Relevance of sleep and associated structural changes in GBA1 mouse to human rapid eye movement behavior disorder

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Rapid eye movement (REM) sleep behaviour disorder (RBD) is a REM parasomnia that often predicts the later occurrence of alpha-synucleinopathies. Variants in the gene encoding for the lysosomal enzyme glucocerebrosidase, GBA, strongly increase the risk of RBD. In a GBA1-mouse model recently shown to mimic prodromal stages of α-synucleinopathy, we now demonstrate striking REM and NREM electroencephalographic sleep abnormalities accompanied by distinct structural changes in the more widespread sleep neurocircuitry.

Idiopathic (isolated) rapid eye movement (REM) sleep behaviour disorder (iRBD) is a type of REM parasomnia1–3, which represents the prodromal stage of an α-synucleinopathy in more than 80% of patients3–5. Variants in the gene encoding for the lysosomal enzyme glucocerebrosidase, GBA, strongly increase the risk of iRBD6,7. The rate of conversion to neurodegeneration is also increased, and may be faster among severe GBA variant carriers7. This has led some clinicians to argue for more regular screening for iRBD in all healthy carriers of GBA variants in order to identify carriers who will develop α-synucleinopathy in the future6.

Elementary, minor and major body and limb jerks on surface electromyography (REM sleep without atonia) or video polysomnography are a hallmark of iRBD3,8. However, to date, surprisingly little is still known about the (microscopic) sleep architecture in iRBD9–12. Even less is known about its relationship with the GBA-genotype and any associated neuroanatomical or circuitry changes13. In the past, it has been argued that a breakdown in the specific “REM-on” brainstem sleep circuitry residing in the region of the sublaterodorsal tegmental nucleus and the coeruleus/subcoeruleus locus (SLD)14, i.e. the structure analogous to the peri-locus coeruleus-alpha in cat and sublaterodorsalis nucleus (e.g. A6sc) in rodents, might lead to iRBD-associated sleep architectural changes9,15–17. However, the specific changes have by and large remained elusive17. In order to put this concept to the test, whilst accounting for the effect of GBA1-genotype, we used a GBA1-mouse model (heterozygous D409V/WT), recently shown to mimic early prodromal stage of α-synucleinopathy18.

Methods and materials

Three mouse lines were investigated, heterozygous (D409V/WT), homozygous (D409V/D409V) GBA1 mutant mice (D409V/WT) and wild type (C57Bl/6), (further in the text D409V/WT, D409V/D409V and WT respectively), all previously described by us and others18,19 (Supplement Methods; Fig. S1 depicts the research protocol). D409V/WT (heterozygous GBA1 mice) carry one copy of the human D427V point mutation (human equivalent of the murine D409V point mutation) in the murine glucocerebrosidase (Gba) gene. Video-electroencephalogram

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(EEG)-recordings and polysomnography investigations were done according to a strict twelve-hour light–dark cycle protocol, as previously described20. All investigations were performed in accordance with the United Kingdom Home Office Animal Procedures Act (1986) and approved by the King’s College London Animal Welfare Ethical Review Body (AWERB).

Sleep recording and EEG analysis. Ten heterozygous D409V/WT and eight WT mice were implanted with screw electrodes (10–50 kΩ) (Supplement. Fig. S2). The monopolar screw electrodes record the frontal and parietal EEG with ground at occipital bone. In addition, a pair of stainless-steel EMG electrodes was implanted in the dorsal neck muscle. The electrodes were secured with dental cement and standard EEG was recorded for twenty-four hours. EEG/EMG signals were sampled at 200 Hz, amplified 100× and low-pass filtered at 100 Hz using a pre-amplifier (please refer to Supplement Methods for further in depth description). Sleep scoring was performed manually on ten-second epochs using Sirenia Sleep software (Pinnacle Technology Inc.), and signal processing analysis was conducted as previously described21. Further analysis of EEG data in the frequency domain was performed using Fourier transforms. Spectrograms showing the amplitude of EEG signals in the time and frequency domain were generated using either Morlet wavelet or using short-time Fourier transform with 1024 samples window length, as previously described22. EEG signal analyses were conducted using custom codes written in MATLAB, as previously described23–25.

