
<tr>
<td>WT_DPZ - J20_Ctrl</td>
<td>(-0.04, 0.23)</td>
</tr>
<tr>
<td>J20_DPZ - J20_Ctrl</td>
<td>(0.02, 0.29)</td>
</tr>
<tr>
<td>WT_NSTG - J20_Ctrl</td>
<td>(-1.40, -1.13)</td>
</tr>
<tr>
<td>J20_DPZ - WT_DPZ</td>
<td>(-0.07, 0.21)</td>
</tr>
<tr>
<td>WT_NSTG - WT_DPZ</td>
<td>(-1.49, -1.22)</td>
</tr>
<tr>
<td>WT_NSTG - J20_DPZ</td>
<td>(-1.56, -1.29)</td>
</tr>
</tbody>
</table>

### h
**Normal**  
**Paired t-test**  
**Difference of mean IIS rate**  

<p>|                          | (-0.01, 0.03) |</p>
<table>
<thead>
<tr>
<th>Sleep Stage</th>
<th>Patient #1 (AD dementia)</th>
<th>Patient #2 (aMCI)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total Hours in Record</td>
<td>Total Hours</td>
</tr>
<tr>
<td>Wake</td>
<td>4.7</td>
<td>5.2</td>
</tr>
<tr>
<td>NREM1</td>
<td>0.7</td>
<td>1.5</td>
</tr>
<tr>
<td>NREM2</td>
<td>2.1</td>
<td>3.8</td>
</tr>
<tr>
<td>NREM3</td>
<td>1.4</td>
<td>3.1</td>
</tr>
<tr>
<td>REM</td>
<td>0</td>
<td>0.7</td>
</tr>
</tbody>
</table>