



Behavioural Neurology

Mind-wandering in Parkinson's disease hallucinations reflects primary visual and default network coupling



Ishan C. Walpola^a, Alana J. Muller^a, Julie M. Hall^{a,b},
 Jessica R. Andrews-Hanna^c, Muireann Irish^{a,d,e}, Simon J.G. Lewis^a,
 James M. Shine^a and Claire O'Callaghan^{a,f,*}

^a Brain and Mind Centre and Central Clinical School, Faculty of Medicine and Health, University of Sydney, Sydney, Australia

^b School of Social Sciences and Psychology, Western Sydney University, Sydney, Australia

^c Department of Psychology and Cognitive Science Program, University of Arizona, Tucson, AZ, USA

^d Australian Research Council Centre of Excellence in Cognition and its Disorders, Sydney, Australia

^e School of Psychology, University of Sydney, Sydney, Australia

^f Department of Psychiatry and Behavioural and Clinical Neuroscience Institute, University of Cambridge, Cambridge, UK

ARTICLE INFO

Article history:

Received 26 July 2019

Reviewed 23 September 2019

Revised 26 September 2019

Accepted 22 December 2019

Action editor Stefano Cappa

Published online 22 January 2020

Keywords:

Parkinson's disease

Visual hallucinations

Mind-wandering

Default network

Resting state functional magnetic resonance imaging

ABSTRACT

Visual hallucinations are an underappreciated symptom affecting the majority of patients during the natural history of Parkinson's disease. Little is known about other forms of abstract and internally generated cognition – such as mind-wandering – in this population, but emerging evidence suggests that an interplay between the brain's primary visual and default networks might play a crucial role in both internally generated imagery and hallucinations. Here, we explored the association between mind-wandering and visual hallucinations in Parkinson's disease, and their relationship with brain network coupling. We administered a validated thought-sampling task to 38 Parkinson's disease patients (18 with hallucinations; 20 without) and 40 controls, to test the hypothesis that individuals with hallucinations experience an increased frequency of mind-wandering. Group differences in the association between mind-wandering frequency and brain network coupling were also examined using resting state functional magnetic resonance imaging. Our results showed that patients with hallucinations exhibited significantly higher mind-wandering frequencies compared to non-hallucinators, who in turn had reduced levels of mind-wandering relative to controls. At the level of brain networks, inter-network connectivity and seed-to-voxel analyses identified that increased mind-wandering in the hallucinating versus non-hallucinating group was associated with greater coupling between the primary visual cortex and dorsal default network. Taken together, our results suggest a relative preservation of mind-wandering in Parkinson's disease patients who

* Corresponding author. Herchel Smith Building for Brain & Mind Sciences, Cambridge Biomedical Campus, Cambridge, CB2 0SZ, UK.

E-mail address: claire.ocallaghan@sydney.edu.au (C. O'Callaghan).

<https://doi.org/10.1016/j.cortex.2019.12.023>

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experience visual hallucinations, which is associated with increased visual cortex-default network coupling. We propose that the preservation of florid abstract and internally generated cognition in the context of the Parkinson's disease can contribute to visual hallucinations, whereas healthy individuals experience only the vivid images of the mind's eye. These findings refine current models of visual hallucinations by identifying a specific cognitive phenomenon and neural substrate consistent with the top-down influences over perception that have been implicated in hallucinations across neuropsychiatric disorders.

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1. Introduction

Hallucinations that are predominantly visual in nature affect over 40% of patients in the early stages of Parkinson's disease, and upwards of 80% as the disease progresses (Ffytche et al., 2017). Parkinson's disease visual hallucinations have been associated with increased activity and connectivity in the default network, both in the resting state (Franciotti et al., 2015; Yao et al., 2014) and during recorded visual misperceptions (Shine, Muller et al., 2015). The default network is situated most distantly from primary sensory networks along a hierarchical gradient of cortical connectivity (Huntenburg, Bazin, & Margulies, 2018; Margulies et al., 2016). As such, it is a key candidate for generating certain high-level predictions to prepare earlier visual regions for incoming sensory information, thereby increasing perceptual sensitivity and efficiency (de Lange, Heilbron, & Kok, 2018).

Several models of Parkinson's disease visual hallucinations have posited that in the context of poor quality visual input – due to attentional impairments, visual deficits and retinal pathology – higher-order regions involved in the internal generation of mental imagery and perceptual expectancies exert excessive influence upon perception (Collerton, Perry, & McKeith, 2005; Diederich, Goetz, & Stebbins, 2005; Shine, O'Callaghan, Halliday, & Lewis, 2014). An instantiation of such models is that an over-active or unconstrained default network dominates the perceptual processes in a top-down manner, supplying perceptual predictions in the form of internally generated imagery that overrides incoming sensory information (Powers, Kelley, & Corlett, 2016; Shine, O'Callaghan, Halliday, & Lewis, 2014; O'Callaghan, Kveraga, Shine, Adams, & Bar, 2017; Onofri, Espay, Bonanni, Delli Pizzi, & Sensi, 2019). This may be further exacerbated in Parkinson's disease due to the numerous sources of visual and attentional dysfunction that render incoming sensory information less reliable (Weil et al., 2016). Such an imbalance between top-down influences and incoming sensory input can be interpreted in a Bayesian predictive coding framework, in which increased precision (i.e., relative weighting) is consistently afforded to prior beliefs (i.e., top-down expectations or predictions), such that they override incoming sensory evidence and dominate the ultimate percept (Adams, Stephan, Brown, Frith, & Friston, 2013; Fletcher & Frith, 2009; Friston, 2005). Patients with Lewy body disease and

hallucinations have been shown to rely more heavily on prior knowledge during perceptual inference, relative to non-hallucinators (Zarkali et al., 2019). Attentional network dysfunction – specifically involving the dorsal attention network – has been established in Parkinson's disease with visual hallucinations, and may reflect an inability to dynamically modulate precision at the mesoscale of brain function (Hall et al., 2019; Shine, Muller et al., 2015; Shine, Keogh et al., 2015). This is in keeping with a recent finding that in Parkinson's disease with visual hallucinations, accumulation of sensory evidence is slow and inefficient – and is therefore less informative – which may cause it to be down-weighted in favour of relatively preserved perceptual priors (O'Callaghan et al., 2017).

