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Consistent abnormalities in metabolic network activity in idiopathic rapid eye movement sleep behaviour disorder

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Rapid eye movement sleep behaviour disorder has been evaluated using Parkinson’s disease-related metabolic network. It is unknown whether this disorder is itself associated with a unique metabolic network. [18F]-fluorodeoxyglucose positron emission tomography was performed in 21 patients (age 65.0 ± 5.6 years) with idiopathic rapid eye movement sleep behaviour disorder and 21 age/gender-matched healthy control subjects (age 62.5 ± 7.5 years) to identify a disease-related pattern and examine its evolution in 21 hemi-parkinsonian patients (age 62.6 ± 5.0 years) and 16 moderate parkinsonian patients (age 56.9 ± 12.2 years). We identified a rapid eye movement sleep behaviour disorder-related metabolic network characterized by increased activity in pons, thalamus, medial frontal and sensorimotor areas, hippocampus, supramarginal and inferior temporal gyri, and posterior cerebellum, with decreased activity in occipital and superior temporal regions. Compared to the healthy control subjects, network expressions were elevated (P < 0.0001) in the patients with this disorder and in the parkinsonian cohorts but decreased with disease progression. Parkinson’s disease-related network activity was also elevated (P < 0.0001) in the patients with rapid eye movement sleep behaviour disorder but lower than in the hemi-parkinsonian cohort. Abnormal metabolic networks may provide markers of idiopathic rapid eye movement sleep behaviour disorder to identify those at higher risk to develop neurodegenerative parkinsonism.

Keywords: rapid eye movement sleep behaviour disorder; Parkinson’s disease; [18F]-fluorodeoxyglucose; positron emission tomography; network biomarkers

Abbreviations: FDG = [18F]-fluorodeoxyglucose; PDRP = Parkinson’s disease-related pattern; RBD = rapid eye movement sleep behaviour disorder; RBDRP = rapid eye movement sleep behaviour disorder-related pattern; REM = rapid eye movement

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Introduction

Rapid eye movement sleep behaviour disorder (RBD) is a para-somnia characterized by dream enactment behaviours emerging during REM sleep and associated with a loss of normal skeletal muscle atonia (Boeve, 2010). Abnormal neural circuits resulting from damage to brainstem areas may have precipitated idiopathic RBD leading to the loss of motor inhibition during rapid eye movement (REM) sleep (Boeve et al., 2007; Lu and Saper, 2011). Idiopathic RBD is regarded as a prodromal stage of synucleinopathies such as Parkinson’s disease, dementia with Lewy bodies and multiple system atrophy because it often precedes the development of motor, cognitive, neuropsychiatric and autonomic features of these syndromes by years or decades (Iranzo et al., 2006; Postuma et al., 2009; Claassen et al., 2010; Schenck et al., 2013). Functional neuroimaging studies with Single Photon Emission Computed Tomography (SPECT) or PET provided further support with reduced nigrostrial dopaminergic activity (Eisensehr et al., 2000; Iranzo et al., 2011), decreased cardiac adrenergic activity (Miyamoto et al., 2006; Fujishiro et al., 2010) and impaired brain perfusion (Hanyu et al., 2011; Vendette et al., 2011) found in patients with idiopathic RBD.

Metabolic imaging with FDG PET and spatial covariance analysis has provided valuable information concerning the abnormal regional brain function underlyng neurodegenerative disorders (Eidelberg, 2009). Abnormal metabolic brain networks have been identified for Parkinson’s disease, multiple system atrophy and progressive supranuclear palsy for accurate classification of these parkinsonian disorders at the single-patient level, even at early disease stages (Ma et al., 2007; Eckert et al., 2008; Tang et al., 2010). We have also identified a Parkinson’s disease-related covariance pattern (PDRP) in a Chinese cohort (Wu et al., 2013) that closely resembled the original topography identified in an analogous North American cohort (Niethammer and Eidelberg, 2012).

Abnormal changes in PDRP expression have recently been reported in two idiopathic RBD cohorts and, on an individual basis, higher baseline expression was associated with a greater likelihood of developing Parkinson’s disease or dementia with Lewy bodies (Holtbernd et al., 2014). It is not known, however, whether idiopathic RBD is itself associated with a unique metabolic topography that would evolve before symptom onset. We sought to identify a RBD-related covariance pattern (RBDRP) and determined its relationships with concurrent PDRP measurements using our network-based algorithm (Ma et al., 2007; Spetsieris et al., 2013). We also assessed the evolution of these network expressions in patients with idiopathic RBD and in patients with very early and moderate Parkinson’s disease.