All statistical tests were performed in “GraphPad Prism” or IBM SPSS Statistics version 25.0. Kolmogorov–Smirnov test was used for normality. Data are represented as the mean ± SEM, unless otherwise stated. Two-way ANOVA (time and treatment factors) and two-tailed unpaired t-test were used for the analysis of the sleep and EEG spectrum power data. P values are shown when they are less than 0.05 (please refer to Supplement for further in-depth description and the full list of methodological references).

Neuroimaging methodology. MRI and Statistical Analyses: The MR images were acquired on 9.4 T scanner from ex vivo brains and processed using a combination of FSL, ANTs and the QUIT toolbox, as previously described by our group26. Tensor based morphometry (TBM) was used to assess the effect of genotype on regional brain volumes. A group comparison was carried out on the Jacobian determinant images with permutation tests and Threshold-Free Cluster Enhancement (TFCE) using FSL randomise. Voxelwise differences data (Supplement, Figs. S4–S6) are displayed on the mouse template image. We also performed a region of interest (ROI) analysis using 71 anatomical ROIs derived from a modified Allen brain atlas, that we co-registered to the study template used for TBM. Volumes of each ROI were calculated by summing the Jacobian determinants within the ROI, and univariate pairwise group comparisons performed using Mann–Whitney U-tests. (please refer to Supplement for the full list of methodological references).

Further detailed methodological description of experimental and study procedures, including the undertaken statistical and MR analyses and all the pertinent methodological references, are available in the Supplement. The study is reported in accordance with ARRIVE guidelines27.

Results
Firstly, in order to explore putative similarities of macroscopic and microscopic sleep structure in D409V/WT mice18 to those previously reported in iRBD19–21, 24-h video-EEG recordings of aged-matched D409V/WT and wild type (WT) mice were undertaken. Increased (sleep) latency to NREM, longer and deeper bouts of NREM, overall reduced amount of REM sleep, accompanied by change in REM’s theta power and range, were all found in D409V/WT by comparison to WT mice (Fig. 1a–e).

Specifically, the time spent in NREM throughout the 24-h recording was similar between the two groups (Supplement Fig. S3). D409V/WT mice displayed a longer latency to NREM during the twelve hours lights-on (the “sleep” period) (361.6 ± 24.76 vs 505. ± 52.19 s; P = 0.035, t = 2.29, DF = 16; t-test) (Fig. S3b). However, once in NREM, D409V/WT mice remained in this state longer, as shown by longer duration of the average NREM episodes during the “sleep” period (296.5 ± 20.77 vs 357.2 ± 16.35 s; P = 0.033, t = 2.33, DF = 16; t-test) (Fig. S3c).

Accordingly, sleep fragmentation and the overall number of arousals were similarly reduced (172.8 ± 10.67 vs 137.4 ± 3.53; P = 0.0024, t = 3.56, DF = 17; t-test) (Fig. S3d,e).

Strikingly, D409V/WT mice also spent significantly less time in REM (P(genotype)WT-D409V/WT = 0.035; F(genotype)WT-D409V/WT = 6.098; Fig. 1c), predominantly due to a reduction in the number of REM episodes (lights-on: 47.38 ± 3.32 vs 33.10 ± 2.19; t = 3.71, P = 0.001; lights-off: 25.13 ± 2.34 vs 17.40 ± 1.61, P = 0.0012; t = 2.79; DF = 16; t-test), rather than their duration (Fig. 1c).

Further spectral EEG analyses indicated a significant reduction in the theta frequency (6–9 Hz) both at the NREM-REM transitions (105.1 ± 3.76 vs 92.61 ± 2.45, P = 0.011, t = 2.87, DF = 16; t-test) and during the subsequent REM (211.6 ± 3.52 vs 190.1 ± 5.91, P = 0.009, t = 2.92; DF = 16; t-test) (Fig. 1b,d,e). Increased tonic and phasic EMG activity during REM was also observed in D409V/WT mice by comparison to WT (Supplement Fig. S4).

Secondly, we set to investigate how GBA genotype might affect the (sleep) neurocircuitry by conducting a high resolution ex-vivo magnetic resonance imaging (MRI) of D409V/WT, homozygous (D409V/D409V) and WT mice (Figs. 2, S5–S8).