The hierarchical nature of the brain's visual processing system supports the reciprocal feed-forward/feed-back flow of information between early visual and higher-order transmodal regions across the cerebral cortex (Gilbert & Li, 2013). More specifically, regions within the default network have been identified as sources of top-down influence over visual perception. These regions include the orbitofrontal and medial prefrontal cortices, which use early low spatial frequency information to generate expectations that constrain ongoing visual processing (Bar, 2003; Bar et al., 2006; Chaumon, Kveraga, Barrett, & Bar, 2014; Kveraga, Boshyan, & Bar, 2007; Summerfield et al., 2006); hippocampal pattern completion mechanisms that supply memory-based expectations to the visual cortex (Hindy, Ng, & Turk-Browne, 2016); parahippocampal and retrosplenial cortices supporting the rapid activation of contextual associations during visual processing (Aminoff, Kveraga, & Bar, 2013; Kveraga et al., 2011); and, distinct populations of neurons in the inferior temporal cortex that encode predictions and prediction errors in relation to incoming visual input (Bell, Summerfield, Morin, Malecek, & Ungerleider, 2016; Kok, 2016). The temporal properties of supramodal regions such as the default network are also consistent with the unfolding of high-level predictions over longer timescales (Baldassano et al., 2017; Margulies et al., 2016; Weillnhammer, Stuke, Sterzer, & Schmack, 2018), which fits well with the complex and temporally extended hallucinations that occur in Parkinson's disease (Ffytche et al., 2017). Yet, despite a number of established routes by which the default network may influence visual perception, and evidence of unconstrained default network activity in Parkinson's disease visual hallucinations, we know very little about the

behavioural consequences of an over-engaged default network and how this might contribute to hallucinations.

In keeping with its role as a source of top-down influence over visual perception, the default network is implicated in many cognitive processes relevant for internally generating expectations about the sensory environment, including mental imagery and scene construction, autobiographical memory, prospection, and retrieval of contextual associations (Andrews-Hanna, Reidler, Sepulcre, Poulin, & Buckner, 2010; Buckner, Andrews-Hanna, & Schacter, 2008; Kveraga et al., 2011; Schacter et al., 2012; Spreng, Mar, & Kim, 2009). One common experience that draws upon the aforementioned cognitive processes, and is strongly linked to activity in the default network, is mind-wandering. Mind-wandering is often characterised as thoughts that are decoupled from the immediate perceptual environment and unrelated to ongoing task demands (Seli et al., 2018; Smallwood & Schooler, 2015). Recent frameworks further emphasise that mind-wandering is a mental state that arises spontaneously, in which thoughts are unguided and unconstrained (Christoff, Irving, Fox, Spreng, & Andrews-Hanna, 2016; Irving, 2016).

There is a striking similarity between certain core characteristics of both mind-wandering and visual hallucinations: both are transient forms of spontaneous cognition, relatively unconstrained by sensory input, and are underpinned by dynamic shifts in the interactions both within and between similar large-scale brain networks (Christoff et al., 2016; Collerton et al., 2016; Kucyi, 2017; Zabelina & Andrews-Hanna, 2016). Given the shared phenomenology and evidence for an overlapping neural basis, we predicted that clinical subgroups with visual hallucinations would also show changes in their propensity for mind-wandering. While such definitive studies have not been conducted to date, in patients with schizophrenia higher frequencies of task-unrelated thought have been observed, correlating with the severity of their positive symptoms, including hallucinations (Shin et al., 2015). Additionally, previous work in Parkinson's disease has demonstrated that individuals with visual hallucinations exhibit stronger mental imagery – a prominent feature of mind-wandering (Shine, Keogh et al., 2015). Taken together, these findings suggest that increased mind-wandering may provide a cognitive correlate for the predisposition to experience hallucinatory phenomena across disease states.

To address the question of whether mind-wandering is related to hallucinations in Parkinson's disease, the present study utilised a validated thought sampling task designed for clinical populations with cognitive impairment (O'Callaghan, Shine, Lewis, Andrews-Hanna, & Irish, 2015). Using this task, we measured mind-wandering frequencies in Parkinson's disease patients with and without visual hallucinations, and healthy controls. To explore the neural correlates of mind-wandering frequency, we used network-level and seed-to-voxel analysis of resting-state functional magnetic resonance imaging. We were primarily interested in between-group differences in the association between mind-wandering frequency and network interactions in patients, to determine the features uniquely related to hallucination predisposition in Parkinson's disease. Based on previous work, we anticipated that connectivity between the default network and visual areas would be related to elevated mind-wandering

in patients with visual hallucinations. In our study, this is operationalised more specifically as the prediction that hallucinating patients would show a stronger (positive) association between visual-default network coupling and mind-wandering frequency. This would be consistent with increased mind-wandering in hallucinating patients being associated with greater visual-default network coupling. Contrasts with healthy controls were of secondary interest and were used to inform differential patterns of brain connectivity and behaviour potentially indicative of differential disease trajectories in the Parkinson's disease groups. Overall, our aim was to investigate whether elevated mind-wandering and its associated neural correlates may be identifiable traits in a clinical population prone to visual hallucinations.

2. Methods and materials

We report how we determined our sample size, all data exclusions (if any), all inclusion/exclusion criteria, whether inclusion/exclusion criteria were established prior to data analysis, all manipulations, and all measures in the study.

2.1. Case selection

Thirty-eight individuals with Parkinson's disease were recruited from the Parkinson's disease research clinic, University of Sydney, Australia. Question two of the MDS-UPDRS (Goetz et al., 2008) was used to identify visual hallucinations (i.e., "Over the past week have you seen, heard, smelled or felt things that were not really there? If yes, examiner asks the patient or caregiver to elaborate and probes for information"). If an individual scored ≥ 1 on this item and if their subsequent description was consistent with visual hallucinatory phenomena, including minor (passage or illusions) or complex hallucinations, they were included in the hallucinating group. This resulted in 18 patients with hallucinations and 20 without hallucinations.

All individuals with Parkinson's disease satisfied the United Kingdom Parkinson's Disease Society Brain Bank criteria and did not meet criteria for dementia, scoring above the recommended cut-off of ≥ 26 on the Montreal Cognitive Assessment (MoCA) (Dalrymple-Alford et al., 2010). Motor severity was determined by the Hoehn and Yahr Scale and the unified Parkinson's disease rating scale (MDS UPDRS-III) (Goetz et al., 2008). Mood was assessed via the self-reported Beck Depression Inventory-II (BDI-II) (Beck, Steer, & Brown, 1996). General neuropsychological measures assessed working memory (backwards digit-span), attentional set-shifting (Trail Making Test, Part B minus Part A), and memory (story retention on the Logical Memory component of the Wechsler Memory Scale). All clinical and neuropsychological assessments, as well as neuroimaging, were performed with participants on their regular antiparkinsonian medication. Dopaminergic dose equivalence (DDE) scores were calculated, and no participants were taking antipsychotic medication or cholinesterase inhibitors. All individuals with Parkinson's disease underwent neuroimaging.