Materials and methods

Subjects

For identification of RBDRP, 21 patients with idiopathic RBD [male/female: 17/4, age: 65.0 ± 5.6 (mean ± standard deviation, SD) years] and 21 gender/age-matched healthy control subjects (male/female: 17/4, age: 62.5 ± 7.5 years) were recruited from Huashan Hospital for 18F-Fluorodeoxyglucose (FDG) PET imaging (Cohort A). Diagnosis of idiopathic RBD was made by sleep medicine experts (H.Y. and Y.D.) requiring a history of dream-enacting behaviours and video-polysomnographic demonstration of either increased EMG activity or abnormal behaviours during REM sleep (Boeve, 2010). Detailed polysomnographic procedures are included in the online Supplementary material.

For prospective validation of RBDRP network expression we included Cohort B with 15 patients with idiopathic RBD (male/female: 12/3, age: 65.5 ± 6.0 years) and 15 gender/age-matched healthy control subjects (male/female: 10/5, age: 60.5 ± 7.6 years). This idiopathic RBD group comprised nine patients from Cohort A who were scanned again at 1 year (male/female: 7/2, age: 65.2 ± 6.9 years) and six newly recruited subjects with idiopathic RBD (male/female: 5/1, age: 66.0 ± 5.0 years). Clinical and polysomnographic parameters did not differ significantly between the patients with idiopathic RBD in Cohorts A and B (Table 1).

We also included Cohort C with 21 age-matched patients with hemi-Parkinson’s disease (male/female: 12/9, age: 62.6 ± 5.0 years, Hoehn and Yahr 1) and 16 patients with moderate Parkinson’s disease (male/female: 5/11, age: 56.9 ± 12.2 years, Hoehn and Yahr: 2.5 ± 0.5, range 2–3) to assess the correlations between the RBDRP and PDRP networks. All normal and parkinsonian subjects were recruited as described previously (Wu et al., 2013).

Ethical permission for the study was obtained from the Institutional Review Board of Huashan Hospital. Written consent was obtained from each subject after detailed explanation of the procedures.

PET imaging and processing

The imaging procedure and the data preprocessing were performed as described in detail previously (Wu et al., 2013). Briefly, FDG PET images of relative glucose metabolism were acquired with a Siemens Biograph 64 PET/CT (Siemens) and spatially normalized into a standard brain space by statistical parametric mapping software (SPM5; Institute of Neurology, London, UK).

Network analysis and prospective comparisons

RBDRP was identified from the combination of idiopathic RBD and control subjects in Cohort A by an automated voxel-based network analysis over the whole brain (Ma et al., 2007; Spetsieris et al., 2013). This was done within a grey matter mask defined with all FDG PET images by applying scaled subprofile model/principal component analysis (SSM/PCA; available on http://www.feinsteinneuroscience.org at the Centre for Neuroscience, the Feinstein Institute for Medical Research, Manhasset, NY, USA). The reliability of the resulting topography was evaluated by using a bootstrapping resample scheme described previously (Habecock et al., 2008; Peng et al., 2014).

The RBDRP was compared topographically to the PDRP (Wu et al., 2013) using volume- or voxel-wise correlations over the whole brain. The expressions of RBDRP or PDRP were quantified in all subjects with a voxel-based algorithm (Ma et al., 2007; Spetsieris et al., 2013). The network expression was Z-transformed using subject scores of the controls in the corresponding derivation cohort for each network.
**Table 1 Clinical characteristics and polysomnographic measures in patients with idiopathic RBD**