Structural comparisons have highlighted GBA-genotype-driven hypotrophic changes in widespread cortical, subcortical and white matter regions (Fig. 2). Comparative enlargements were, however, also noted. For example, hypertrophic plastic changes were conspicuous in the infralimbic cortices, the pons, pedunculopontine (PPT) and the dorsal raphe nuclei (DRN; Figs. 2, S5–S8). Of note, preliminary immunohistochemistry findings suggest that hypertrophic changes in the PPT, albeit not in DRN, could be, at least in part, driven by the glial proliferation (Supplement Fig. S9a; Iba1 (mean ± S.D.): WT: 138.6 ± 8.29; D409V/WT: 168.3 ± 17.78; D409V/D409V: 181 ± 36.050; F_D409V/WT-D409V = 0.014 and F_D409V/D409V = 0.021, Mann–Whitney U). Conversely, some
of the recorded hypotrophic changes were associated with the plastic changes in the brainstem’s catecholaminergic neurocircuitry (Supplement, Fig. S9b–d). More specifically, in the dopaminergic region of the ventral tegmental area (VTA) and substantia nigra pars compacta (SNpc), a reduction in tyrosine hydroxylase-positive (TH +) area was recorded (D409V/WT: 13.332 ± 2.708%; D409V/D409V: 12.538 ± 1.171%; WT: 14.666 ± 1.017%; \( P_{D409V/D409VvsWT} = 0.027 \), Mann–Whitney U).

Overall, the observed changes in D409V/WT mice followed the same trend towards the volume loss or gain in all regions of interest (ROIs) as recorded for D409V/D409V (Fig. 2b), suggesting a ‘dose–effect’ of the GBA-mutation on putative neurocircuitry phenotypes. However, D409V/WT-specific structural changes were the most prominent in the regions of medial geniculate nuclei, thalamus and RN, with further distinct cellular changes localised to the noradrenergic region of subcoeruleus/sublaterodorsalis nuclei A6sc (\( P_{D409V/WTvsWT} = 0.046 \), Mann–Whitney U; Supplement Fig. S9c).

Discussion
In summary, we demonstrate here for the first time the EEG phenotype\(^9,10\) of GBA-mutation (D409V/WT) mouse carriers, accompanied by specific structural changes in the more widespread sleep neurocircuitry (Fig. 2). Notably, our findings are in keeping with human data for patients with GBA-mutation, where increased grey matter changes in the brain regions of DRN, midbrain, midbrain reticular formation and PPT have been recently reported\(^13\).

Of particular note to our study are also findings of two very recent studies\(^12,28\). The preliminary results of the first study, conducted in RBD patients, successfully argue for microstates of REM (e.g. phasic versus tonic REM) as a so far ignored, and yet likely highly informative and relatively widely accessible, EEG biomarker of RBD\(^12\). In a second study, Ramirez-Villegas and colleagues\(^28\) elegantly argue for brainstem nuclei (e.g. parabrachial and SLD; Supplement Fig. S9) as pivotal generators of the two types of ponto-geniculate-occipital (PGO) waves that subsequently direct widespread cortical NREM-REM sleep state changes\(^28\). It is here also suggested that putative mechanisms for such selective widespread neuronal modulations might be twofold, via cholinergic...
neuromodulation associated with the ascending brainstem–hippocampal synchronizing pathway that terminates in the medial septum and diagonal band of Broca, both shown as affected in our mice (Fig. 1), and secondly via direct pontine input to the hippocampus²⁸. In light of these two studies, our observation of a reduction in REM-derived theta, with similar qualitative change in NREM-REM band transients in D409V/WT mice (Fig. 1), could be taken to reflect underlying changes in two types of PGO waves²⁸. Moreover, this possibly also suggests an already defective pontine coordination mechanism in D409/WT mice, with likely aberrant systems and synaptic memory consolidation, as well as synaptic homeostasis. This notion is arguably also supported by our distinct neuroanatomical and histological findings that are further suggestive of an already altered function of cholinergic and noradrenergic brain networks (Fig. 1; Supplement).

In past studies of iRBD, lower number of awakenings, lower percentage of wakefulness after sleep onset, higher sleep efficiency and a higher number of sleep stage shifts have all been demonstrated in iRBD patients, which is in broad keeping with our EEG findings for the D409/WT model (Figs. 1, S3)⁹,¹¹. However, in contrast to our findings are results from human studies that overwhelmingly report a relatively preserved sleep architecture in iRBD patients⁹,²⁸, albeit with varied degrees of age-dependant EEG slowing during wake and REM sleep¹⁰,³⁰,³¹. Notwithstanding, contradictory and varied human sleep EEG findings have, however, been reported over the years⁹,¹⁰. For example, in opposition, other studies have reported an altered density of NREM sleep spindles³² and an increased percentage of slow-wave sleep in patients with iRBD³³,³⁴. Similarly, others have demonstrated