Forty age- and education-matched healthy controls were included to provide a large normative dataset for the mind-

wandering task (a subset of 20 of these controls also underwent neuroimaging). Controls were screened for a history of neurological or psychiatric disorders. The study was approved by the local Ethics Committees and all participants provided written informed consent in accordance with the Declaration of Helsinki. See Table 1 for demographic details and clinical characteristics. No part of the study procedures was pre-registered prior to the research being conducted. Sample sizes were determined based on a convenience sample of patients and controls enrolled in ongoing studies and regular clinics, and inclusion criteria were established prior to recruitment. The conditions of our ethics approval do not permit public archiving of individual anonymised raw data. Readers seeking access to the data should contact the corresponding author (CO). Access will be granted to named individuals in accordance with ethical procedures governing the reuse of sensitive data. Specifically, requestors must meet the following conditions to obtain the data: completion of a formal data sharing agreement; use of the data only for non-commercial research purposes.

2.2. Mind-wandering experimental task & scoring procedures

The thought-sampling task was designed for use in patient populations with cognitive impairment and has previously been validated in older adults (O'Callaghan et al., 2015). The task involved 9 trials. In each trial, a 2-dimensional coloured shape (e.g., blue square, yellow circle, etc.) was presented on the screen for varying durations (Short: ≤ 20 sec, Medium: 30–60 sec, Long: ≥ 90 sec). At the outset, participants were told

Table 1 – Mean (standard deviation) values for demographics, clinical characteristics and background neuropsychology.

Demographics, clinical characteristics & general neuropsychology	Controls	PD + VH	PD-VH	p value
N	40	18	20	–
Sex (M:F)	21:19	14:4	17:3	–
Age	66.3 (6.2)	67.5 (6.7)	63.7 (6.6)	n.s.
Education	14.7 (2.4)	13.3 (3.3)	14.6 (2.4)	n.s.
MoCA	–	27.9 (1.3)	27.9 (1.1)	n.s.
Duration (yrs diagnosed)	–	7.6 (5.0)	5.7 (3.2)	n.s.
DDE (mg/day)	–	832.8 (395.7)	691.5 (440.5)	n.s.
Hoehn & Yahr stage	–	2.2 (.57)	2.1 (.36)	n.s.
UPDRS III	–	33.4 (15.9)	28.4 (13.4)	n.s.
BDI-II	–	11.6 (10.2)	8.1 (5.9)	n.s.
Neuropsychology				
TMT B-A (seconds)	–	76.6 (57.3)	35.2 (22.7)	**
Digit span backward	–	7.0 (1.6)	6.6 (2.3)	n.s.
Logical memory % retention	–	75.6 (17.7)	87.1 (10.9)	*

n.s. = non significant; ** = $p < .01$; * = $p < .05$. Significance tests refer to two-sided Welch's Independent samples t-tests or one-way ANOVAs with Tukey post hoc tests. PD-VH = non-hallucinators; PD + VH = hallucinators; MoCA = Montreal Cognitive Assessment; DDE = Dopaminergic dose equivalence; BDI-II = Beck Depression Inventory-II; TMT B-A = Trail Making Test part B minus part A.

that they would be shown a series of shapes and to just relax and continue looking at the shape. Immediately after each shape was presented, the participant was prompted to describe aloud what they were thinking about during the presentation of the shape stimulus.

Participants' reported thoughts were scored on a continuum, ranging from Level 1 to Level 4. Level 1 represents stimulus-bound/impoverished thought, including thinking about the stimulus, e.g., "a blue square," or describing thinking of "nothing". In contrast, Level 4 responses bear no obvious relationship to the stimulus, the task at hand, or the immediate testing environment, indicating thought content that is stimulus-independent and task-unrelated – a class of cognition often referred to as mind-wandering (Smallwood & Schooler, 2015). Examples of mind-wandering include, "I thought about the people I saw today and how we chatted with them outside the unit"; "I thought of a sailing boat in the Greek Islands". Levels 2 and 3 represent intermediary responses, which do not qualify as fully-fledged instances of mind-wandering, as they still bear a discernible relationship to the presented stimulus or immediate environment. These levels capture the transition from stimulus-related to increasingly stimulus-independent responses. See Fig. 1 and see Supplementary Material for a detailed description of the scoring levels and example responses. Task materials and instructions are available online at https://github.com/claireocallaghan/MindWandering_PD_VH.

The final score awarded for each trial was the highest level achieved on that trial, ranging from 1 to 4. Instances of each scoring level achieved were counted across the 9 trials and transformed into a total percentage for each level during the task (i.e., total instances of levels 1, 2, 3 or 4 divided by 9 multiplied by 100). The primary analysis focused on differences in mind-wandering frequency between Parkinson's disease with hallucinations, Parkinson's disease without hallucinations, and controls. Therefore, the proportion of Level 4 responses – referred to as the mind-wandering frequency – was compared across the groups and used as a covariate in the neuroimaging analyses. Secondary behavioural analyses were conducted to determine overall performance of the three groups on the task. This involved comparing the proportion of responses that each group achieved across the four scoring levels of the task.

2.3. Statistical analysis

Analyses were performed in R version 3.3.1 (<http://www.r-project.org/>). For performance on the mind-wandering task, homogeneity of variance was verified using Levene's test and values were checked for normality by inspection of normal Q–Q plots and the Shapiro–Wilk test. To reduce skew in the data, a square root transformation was applied to the mind-wandering scores. Scores were then analysed using mixed-effects ANOVAs, implemented in the "lme4" package (Bates, Mächler, Bolker, & Walker, 2014). Where appropriate, group, score level and trial duration were specified as fixed effects, and subject was entered as a random effect. Where a factor only had three levels, post hoc analysis of significant main effects was performed using the Fisher's Least Significant Difference (LSD) procedure (Cardinal & Aitken, 2013). Where