<table>
<thead>
<tr>
<th></th>
<th>Cohort A</th>
<th>Cohort B</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex (male/female)</td>
<td>17/4</td>
<td>12/3</td>
</tr>
<tr>
<td>Age (years)</td>
<td>65.0 ± 5.6</td>
<td>65.5 ± 6.0</td>
</tr>
<tr>
<td>Age at onset of RBD (years)</td>
<td>59.3 ± 6.9</td>
<td>60.1 ± 7.6</td>
</tr>
<tr>
<td>RBD duration (years)</td>
<td>5.7 ± 3.5</td>
<td>5.5 ± 4.0</td>
</tr>
<tr>
<td>UPDRS III</td>
<td>2.4 ± 2.1</td>
<td>2.8 ± 1.9</td>
</tr>
<tr>
<td>Body mass index (kg/m²)</td>
<td>23.8 ± 2.7</td>
<td>23.8 ± 2.5</td>
</tr>
<tr>
<td>Polysomnographic data</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total sleep time (min)</td>
<td>329.2 ± 47.0</td>
<td>325.3 ± 39.5</td>
</tr>
<tr>
<td>Sleep efficiency (%)</td>
<td>74.9 ± 9.4</td>
<td>73.5 ± 10.7</td>
</tr>
<tr>
<td>Sleep onset latency(min)</td>
<td>26.7 ± 32.2</td>
<td>21.1 ± 24.9</td>
</tr>
<tr>
<td>REM sleep latency(min)</td>
<td>81.4 ± 38.0</td>
<td>96.1 ± 37.2</td>
</tr>
<tr>
<td>Non-REM sleep 1 (%)</td>
<td>17.2 ± 8.7</td>
<td>17.1 ± 9.4</td>
</tr>
<tr>
<td>Non-REM sleep 2 (%)</td>
<td>46.2 ± 9.2</td>
<td>47.7 ± 7.5</td>
</tr>
<tr>
<td>Non-REM sleep 3 (%)</td>
<td>19.5 ± 8.2</td>
<td>18.0 ± 8.5</td>
</tr>
<tr>
<td>REM sleep (%)</td>
<td>17.2 ± 5.1</td>
<td>17.1 ± 5.8</td>
</tr>
<tr>
<td>Microarousals index</td>
<td>15.2 ± 7.4</td>
<td>16.1 ± 4.9</td>
</tr>
<tr>
<td>Phasic EMG activity index/chin</td>
<td>17.9 ± 9.8</td>
<td>17.5 ± 8.5</td>
</tr>
<tr>
<td>Tonic EMG activity index /chin</td>
<td>16.0 ± 11.9</td>
<td>17.8 ± 16.4</td>
</tr>
<tr>
<td>Phasic EMG activity index /legs</td>
<td>12.7 ± 7.1</td>
<td>13.5 ± 8.2</td>
</tr>
<tr>
<td>Tonic EMG activity index /legs</td>
<td>8.1 ± 6.3</td>
<td>8.0 ± 7.3</td>
</tr>
<tr>
<td>Patients with either abnormal phasic or tonic chin EMG activity in REM sleep (%)</td>
<td>10 (47.6%)</td>
<td>7 (46.7%)</td>
</tr>
<tr>
<td>Apnoea-hypopnoea index</td>
<td>3.0 ± 4.0</td>
<td>3.8 ± 2.7</td>
</tr>
<tr>
<td>Mean SaO²</td>
<td>94.5 ± 0.9</td>
<td>94.5 ± 1.2</td>
</tr>
</tbody>
</table>

Values are given as means ± SD.

**Statistical analysis**

Differences in network expression between patients and normal control subjects were assessed by Student’s t-tests with their diagnostic power for group discrimination evaluated by receiver operating characteristic analysis. One-way ANOVA was performed for each network measure across the subject groups with between-group differences assessed using post hoc Tukey’s HSD tests. Group × network activity interaction was evaluated by two-way ANOVA to determine whether any differences exist between changes for each of these network measures over the subject groups. Correlations between RBDRP and PDRP expressions as well as those with clinical measures in different subject groups were separately assessed by computing Pearson correlation coefficients. All analyses were performed using SPSS software (SPSS Inc.) and considered significant for \( P < 0.05 \).

**Results**

**RBD-related pattern identification and disease discrimination**

In Cohort A, the RBDRP was identified as the first principal component accounting for 14.0% of subject × voxel variance. The pattern was characterized by increased metabolic activity in pons, thalamus, precentral gyrus, supplementary motor area, medial frontal gyrus, hippocampus/parahippocampal gyrus, supramarginal and inferior temporal gyrus, and posterior cerebellar tonsil, associated with decreased metabolic activity in occipital regions, midbrain (red nucleus) and superior/middle temporal gyrus (Fig. 1A; Supplementary Table 1). RBDRP expression was abnormally elevated \( (P < 0.0001) \) in the patients with idiopathic RBD compared with the control subjects (Fig. 1B) and significantly discriminated \( (P < 0.0001) \) the subject groups, with an area under the curve of 0.989 ± 0.013 and a sensitivity of 95.2% and a specificity of 95.2%.