Figure 2. GBA1 Genotype-Driven Neuroanatomical Phenotypes. ROIs (a,b) with significant differences (P < .05, FDR corrected) in volume, expressed as % of the averaged wildtype group. Note relative volumes decrease with genotype (WT > D409/WT > D409/D409), except in the dorsal raphe nucleus, pedunculopontine nuclei and infralimbic cortices where there is an opposite trend (WT < D409/WT < D409/D409).
an increased all-night δ power in iRBD\textsuperscript{35}. Correspondingly, a reduced decline in SWA from early to late NREM sleep, along with reduced change in the distribution of the amplitude of slow waves have recently been reported\textsuperscript{32,33}. Nonetheless, to date, the majority of studies appear to report only minimal or absent abnormality of EEG during NREM\textsuperscript{36}.

In regards to REM sleep, several earlier studies have demonstrated substantial EEG abnormality during REM in iRBD\textsuperscript{30}. For instance, impaired cortical activation during both wakefulness and REM in iRBD has been shown, including progressive NREM, REM and transitions instability; although changes in sleep macroarchitecture have been far less consistently demonstrated\textsuperscript{9,10,30,31}. Thus, despite some intriguing similarities, the EEG findings of D409V/WT mice also differ to those previously reported in iRBD. Arguably, this may, at least in part, reflect the distinct sleep pathology of D409V/WT model.

Therefore our findings may, conceivably, represent some of the earliest pathologic EEG changes that occur in distinct relation to GBA1 mutation. As such, they may essentially diverge from those that occur in α-synucleinopathies of other aetiologies. Thus far, very little is understood about the relationship between α-synuclein and GBA, although a strong link has been established between its deficiency and the deposition of aggregated α-synuclein\textsuperscript{32}. However, it is far from clear whether this relationship is consequence of a loss, or gain of, function of GBA due to heterogeneous GBA1 mutation\textsuperscript{25,26}, and even less is known of its impact on the sleep physiology.

Finally, the loss of REM sleep atonia control is well accepted in iRBD patients\textsuperscript{10}, although the underlying mechanism remains ambiguous\textsuperscript{1}. Previous studies have linked a dysfunction of pontine sublateral dorsal/subcoeruleus nucleus, also shown as affected in our study (Supplement, Fig. S9), with loss of REM atonia\textsuperscript{3,39}. According to the current human sleep scoring regulations, in order to distinguish RBD patients reliably, (tonic/phasic) chin EMG activity should be scored in 30-s epochs combined with bilateral phasic activity of the flexor digitorum superficialis muscles in more than 27% of REM sleep\textsuperscript{13}. As yet, no such scoring guidance exists for animal (rodent) RBD model. In our study, though, we observed an intermittent excessive fragmentary myoclonus in nuchal muscles of D409V/WT mice (Fig. S4), presumably in keeping to that reported to occur in prodromal iRBD\textsuperscript{1}. Nonetheless, we find that due to the technical limitations of our study any causal extrapolation is herein limited. For instance, we did not map or analyse the (age-related) phasic and tonic involvement of various rodent muscle groups. This would be important to do in future studies in order to obtain the most optimal EMG biomarker that meaningfully reflects presumed neurodegenerative changes.

Despite obvious limitations of our preliminary and cross-sectional study, we believe that the findings posit important translational questions in regards to whether the reported changes in REM-sleep and its theta-frequency-range power\textsuperscript{20}, along with involvement of core circuitry structures such are PPT, RN and SLD suggest a differential phenotype of any early non-motor (e.g, cognitive, memory and mood) symptoms in non-manifesting human GBA carriers. Similarly, we believe that these findings also advance the relevance of the GBA1 mice as a valuable model for future exploration of neurostructural and physiological targets in iRBD, and its links with other α-synucleinopathies. Future studies should aim to delineate authoritative age-related EEG changes in this animal model, ideally guided and informed by the concomitant multidisciplinary and multi-centre translational trials of RBD patients with different GBA variants.

**Data availability**

All data that support the findings of this study are available upon reasonable request from the corresponding author.

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Author contributions
C.G., D.C., P.F. and I.R. designed the study. C.G., D.C., C.S., E.K., K.I. conducted and analysed data, all authors were involved in preparation and review of the manuscript.

Competing interests
The authors declare no competing interests.

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