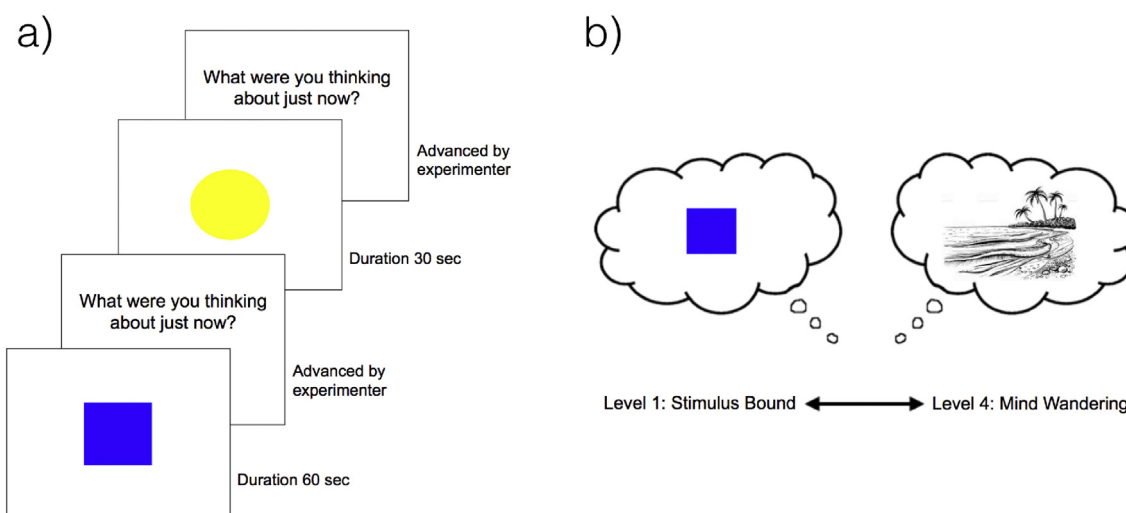


Fig. 1 – Task structure and schematic of scoring system. a) An example of two trials in the thought sampling task; b) Responses were scored from 1 to 4, with Level 1 consisting of a stimulus-bound response, such as reporting thoughts about the displayed shape, and Level 4 corresponding to thoughts completely unrelated to the task or immediate environment.

factors had more than three levels, main effects were analysed using post-hoc t-tests with the Sidak correction for multiple comparisons. In these cases, post-hoc analyses of interactions were conducted using separate univariate ANOVAs to establish simple effects. Behavioural analysis scripts are available at https://github.com/claireocallaghan/MindWandering_PD_VH.

2.4. Imaging acquisition

The 38 individuals with Parkinson's disease and a subset of 20 controls underwent magnetic resonance imaging (MRI) to acquire T1-weighted structural images and resting-state blood-oxygenation level dependent (BOLD) functional scans. Imaging was conducted on a 3 T MRI (General Electric, Milwaukee, USA). Whole-brain three dimensional T1-weighted sequences were acquired as follows: coronal orientation, matrix 256×256 , 200 slices, $1 \times 1 \text{ mm}^2$ in-plane resolution, slice thickness 1 mm, TE/TR = 2.6/5.8 msec. T2*-weighted echo planar functional images were acquired in interleaved order with repetition time (TR) = 3 sec, echo time (TE) = 32 msec, flip angle 90°, 32 axial slices covering the whole brain, field of view (FOV) = 220 mm, interslice gap = .4 mm, and raw voxel size = $3.9 \times 3.9 \times 4 \text{ mm}$ thick. Resting state scan acquisition lasted 7 min. During the resting-state scan, patients were instructed to lie awake with their eyes closed.

2.5. Resting state fMRI preprocessing and motion correction

Functional magnetic resonance imaging (fMRI) pre-processing and analysis was performed using Statistical Parametric Mapping software (SPM12, Wellcome Trust Centre for Neuroimaging, London, UK, <http://www.fil.ion.ucl.ac.uk/spm/software/>). We used a standard pre-processing pipeline that included slice-timing correction, rigid body realignment,

spatial smoothing with a Gaussian kernel (FWHM) of 6 mm, and registration of the anatomical scans to the Montreal Neurological Institute standard brain space. Pre-processed images were imported into CONN: The Functional Connectivity toolbox (<https://www.nitrc.org/projects/conn>) in MATLAB for all functional connectivity analyses.

To compensate for motion-related artefacts we performed the “scrubbing” procedure in an effort to effectively remove time points with excessive head motion (i.e., framewise displacement in x, y, or z direction > 2 mm from the previous frame; global intensity > 9 standard deviations from mean image intensity of the entire resting state scan) (Power, Barnes, Snyder, Schlaggar, & Petersen, 2012). This approach (Artefact Detection Tools; <https://www.nitrc.org/projects/artefact-detect>) effectively removes outlier frames by including them as dummy-coded regressors during the denoising procedure, so as to avoid discontinuities in the time-series. We also tested for significant differences in maximum motion and number of frames scrubbed between the Parkinson's disease groups. No significant differences were found in maximum framewise displacement ($p = .50$), maximum frames scrubbed ($p = .25$), mean framewise displacement ($p = .59$), or mean number of frames scrubbed ($p = .25$). Other noise sources in the BOLD signal (i.e., from white matter and cerebrospinal fluid) were corrected for by using a principle component-based ‘aCompCor’ method (Whitfield-Gabrieli & Nieto-Castañón, 2012). We applied a band-pass filter (.008–.09 Hz) to limit the effect of low-frequency drift and high-frequency noise on the BOLD signal time-series.

2.6. Relationship between inter-network functional connectivity and mind-wandering frequency

To assess differences in the association between mind-wandering frequency and inter-network functional

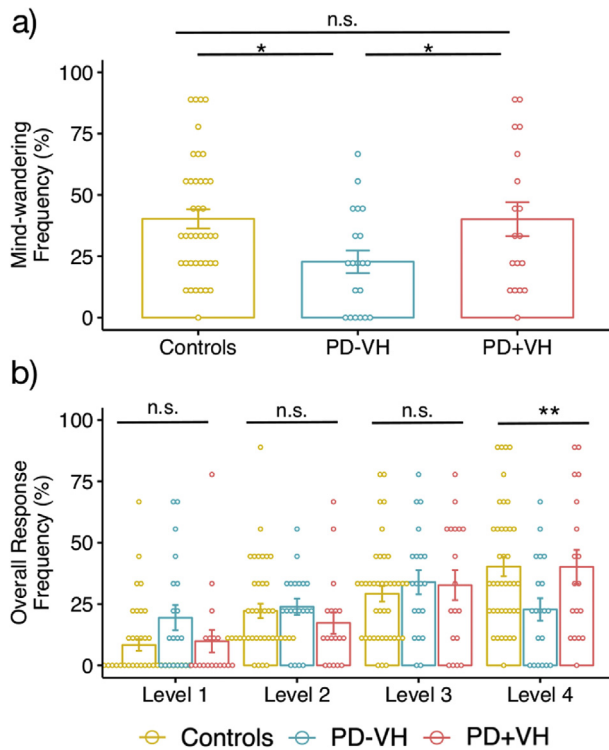


Fig. 2 – Performance on the mind-wandering task. a) Mind-wandering frequency. Parkinson's disease with hallucinations and controls both exhibited higher frequencies of mind-wandering (i.e., had significantly more Level 4 responses on the thought sampling task) compared to non-hallucinators; b) Frequency of responses at each of the four scoring levels. Group differences were only found for Level 4 responses (i.e., mind-wandering frequency). Frequencies of responses across the three groups were not significantly different for Levels 1, 2 and 3. Error bars show standard error of the mean; PD-VH = non-hallucinators; PD + VH = hallucinators; n.s. = not significant; * = $p < .05$; ** = $p < .01$.

connectivity across the two patient groups and controls, we used 12 individual intrinsic connectivity network regions of interest, which were defined on a functional basis (Shirer, Ryali, Rykhlevskaia, Menon, & Greicius, 2012). The networks included in our analysis are shown in Fig. 3a. We did not include the auditory or language networks, as the auditory network is not implicated in network-based models of visual hallucinations and, as others have noted, there is substantial overlap between the language network and aspects of the default/limbic networks (Zabelina & Andrews-Hanna, 2016). We corrected for multiple comparisons by using a False Discovery Rate (FDR) threshold of $p < .05$.