**Comparisons between spatial topographies of RBD- and Parkinson’s disease-related patterns**

Both RBDRP and PDRP showed varying degrees of overlap in several brain areas including the thalamus, sensorimotor cortices and occipital regions (Figs 1A and 2A). Their region-weights correlated significantly with each other using both volumetric \( r = 0.634, P < 0.002 \) and voxelized \( r = 0.680; P < 0.0001 \) analyses.

**RBD-related pattern expressions in patients with Parkinson’s disease**

RBDRP expressions in different patient cohorts were presented in Table 2 and Fig. 1B. RBDRP activity differed significantly among the multiple subject groups \( F(3,75) = 51; F(3,63) = 32, P < 0.0001; \) one-way ANOVA across Cohorts A and C or B and C, respectively). RBDRP expression in the patients with idiopathic RBD was similarly elevated \( (P < 0.0001) \) relative to the controls in Cohort A/B. RBDRP scores were also elevated \( (P < 0.0001) \) in the hemi-parkinsonian patients but slightly lower \( (P > 0.18) \) than these patients with idiopathic RBD. However, these scores decreased significantly in the moderate parkinsonian patients compared to the idiopathic RBD patients in Cohort A/B \( (P = 0.032/0.014) \). RBDRP scores did not differ between the two parkinsonian groups \( (P > 0.22) \).

**Parkinson’s disease-related pattern expressions in patients with idiopathic RBD**

PDRP expressions in different patient cohorts were presented in Table 2 and Fig. 2B. PDRP activity differed significantly among the multiple subject groups \( F(3,75) = 42; F(3,63) = 21, P < 0.0001; \) one-way ANOVA across Cohorts A and C or B and C, respectively). PDRP scores in the idiopathic RBD patients in Cohort A/B were similarly elevated \( (P < 0.0001) \) relative to the controls, decreased not significantly \( (P > 0.74) \) versus the hemi-parkinsonian patients but significantly \( (P = 0.001/0.047) \) versus the moderate parkinsonian patients. PDRP scores increased significantly \( (P = 0.014/0.035) \) comparing the moderate to the hemi-parkinsonian groups.
Figure 1 (A) RBDRP identified by network analysis of FDG PET scans from 21 patients with idiopathic RBD and 21 age-matched normal controls in Cohort A. This spatial covariance pattern was characterized by relative increases in sensorimotor, superior frontal, cingulate, thalamic, pontine, and cerebellar metabolism, associated with decreases in occipital, midbrain and superior temporal metabolism. The pattern was overlaid on a standard MRI brain template to display voxels that were reliable at $P < 0.01$ based on the bootstrapping algorithm. Voxels with positive region weights (metabolic increases) are colour-coded red; those with negative region weights (metabolic decreases) are colour-coded blue. (B) RBDRP expressions (subject scores) were increased in the patients with idiopathic RBD relative to the normal subjects in Cohort A as well as the patients with moderate Parkinson’s disease in Cohort C. These scores were also elevated when compared to those in the hemi-parkinsonian patients in Cohort C but did not reach significance. Subject scores of the parkinsonian patients in Cohort C were both elevated ($P < 0.0001$; post hoc tests) versus the normal controls. The error bars represent standard errors of the mean. NL = normal control; HY = Hoehn and Yahr; SMA = supplementary motor area.

Figure 2 (A) PDRP identified by network analysis of FDG PET scans from 33 patients with Parkinson’s disease and 33 age-matched normal control subjects (Wu et al., 2013). This spatial covariance pattern was characterized by relative increases in sensorimotor, pallidothalamic, pontine, and cerebellar metabolism, associated with decreases in premotor and posterior parietal-occipital metabolism. The pattern was overlaid on a standard MRI brain template to display voxels that were reliable at $P < 0.001$ based on the bootstrapping algorithm. Voxels with positive region weights (metabolic increases) are colour coded red; those with negative region weights (metabolic decreases) are colour coded blue. (B) PDRP expressions (subject scores) were increased in the patients with idiopathic RBD relative to the normal subjects in Cohort A but decreased relative to the moderate parkinsonian patients in Cohort C. These scores were also decreased slightly but not significantly when compared to the hemi-parkinsonian patients in Cohort C. Subject scores of the parkinsonian patients in Cohort C were both elevated ($P < 0.0001$; post hoc tests) versus the normal controls. The error bars represent standard errors of the mean. GP = globus pallidus; PU = putamen; NL = normal control; HY = Hoehn and Yahr; SMA = supplementary motor area; SMC = sensorimotor cortex.
Comparisons of network activities and clinical correlations in patients with idiopathic RBD and Parkinson’s disease

