Visual hallucinations in Lewy body disease relate to Lewy bodies in the temporal lobe

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Summary
Consensus opinion characterizes dementia with Lewy bodies (DLB) as a progressive dementing illness, with significant fluctuations in cognition, visual hallucinations and/or parkinsonism. When parkinsonism is an early dominant feature, consensus opinion recommends that dementia within the first year is necessary for a diagnosis of DLB. If dementia occurs later, a diagnosis of Parkinson’s disease with dementia (PDD) is recommended. While many previous studies have correlated the neuropathology in DLB with dementia and parkinsonism, few have analysed the relationship between fluctuating cognition and/or well-formed visual hallucinations and the underlying neuropathology in DLB and PDD. The aim of the present study was to determine any relationship between these less-studied core clinical features of DLB, and the distribution and density of cortical Lewy bodies (LB). The brains of 63 cases with LB were obtained over 6 years following population-based studies of dementia and parkinsonian syndromes. Annual, internationally standardized, clinical assessment batteries were reviewed to determine the presence and onset of the core clinical features of DLB. The maximal density of LB, plaques and tangles in the amygdala, parahippocampal, anterior cingulate, superior frontal, inferior temporal, inferior parietal and visual cortices were determined. Current clinicopathological diagnostic criteria were used to classify cases into DLB (n = 29), PDD (n = 18) or parkinsonism without dementia (n = 16) groups. Predictive statistics were used to ascertain whether fluctuating cognition or visual hallucinations predicted the clinicopathological group. Analysis of variance and regressions were used to identify any significant relationship(s) between the presence and severity of neuropathological and clinical features. Cognitive fluctuations and/or visual hallucinations were not good predictors of DLB in pathologically proven patients, although the absence of these features early in the disease course was highly predictive of PDD. Cases with DLB had higher LB densities in the inferior temporal cortex than cases with PDD. There was no association across groups between any neuropathological variable and the presence or absence of fluctuating cognition. However, there was a striking association between the distribution of temporal lobe LB and well-formed visual hallucinations. Cases with well-formed visual hallucinations had high densities of LB in the amygdala and parahippocampus, with early hallucinations relating to higher densities in parahippocampal and inferior temporal cortices. These temporal regions have previously been associated with visual hallucinations in other disorders. Thus, our results suggest that the distribution of temporal lobe LB is more related to the presence and duration of visual hallucinations in cases with LB than to the presence, severity or duration of dementia.

Keywords: cortical Lewy bodies; dementia with Lewy bodies; fluctuating cognition; Parkinson’s disease; visual hallucinations

Abbreviations: CDR = clinical dementia rating; DLB = dementia with Lewy bodies; LB = Lewy bodies; NPV = negative predictive value; PDD = Parkinson’s disease with late dementia; PPV = positive predictive value

Introduction
Dementia with Lewy bodies (DLB) is a clinicopathological diagnosis for a progressive dementing illness with fluctuating cognition, visual hallucinations and/or parkinsonism in association with Lewy bodies (LB) in the brain (McKeith et al., 1996). However, LB pathology in the midbrain has been most closely associated with idiopathic Parkinson’s...
disease and is currently mandatory for a definitive clinicopathological diagnosis of this disease (Gelb et al., 1999). While there is still debate concerning the pathological overlap between these distinct clinical phenotypes, midbrain LB accumulate independently from those found in the cortex in DLB (Gómez-Tortosa et al., 1999) and high densities of temporal lobe LB differentiate Parkinson’s disease from cases with dementia (Harding and Halliday, 2001).

The consensus criteria for DLB require progressive cognitive decline, but do not require dementia as the initial symptom (McKeith et al., 1996, 1999). The criteria state: ‘The central feature required for a diagnosis of DLB is progressive cognitive decline of sufficient magnitude to interfere with normal social or occupational function. Prominent or persistent memory impairment may not necessarily occur in the early stages, but is usually evident with progression’ (McKeith et al., 1996). DLB is differentiated from Parkinson’s disease with late developing dementia (PDD) by the early presentation of cognitive decline within a year of the initial presentations of the disease (most often parkinsonism) (McKeith et al., 1996; McKeith, 1999).