Based on the results from the above analysis, we performed a post-hoc seed-to-voxel functional connectivity analysis. This was to investigate whether the between-group differences (i.e., Parkinson's disease hallucinators vs non-hallucinators and controls vs non-hallucinators) that were identified in the association between mind-wandering and primary visual network (V1) functional connectivity extended

beyond the dorsal default network. We calculated a correlation between the average filtered BOLD signal in V1 and all other voxels in the brain for each group. Then, a between-group contrast of the regression coefficient, capturing the association between mind-wandering frequency and seed-to-voxel functional connectivity, was carried out with a height threshold of $p < .001$ and a cluster-size threshold of $p < .05$, corrected using FDR. Seed-to-voxel statistical tests for this post-hoc analysis were one-sided, as we were specifically interested in the spatial boundaries of the directional V1 connectivity demonstrated in the prior inter-network analysis.

3. Results

3.1. Demographics, clinical characteristics and background neuropsychology

Parkinson's disease and control groups were matched for age and education. The Parkinson's disease groups were matched on all demographic variables; however, the hallucinating group performed worse than the non-hallucinators on measures of attentional set-shifting and memory retention, although working memory performance did not differ significantly between the groups. See Table 1 and see Supplementary Material for detailed results.

3.2. Hallucinators had a higher frequency of mind-wandering compared to non-hallucinators

As shown in Fig. 2a, mind-wandering occurred significantly more frequently in hallucinators than non-hallucinators. There was a main effect of group [$F(2,75) = 5.34, p < .01$], and post-hoc comparisons revealed that controls and Parkinson's disease with hallucinations did not differ, but both groups exhibited higher frequencies of mind-wandering than the non-hallucinating group (Fisher's LSD, $p < .05$).

3.3. Overall performance on the mind-wandering task revealed that significant group differences emerged exclusively for mind-wandering responses

Fig. 2b shows the frequency of responses across all scoring levels, ranging from Level 1 (stimulus-bound thought) to Level 4 (mind-wandering). No significant main effect of group was evident [$F(2,300) = .28, p = .76$], however, a significant main effect of response level [$F(3,300) = 28.57, p < .0001$] was observed. Post hoc t-tests with Sidak correction showed that, regardless of group, Level 1 was the least frequent response, relative to Levels 2, 3 and 4 (p values $< .0001$), and higher frequencies of Level 4 were obtained relative to Level 2 ($p < .01$). The other Levels did not differ significantly from each other (p values $> .05$).

The Level \times Group interaction was significant [$F(6,300) = 3.06, p < .01$]. We followed this interaction with tests of simple effects to determine whether the groups differed in their response frequencies at any Level apart from Level 4, which we showed in our focused analysis above. Follow-up tests of simple effects revealed the frequency of responses

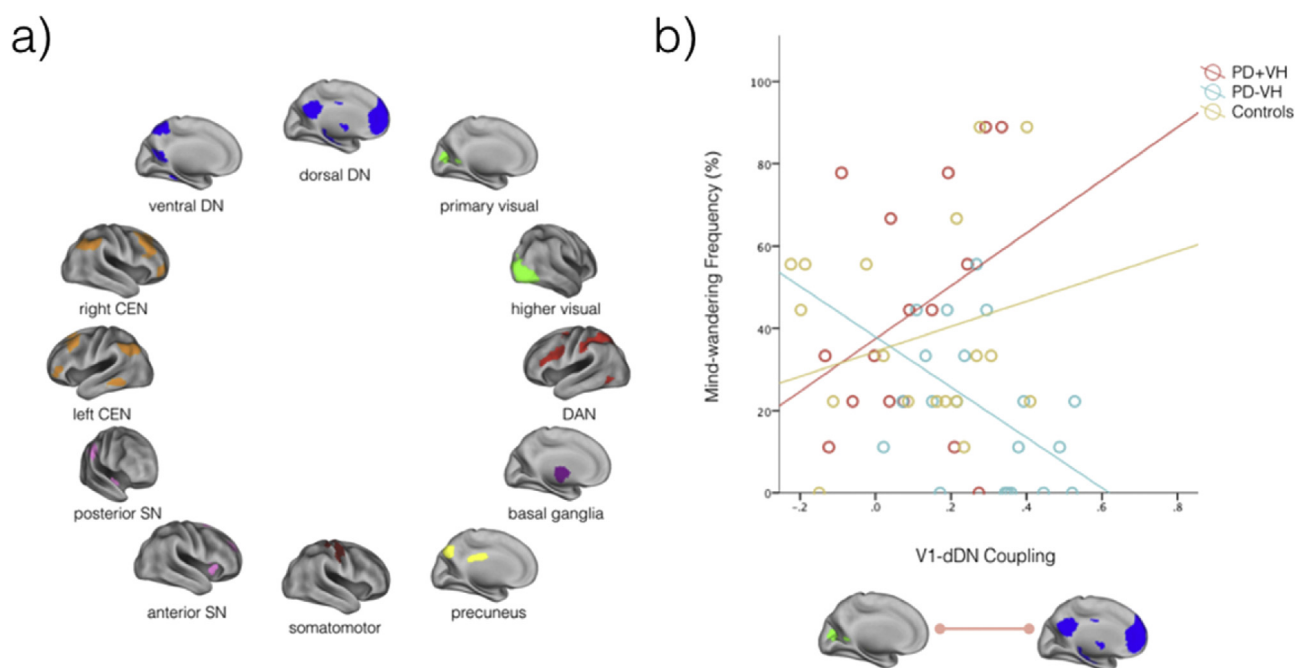


Fig. 3 – Association between mind-wandering frequency and inter-network coupling. a) Stanford atlas networks included in the analysis examining the association between inter-network functional connectivity and mind-wandering frequency; b) Individuals with visual hallucinations (PD + VH) had a significantly stronger positive association between mind-wandering frequency and V1-dDN coupling, compared to those without hallucinations (PD-VH). dDN = dorsal default network; vDN = ventral default network; CEN = central executive network; SN = salience network; DAN = dorsal attention network.

across the groups did not differ significantly for Level 1 [$F(2,75) = 2.58, p = .08$], Level 2 [$F(2,75) = 1.16, p = .32$] or Level 3 [$F(2,75) = .20, p = .82$]. The groups only differed significantly in their Level 4 responses (i.e., % mind-wandering frequency) [$F(2,75) = 5.34, p < .01$]. In addition, all groups showed an increased tendency towards mind-wandering on longer trials (See [Supplementary material and Figure S1](#) for analyses and results), replicating previous studies using the task (Geffen et al., 2017; O’Callaghan, Shine, Hodges, Andrews-Hanna, & Irish, 2019).