RBDPR and PDRP scores disclosed an effect of interaction across the multiple subject groups ($F = 9.4/9.3$, $P = 0.003$; two-way ANOVA across Cohorts A and C or B and C, respectively). RBDPR and PDRP scores correlated in the idiopathic RBD patients in Cohort A ($r = 0.710$, $P < 0.0001$) and in the parkinsonian cohort ($r = 0.387$, $P = 0.018$). UPDRS motor ratings correlated with PDRP ($r = 0.481$, $P = 0.003$) but not RBDPR ($r = 0.14$, $P = 0.41$) scores in the parkinsonian cohort. Moreover, neither network scores correlated with clinical and polysomnographic parameters in patients with idiopathic RBD.

Discussion

This report describes the first study to identify a characteristic metabolic brain network (i.e., RBDPR) with FDG PET in an idiopathic RBD population. RBDPR featured a highly symmetrical topography suggesting the existence of a widely-distributed brain network of abnormal metabolic activity underlying this disorder. RBDPR expression accurately discriminated patients with RBD from normal control subjects in both derivation and validation samples. Moreover, RBDPR activity decreased but PDRP activity increased in early to moderate parkinsonian patients compared to patients with idiopathic RBD.

Comparisons of brain network characteristics associated with idiopathic RBD and Parkinson’s disease

We have noted that RBDPR and PRDP share some similarities. Firstly, the two topographic patterns showed moderate overlap. Secondly, both RBDPR and PDRP expressions were elevated and correlated with each other in patients with RBD. Thirdly, elevated PDRP expression in the patients with idiopathic RBD was lower or comparable to the hemi-parkinsonian RBD was lower or comparable to the hemi-parkinsonian patients, in agreement with the recent report (Holtbernd et al., 2014).

We have also observed unique differences between RBDPR and PRDP. Firstly, the RBDPR included several specific regions such as hippocampus and midbrain, but excluded the putamen and pallidum (Figs 1A and 2A). Secondly, RBDPR activity declined while PRDP activity increased steadily in patients with idiopathic RBD, hemi- and moderate Parkinson’s disease, given the elevation of both network activities in each of these patient cohorts relative to the controls. Thirdly, not surprisingly, subject scores of PDRP (not RBDPR) correlated moderately with clinical motor ratings in the combined patient cohort of very early to moderate Parkinson’s disease.

Relationships with dementia with Lewy bodies or multiple system atrophy

Occipital and parietotemporal hypometabolism evident in both RBDPR and PDRP is also a finding frequently seen on FDG PET and perfusion SPECT in patients with probable dementia with Lewy bodies (Mosconi et al., 2008; Inui et al., 2014). Thus, a PDRP-like network may underlie dementia with Lewy bodies to a large degree and an elevated PDRP activity in idiopathic RBD may also point to the subsequent development of this disorder with disease progression (Holtbernd et al., 2014). Of the 17 subjects with idiopathic RBD followed clinically in that study, eight (47%) who had similarly elevated PDRP scores at baseline were diagnosed with Parkinson’s disease or dementia with Lewy bodies after 4.6 ± 2.5 years, in line with the estimated risk in the same period (Franzo et al., 2006; Postuma et al., 2009). That said, it may still be necessary to assess other likely clinical outcomes using different spatial covariance patterns associated with Parkinson’s disease-related cognitive impairment (Huang et al., 2007) and multiple system atrophy (Eckert et al., 2008; Teune et al., 2013).

Evolving metabolic brain networks and activities in patients with idiopathic RBD with disease progression

RBDPR depicts an insightful spatial covariance map for describing intrinsic connectivity between regional abnormalities and related pathways. The topography of RBDPR includes increased metabolic activity in the pons, thalamus and hippocampus, consistent with regional abnormalities measured with FDG PET and perfusion SPECT in several idiopathic RBD cohorts (Vendette et al., 2011; Dang-Vu et al., 2012; Holtbernd et al., 2014). Notably, in addition to increases in the putamen, the latter studies also found that increased metabolism and perfusion in the pons and/or hippocampus at baseline were predictive of the eventual onset of Parkinson’s disease or dementia with Lewy bodies in 3–5 years. Overall, the present study has revealed several dysfunctional neural circuits linking brainstem, midbrain, cerebellum, thalamus and fronto-occipital cortices, in line with the pathophysiology of idiopathic RBD and related disorders (Boeve et al., 2007; Lu and Saper, 2011; Del Tredici and Braak, 2013).