Our recent comparison of the distribution and density of cortical LB in DLB, PDD and Parkinson’s disease revealed considerable overlap between DLB and PDD, suggesting that the timing of the onset of dementia does not relate to the quantity or distribution of cortical LB (Harding and Halliday, 2001). Surprisingly, there have been very few studies of the relationship between the early presence of other differentiating clinical features of DLB (visual hallucinations and/or fluctuating cognition) and the amount of cortical LB. In the absence of a dominant dementia syndrome at presentation, these additional features reliably predict the clinicopathological phenotype of DLB. However, visual hallucinations are also found in approximately 25% of patients with Parkinson’s disease (Aarsland et al., 1999a, b; Fénelon et al., 2000). It may be that the distribution and amount of cortical LB pathology relate more to additional and overlapping clinical features in such patient groups.

The present study analyses a large cohort of cases with LB pathology. Homogeneous clinicopathological groups of patients with DLB, PDD and Parkinson’s disease were segregated and the predictive value of visual hallucinations and/or fluctuating cognition assessed. Correlations between the distribution of cortical LB and these clinical features were also performed. The literature would predict a close association between these variables. Recent analyses of specific hallucination categories identify three psychosyndromes that reflect activation of specific visual pathways (Santhouse et al., 2000). In particular, the ventral temporal lobe is specialized for complex features relating to people, objects and landscapes (Tanaka, 1996; Halgren et al., 1999; Santhouse et al., 2000)—features characteristic of the well-formed visual hallucinations in DLB and some Parkinson’s disease patients. It is well known that LB concentrate in the ventral temporal lobe in DLB, particularly in the amygdala (Mattila et al., 1998, 1999, 2000; Hamilton, 2000), an area of the ventral temporal lobe activated by emotional facial interactions (Morris et al., 1996). Additionally, our recent analysis of cortical LB densities in DLB, PDD and Parkinson’s disease highlights the importance of parahippocampal LB in distinguishing cases with dementia from those with clinical Parkinson’s disease (Harding and Halliday, 2001).

In this study, we assess whether this distribution of cortical LB reflects the presence of these additional core features of DLB in homogeneous groups of cases followed to autopsy.

Methods

Brains were obtained over a 6-year period from participants in a number of clinical population-based studies of dementia (Broe et al., 1990; Reid et al., 1996a; Waite et al., 1996) and parkinsonian syndromes (Hely et al., 1999; Reid et al., 1996b) in the Sydney area. For all participants, regular annual clinical assessments using protocols devised from internationally standardized assessment batteries were available for analysis. Cases with ocular abnormalities were excluded. Immediately following death, information on the presence and severity of all clinical features was verified using standardized questionnaires completed by the subject’s specialist and general medical practitioners and a relative/carer/friend. Patients who significantly declined between their last assessment and death were excluded. Procedures for all studies (including consent) have been approved by the ethics review committees of the Central and South Eastern Sydney Areas Health Services and the Universities of Sydney and New South Wales.

For each case, the brain was removed, weighed and fixed by immersion in 15% buffered formalin for 2 weeks. The brain was routinely prepared for neuropathological analysis using current diagnostic criteria (Braak and Braak, 1991; Mirra et al., 1991; McKeith et al., 1996; National Institute on Aging and Reagan Institute Working Group on Diagnostic Criteria for the Neuropathological Assessment of Alzheimer’s Disease, 1997; Gelb et al., 1999; Harding and Halliday, 1998, 2001). Cases with prior head injury, cerebral infarction, frontotemporal dementia or progressive supranuclear palsy were excluded. Only cases with brainstem LB (McKeith et al., 1996; Harding and Halliday, 1998, 2001) were included (98 of 185 participants coming to autopsy). Of the 98 LB cases, 32 had additional neuropathology excluding them from further analysis. Cases were then assessed clinicopathologically using our new screening algorithm to separate cases with Parkinson’s disease only from those with dementia (Harding and Halliday, 2001). Of the cases with a maximum parahippocampal LB density of >1/field (n = 42), three cases were not demented (all with maximum of two LB/field). These cases were excluded in order to define homogeneous clinicopathological groups of both non-demented and demented patients. Sixty-three cases satisfied all criteria and were included in the study.
Quantitation of cerebral pathology
Five cortical regions [frontal (Brodmann area 9), temporal (area 20), parietal (area 39), occipital (areas 17 and 18) and anterior cingulate (area 24) cortices], as well as the hippocampus and parahippocampus at the level of the lateral geniculate nucleus and the amygdala at its greatest cross-sectional area were assessed quantitatively. Standard application of the CERAD (Consortium to Establish a Registry for Alzheimer’s Disease) method was initially used to determine the semi-quantitative levels of neuritic plaque within the cortex of each case (Mirra et al., 1991).