3.4. A stronger association was identified between primary visual-dorsal default inter-network coupling and mind-wandering frequency in Parkinson’s disease hallucinators versus non-hallucinators

Results of the inter-network coupling analysis were consistent with our prediction of a stronger association between primary visual network and dorsal default network coupling and the degree of mind-wandering in patients with visual hallucinations. Of all network pairs (Fig. 3a), only primary visual-dorsal default network functional connectivity (i.e., primary visual-dorsal default network coupling) and its association with mind-wandering frequency differed significantly between patient groups ($p < .05$, FDR; Fig. 3b). Parkinson’s disease hallucinators had a significantly stronger positive association between mind-wandering frequency and primary visual-dorsal default network coupling compared with non-hallucinators. Controls also exhibited a stronger

positive association between primary visual-dorsal default network coupling and mind-wandering frequency relative to Parkinson’s disease non-hallucinators, but this did not survive correction for contrasts among all network pairs. Only at a more lenient correction threshold – correcting for the number of contrasts between primary visual and all other networks (rather than all comparisons made) – was the association between controls and non-hallucinators statistically different ($p < .05$, FDR; Fig. 3b). As this latter result only survived a more liberal correction for multiple comparisons, we emphasise caution in its interpretation. Statistically significant differences did not emerge between Parkinson’s disease hallucinators and controls.

3.5. Follow-up seed-to-voxel functional connectivity with a primary visual network seed

When the primary visual network was used as a seed, the between-group difference in the association between primary visual network coupling and mind-wandering frequency included brain regions both within and beyond the dorsal default network. Relative to the non-hallucinating group, hallucinators displayed a significantly stronger association between mind-wandering frequency and connectivity of the primary visual network seed to dorsal default network regions (posterior cingulate cortex, medial prefrontal cortex and left inferior parietal lobule), the inferior frontal gyrus, orbito-frontal cortex and high-level visual regions (fusiform gyrus/inferior temporal gyrus). See Fig. 4 and Table 2. The same

analysis contrasting controls and Parkinson's disease non-hallucinators revealed similar core dorsal default network regions (posterior cingulate cortex and medial prefrontal cortex) as well as the orbitofrontal cortex. However, instead of unilateral inferior parietal lobule, changes were found with bilateral angular gyrus connectivity to primary visual cortex. Furthermore, the fusiform gyrus did not display altered primary visual cortex connectivity in this contrast (See Figure S2 and Table S1).

4. Discussion

Our results provide empirical evidence of a link between mind-wandering and hallucinations, revealing that Parkinson's disease patients with visual hallucinations exhibit increased mind-wandering relative to non-hallucinating patients. Elevated mind-wandering may therefore represent a cognitive correlate of the excessive top-down influence upon perception that has previously been hypothesised in Parkinson's disease visual hallucinations. Our resting state analysis revealed a route by which mind-wandering may impact upon early visual processing, as mind-wandering frequency was associated with stronger coupling between the primary visual and dorsal default networks in the Parkinson's disease hallucinators. Together, these findings uncover trait

Table 2 – Peak coordinates from seed-to-voxel connectivity associated with mind-wandering frequency in Parkinson's disease hallucinators versus non-hallucinators.

Brain Region	x	y	z	Voxels
R Orbitofrontal Cortex	40	24	-22	136
L Inferior Frontal Gyrus	-48	26	14	197
L Medial Prefrontal Cortex	-16	54	00	132
	-06	54	18	129
R Fusiform Gyrus	50	-46	-24	94
L Posterior Cingulate Cortex	-10	-42	36	341
L Inferior Parietal Lobule	-44	-46	26	347

R = right; L = left; x, y, z = co-ordinates in MNI standard space.

characteristics: increased mind-wandering frequency related to default-visual network coupling, which may predispose the Parkinsonian brain to hallucinate.

Behavioural analysis of the thought-sampling task revealed that mind-wandering frequency was significantly higher in the hallucinating patient group compared to the non-hallucinating patient group; no significant differences were found between the hallucinating patient group and controls. In contrast, the frequency of mind-wandering was lower in non-hallucinating patients relative to controls, replicating findings from an independent Parkinson's disease

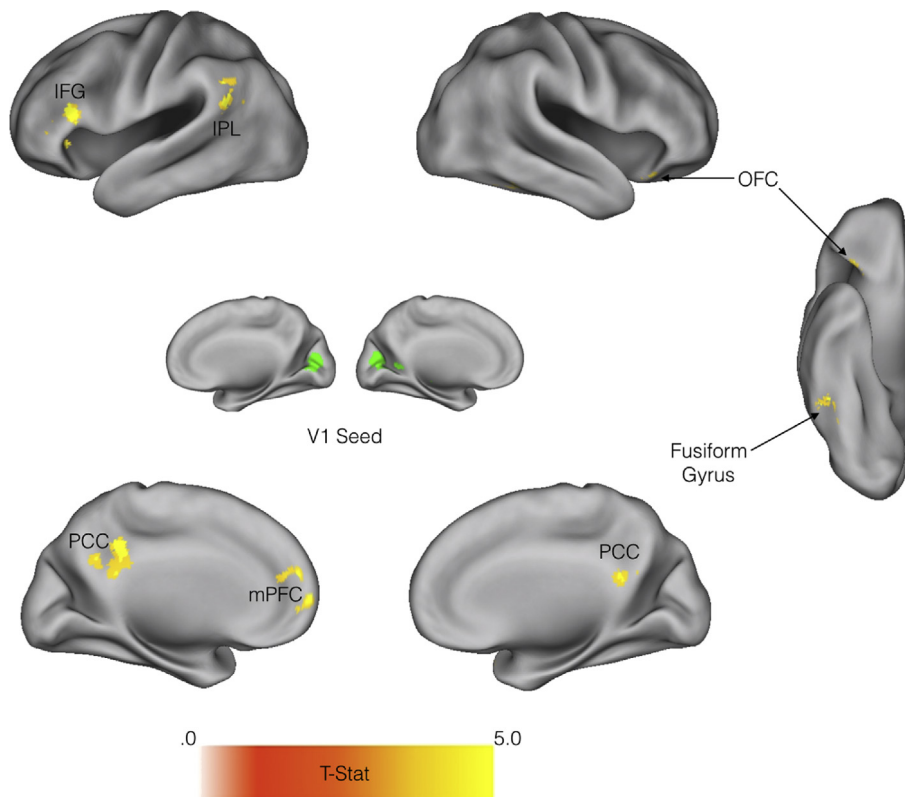


Fig. 4 – Seed-to-voxel connectivity associated with mind-wandering frequency, between V1 seed and whole brain in Parkinson's disease hallucinators versus non-hallucinators. Individuals with hallucinations had a significantly stronger association between mind-wandering frequency and connectivity between V1 and areas of the dorsal default network (PCC, mPFC and IPL), orbitofrontal cortex, inferior frontal gyrus, and high-level visual regions (fusiform gyrus/inferior temporal gyrus), relative to non-hallucinating individuals. PCC = posterior cingulate cortex; mPFC = medial prefrontal cortex; IPL = inferior parietal lobule; OFC = orbitofrontal cortex; IFG = inferior frontal gyrus.

cohort tested on the same task (Geffen et al., 2017). Preserved global cognitive function does not account for the higher levels of mind-wandering in Parkinson's disease hallucinators, as their cognitive abilities were either similar to the non-hallucinators, or mildly reduced.