Table 2 Network expressions in normal controls, patients with idiopathic RBD and Parkinson’s disease

<table>
<thead>
<tr>
<th>Cohort</th>
<th>n</th>
<th>RBDRP scores</th>
<th>PDRP scores</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cohort A</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Normal control</td>
<td>21</td>
<td>0.00 ± 1.00</td>
<td>0.08 ± 0.72</td>
</tr>
<tr>
<td>Idiopathic RBD</td>
<td>21</td>
<td>6.31 ± 2.23</td>
<td>2.25 ± 0.97</td>
</tr>
<tr>
<td>Cohort B</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Normal control</td>
<td>15</td>
<td>1.03 ± 1.48</td>
<td>0.47 ± 0.95</td>
</tr>
<tr>
<td>Idiopathic RBD</td>
<td>15</td>
<td>6.96 ± 1.54</td>
<td>2.52 ± 1.22</td>
</tr>
<tr>
<td>Cohort C</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Parkinson’s disease (Hoehn and Yahr 1)</td>
<td>21</td>
<td>5.72 ± 1.59</td>
<td>2.57 ± 0.93</td>
</tr>
<tr>
<td>Parkinson’s disease (Hoehn and Yahr 2–3)</td>
<td>16</td>
<td>4.97 ± 2.38</td>
<td>3.60 ± 1.38</td>
</tr>
</tbody>
</table>

Values are given as means ± SD.
The findings that RBDRP activity decreased but PDRP activity increased in patients with advancing Parkinson’s disease compared to patients with idiopathic RBD (Figs 1B and 2B) suggest that RBDRP is perhaps relevant only for prodromal RBD cases and likely to break down with disease progression. This is supported by the lack of any correlations between RBDRP activity and clinical motor ratings in either RBD or parkinsonian cohort. Collectively these results pointed out the hypothesis of an evolving abnormal metabolic network with disease progression: from a RBDRP at very early stages (i.e. idiopathic RBD) to a PDRP at later stages when a neurodegenerative process is really advanced to that of Parkinson’s disease. This process is expected to coincide with the appearance in the resulting network of covarying subcortical regions in basal ganglia (i.e. putamen) with increased metabolic activity. This is a hypothesis to be validated by long-term clinical and imaging studies.

This study has some limitations. First, brain metabolism has been assessed not in REM sleep when the abnormal behaviour occurred. However, as currently reported in wake resting state, an increased perfusion in supplementary motor area during a RBD episode in a severe patient affected with multiple system atrophy was also reported in a SPECT study (Dauvilliers et al., 2011). Second, no absolute brain glucose metabolism was quantified to adequately propose a pathophysiological explanation on the network architecture underlying idiopathic RBD. Nevertheless, this study did disclose an abnormal metabolic network implying a specific brain circuitry. Third, elevated RBDRP/PDRP expressions in the patients with idiopathic RBD relative to the control subjects were replicated but not in a truly independent cohort. Regardless this result at least demonstrates that these metabolic network activities are highly reproducible across different idiopathic RBD subjects or metabolic images. Finally, idiopathic RBD was diagnosed by a visual scoring method with excellent sensitivity and specificity for both phasic and tonic chin EMG measures. However, despite a typical history of dream-enacting behaviours in all patients with idiopathic RBD, only ~48% of them had abnormal REM sleep without atonia with the remainder presenting complex movements during REM sleep.

In conclusion, we have reported the first identification of RBDRP with FDG PET, which may be useful in differentiating patients with idiopathic RBD from normal controls and in revealing similarities and unique differences with PDRP. The close association of neuropathological processing between RBDR and Parkinson’s disease suggests that idiopathic RBD is indeed a prodromal phase of Parkinson’s disease along a wide spectrum of disease progression. Further prospective studies are necessary to determine the relationships between baseline measurements in network expression and eventual clinical outcomes of idiopathic RBD patients.

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Supplementary material

Supplementary material is available at Brain online.

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