These findings suggest that reductions in mind-wandering may be common in Parkinson's disease without hallucinations, consistent with observations in ageing (Irish, Goldberg, Alaeddin, O'Callaghan, & Andrews-Hanna, 2019; Jackson & Balota, 2012; McVay, Meier, Touron, & Kane, 2013) and dementia populations including Alzheimer's disease and frontotemporal dementia (Gyurkovics, Balota, & Jackson, 2018; O'Callaghan et al., 2019). By contrast, relatively preserved mind-wandering may be a specific feature of the hallucinating phenotype. In Parkinson's disease hallucinations, this excess of top-down influence is suggested to occur in the context of compromised primary visual and attentional systems, resulting in poor quality sensory evidence accumulation (O'Callaghan et al., 2017; Shine et al., 2014). More broadly, this account is consistent with the idea that relatively strong prior beliefs are implicated in the genesis of hallucinations across modalities and across disorders (Corlett et al., 2018).

Our correlations with imaging reveal a route by which overly strong prior beliefs or internally generated imagery might influence the early perceptual process. The primary finding was a significantly increased association between mind-wandering frequency and primary visual-dorsal default network coupling in the Parkinson's disease hallucinators, relative to the non-hallucinating patients. This result confirmed that an increased propensity to mind-wander in the hallucinating patients was associated with greater connectivity between the default network and early visual regions. It is important to note that we are not describing an absolute difference in visual-default network connectivity strength between hallucinators and non-hallucinators, but rather that the association between this connectivity and mind-wandering is stronger for the hallucinators. This line of enquiry speaks to how differences in the brain's resting state patterns may contribute to differences in cognitive processes. Controls showed a similar positive relationship between mind-wandering frequency and primary visual-dorsal default network coupling, however this association did not differ statistically from Parkinson's disease non-hallucinators. Indeed, in healthy individuals, coupling between the default network and visual cortex has previously been associated with the degree of vivid detail experienced during mind-wandering episodes (Turnbull et al., 2019). Considering the similar patterns between hallucinating patients and controls, this emphasises the overall interpretation of our results as one of a *preserved*, rather than *increased*, association between mind-wandering frequency and primary visual-default network in Parkinson's disease hallucinations. However, unlike controls, this relative preservation is occurring in the context of impaired attention and primary visual dysfunction. When confronted with ambiguous sensory input due to these impairments, Parkinson's disease hallucinators are therefore more likely to rely on top-down predictions to resolve a

percept. This is consistent with previous evidence of increased coupling between the default network and primary visual regions during recorded misperceptions in Parkinson's disease hallucinators (Shine, Muller et al., 2015).

Our results show that, in the context of established Parkinson's disease, patients with visual hallucinations exhibit greater mind-wandering frequency *and* a stronger association between mind-wandering frequency and primary visual-dorsal default network coupling, relative to non-hallucinators. However, when contrasted with healthy controls, Parkinson's disease hallucinators were not statistically distinguishable on either of these variables. In light of the present evidence, we suggest two distinct trajectories in Parkinson's disease: in Parkinson's disease with hallucinations, forms of internally generated mental imagery are relatively better preserved, compared to impaired sensory and attentional abilities, which leads them to dominate perception; in Parkinson's disease without hallucinations, such forms of cognition are compromised, consistent with a more generic course seen in ageing (Maillet & Schacter, 2016) and exacerbated in several neurodegenerative diseases (Geffen et al., 2017; Gyurkovics et al., 2018; O'Callaghan et al., 2019).

Results from the post-hoc seed-to-voxel analysis confirmed a significant difference between the patient groups in the association between mind-wandering frequency and connectivity between the primary visual network and more restricted areas of the dorsal default network (posterior cingulate cortex, medial prefrontal cortex, right temporo-parietal junction and left inferior parietal lobule). Similar to the inter-network analysis, here we describe how the association between mind-wandering frequency and the connectivity between V1 and all other voxels in the brain differs between the groups. This analysis enabled a more specific localisation of regions that might mediate the relationship between mind-wandering and primary visual network connectivity. Our findings implicated specific regions within the default network, including the medial prefrontal cortex, previously identified as a source of top-down influence over visual perception (Bar et al., 2006; Kveraga et al., 2007; Summerfield et al., 2006). Previous work has also shown that stronger resting state coupling between the posterior cingulate/retrosplenial cortex and primary visual cortex correlates with the ability to actively imagine the future, which presumably relies upon scene construction processes (i.e., internally generating a scene or event) (Villena-Gonzalez et al., 2018). We also identified regions outside the default network, including the fusiform gyrus/inferior temporal gyrus, a higher-level region in the ventral visual processing stream; the inferior frontal gyrus, which has been identified (via effective connectivity) as a source of directed top-down influence over V1 during both mental imagery and perception (Dijkstra, Zeidman, Ondobaka, van Gerven, & Friston, 2017); and the orbitofrontal cortex, a source of top-down predictions that refine object recognition during early visual processing (Bar, 2003; Bar et al., 2006; Chaumon et al., 2014). In the context of existing literature, the neural correlates

identified in our analyses overlap with those that have been shown, in healthy people, to be important for coordinating internally generated predictions that influence both mental imagery and early visual perception.

Based on patterns of functional connectivity across the entire cerebral cortex, the default network is located on the opposite end of a principal connectivity gradient from brain regions supporting primary perceptual processing (e.g., the primary visual network) (Huntenburg et al., 2018; Margulies et al., 2016; Murphy et al., 2018). Our results suggest that, in Parkinson's disease with hallucinations, individual differences in mind-wandering frequency are associated with increasing loss of this inherent functional separation between the default network and primary visual areas. This is consistent with a previous task-based study, which found increased visual network coupling to the default network during misperceptions in Parkinson's disease with hallucinations (Shine, Muller et al., 2015). A combination of relatively increased internally generated thought and imagery (as compared to non-hallucinating patients), and an increased association between mind-wandering and default-visual network coupling, may therefore manifest in a neurocognitive endophenotype that is prone to hallucinate. Intriguingly, increased psychopathological features in the general population have been associated with elevated visual network-default network coupling (Elliott, Romer, Knodt, & Hariri, 2018), suggesting that the loss of functional segregation between these networks may play a role across neuropsychiatric disorders.

Given that a cardinal feature of mind wandering is its relative stimulus-independence (Christoff et al., 2016; Mason et al., 2007; Seli et al., 2018), from a predictive processing framework this cognitive phenomenon would depend upon the activation (and possibly the exploration/finishing) of expectations or predictions (i.e., priors). Although the functional importance of mind-wandering remains unclear, it is possible that predictive model optimisation via the pruning of priors (i.e., reduction of model complexity or “overfitting”) during mind-wandering is analogous to predictive processing accounts for the functional importance of dreaming (Hobson & Friston, 2012), but occurs during the waking state with the associated benefits for adaptive fitness.

Previous work has shown that Parkinson's disease patients who experience visual hallucinations are unable to contextually modulate sensory evidence accumulation (O'Callaghan et al., 2017). As the association between mind-wandering frequency and primary visual-dorsal default network coupling differentiated Parkinson's disease patient groups with and without visual hallucinations, this pattern of network coupling may predispose the Parkinsonian brain to hallucinate by allowing relatively strong priors to dominate imprecise sensory evidence during the act of visual perception. Thus, while control subjects do not have significantly different mind-wandering frequencies or associations between mind-wandering frequency and primary visual-dorsal default network coupling compared with Parkinson's disease hallucinators, they may not be prone to visual hallucinations due to intact mechanisms for the attribution of context-dependent precision estimation of sensory evidence. In

keeping with this, previous work has suggested that a breakdown in the co-ordination of networks that support attention and saliency (including the dorsal and ventral attention networks) may be a crucial factor contributing to the inability to resolve imprecise sensory input in Parkinson's disease hallucinations (Shine, Keogh et al., 2015; Shine, Muller et al., 2015). In our cohort, we found that attentional set-shifting was reduced in the hallucinating patients, relative to the non-hallucinators. This is consistent with impairments in attention and saliency systems in hallucinating patients, which would contribute to difficulties in processing visual information. It is also tempting to speculate that an inability to flexibly shift focus could contribute to getting ‘stuck’ in a mind-wandering mode. However, whether these patients show impairments in the dynamic ability to shift in and out of mind-wandering states remains an outstanding question for future research. Our results highlight the importance of future studies that will directly examine the interplay between relatively preserved priors and imprecise sensory evidence in Parkinson's disease hallucinations. One possible line of investigation is to use established tasks for conditioned hallucinations, which have been effective at identifying both relatively strong priors and neural regions supporting these priors in populations with auditory hallucinations (Corlett et al., 2018; Powers, Mathys, & Corlett, 2017, 2016).

Our interpretation of these results was guided by current models of Parkinson's disease hallucinations that emphasise increased top-down influences over perception, in the context of reduced attentional and visual processing (Collerton et al., 2005; Diederich et al., 2005; Shine et al., 2014). However, other frameworks have been proposed, including the intrusion of REM sleep visual imagery into wakefulness (Arnulf et al., 2000) or a reality monitoring deficit in the ability to determine whether imagery is self-generated or drawn from an external source (Barnes, Boubert, Harris, Lee, & David, 2003). These frameworks are broadly consistent with our findings, as misattribution of internally generated imagery reflects a process of over-reliance on top-down expectations during perception. And, intriguingly, other researchers have noted shared phenomenological and neural characteristics across mind-wandering and dreaming (Fox, Nijeboer, Solomonova, Domhoff, & Christoff, 2013), suggesting that hallucinations, dreaming and mind-wandering may exist on a related dimension, becoming less distinctive in certain neuropsychiatric conditions.

In summary, our study has identified mind-wandering frequency as a potential behavioural correlate of the pathological default network, which has previously been implicated in the pathogenesis of visual hallucinations in Parkinson's disease. To our knowledge, these findings provide the first evidence of a relationship between mind-wandering and visual hallucinations in a clinical population. Higher levels of mind-wandering indicate an increased propensity for the internal generation of detailed mental imagery, formation of contextual associations and spontaneous thought – all top-down processes that could furnish the content of visual hallucinations. Our results identify connectivity between the default network and early visual regions as a putative neural

substrate that may be associated with excessive influence from these top-down processes over perception. Thus, the preservation of florid abstract and internally generated cognition in the context of the Parkinson's disease might result in visual hallucinations where healthy individuals experience only the vivid images of the mind's eye.

Open practices

The study in this article earned an Open Materials badge for transparent practices. Materials and data for the study are available at https://github.com/claireocallaghan/Mind_Wandering_PD_VH.

Declaration of Competing Interest

The authors report no conflicts of interest.

CRediT authorship contribution statement

Ishan C. Walpola: Conceptualization, Methodology, Writing - original draft, Investigation, Formal analysis, Visualization. **Alana J. Muller:** Writing - review & editing, Investigation. **Julie M. Hall:** Writing - review & editing, Investigation. **Jessica R. Andrews-Hanna:** Conceptualization, Writing - review & editing. **Muireann Irish:** Conceptualization, Methodology, Writing - review & editing. **Simon J.G. Lewis:** Funding acquisition, Supervision, Conceptualization, Writing - review & editing, Resources, Project administration. **James M. Shine:** Supervision, Conceptualization, Methodology, Writing - review & editing, Investigation, Project administration. **Claire O'Callaghan:** Supervision, Conceptualization, Methodology, Writing - original draft, Writing - review & editing, Investigation, Formal analysis, Visualization, Project administration.

Acknowledgments

We thank Kelly Diederer for providing valuable comments on the manuscript. AJM is supported by an Australian Postgraduate Award through the University of Sydney. JMH is supported by a Western Sydney University Postgraduate Award. JAH is supported by a National Institutes of Aging Arizona Alzheimer's Disease Core Center grant (P30 AG019610). MI is supported by an Australian Research Council Future Fellowship (FT160100096) and an Australian Research Council Discovery Project (DP180101548). SJGL is supported by an NHMRC-ARC Dementia Fellowship (#1110414). JMS is supported by a National Health and Medical Research Council CJ Martin Fellowship (1072403). CO is supported by a National Health and Medical Research Council Neil Hamilton Fairley Fellowship (1091310) and by the Wellcome Trust (200181/Z/15/Z). The study was supported by a Seed Grant from Parkinson's NSW and funding to Forefront, a collaborative research group dedicated to the study of non-Alzheimer disease degenerative dementias,

from the National Health and Medical Research Council of Australia program grant (#1037746 and #1095127).

Supplementary data

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.cortex.2019.12.023>.

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