In each region of each case, the assessment of neurofibrillary tangles in adjacent sections stained with antibodies to tau enabled the semi-quantitative assessment of the distribution of tangles for Braak staging of Alzheimer’s disease (Braak and Braak, 1991). Additionally, this ensured no globule tangles were inadvertently included as LB in the analyses. Alzheimer’s disease (National Institute on Aging and Reagan Institute Working Group on Diagnostic Criteria for the Neuropathological Assessment of Alzheimer’s Disease, 1997) was a permissible pathological diagnosis, although cases with Alzheimer’s disease but without brainstem LB were excluded.

In each region LB were identified in immunohistochemically stained sections using antibodies to ubiquitin or α-synuclein and formic acid pre-treatment (Takeda et al., 1998). Sites of greatest LB density were determined at ×100 magnification; the magnification was then increased to quantify the number of LB/×200 field (field diameter of 1 mm). Paired Student’s t-tests revealed no significant differences in the number of LB identified in sequential sections using either ubiquitin or α-synuclein antibodies (Fig. 1A and B; P > 0.14). Counts from sections containing the highest densities of LB were used.

Evaluation of clinical signs and symptoms
For each case, assessments were performed routinely throughout the disease course, including assessments for the presence or absence of spasticity, brisk reflexes, ankle clonus and/or extrapyramidal features (tremor, bradykinesia, cogwheeling, stooped posture and glabellar tap). The clinical response and dosage of levodopa therapy were documented. The clinical severity of parkinsonism was assessed using the Hoehn and Yahr score (Hoehn and Yahr, 1967). Cases were diagnosed with dementia using the NINCDS–ADRDA (National Institute for Neurological and Communicative Disorders and Stroke and the Alzheimer’s Disease and Related Disorders Association) criteria for probable or possible Alzheimer’s disease (McKhann et al., 1984). Severity and progression of dementia were assessed according to the Mini–Mental State Examination and the Clinical Dementia Rating Scale (Morris, 1993). Functional impairment was rated by the Instrumental Activities of Daily Living (Lawton and Brody, 1969). The Neuropsychiatric Inventory (Cummings et al., 1994) was used to record psychiatric features, including depressive symptoms, hallucinations, disinhibition and apathy. Hallucinations are an under-reported phenomenon (Ballard et al., 1997), and therefore careful repeated enquiries were made regarding this clinical feature. The average interval ± standard deviation from last examination to death was 10 ± 7 months—substantially shorter than in previous analyses (Gómez-Tortosa et al., 1999). The last assessments prior to death (verified or modified by the information received after death from the specialist, general medical practitioners and carers) were used in all correlations. Given all the clinical information available for each case, the present study concentrated on analysing the core clinical features of DLB (McKeith et al., 1996, 1999), including bradykinesia and rigidity, fluctuations or clouding of consciousness, well-formed recurrent visual hallucinations and dementia. If these features were present, the date first noted was included in the analysis.

Visual hallucinations were noted when reported in the presence of the examiner on at least one occasion and/or when there were consistent reports by the subject to a carer. Flashing lights and similar phenomena reported to occur on hallucinogenic drugs (Fénelon et al., 2000) were not considered well-formed visual hallucinations. The types of visual hallucinations considered were mostly complex features relating to people, objects and landscapes. Because there has been some discussion on the difficulty with the reliability of fluctuations in cognition (Ince et al., 1998; McKeith et al., 1999), transient reductions or loss of consciousness were recorded as fluctuations in cognition. These resembled transient ischaemic attacks or delirium. Because of poor reproducibility and reliability, episodic periods of confusion were not considered sufficient to be recorded as true fluctuations in cognition.

Case classification and statistics
Cases were classified according to their clinical presentations. Four cases had no evidence of parkinsonism throughout their course (DLB-PD group in Table 1). Because of their small number, this group was not evaluated in the statistical analyses. All remaining cases (n = 59) had a clinical history of parkinsonism, and were grouped into those without dementia (PD-only group in Table 1), those with early dementia (DLB group, Table 1) and those with late dementia (PDD group, Table 1). All Parkinson’s disease-only and PDD subjects had good clinical improvement with high-dose levodopa therapy as required for a diagnosis of Parkinson’s disease (Gelb et al., 1999). The majority (14 out of 25) of cases in the DLB group were also on dopamine replacement therapy at their last assessment.

Most statistics were generated using the Statview programme (Abacus Concepts, Berkeley, Calif., USA) and a P value of 0.05 was taken as the level of significance. Differences between clinical groups were analysed using ANOVA (analysis of variance) and post hoc protected t-tests.
Positive and negative predictive values were calculated to ascertain whether other core clinical feature(s) predicted the clinicopathological group(s). Positive predictive values (PPVs) refer to the probability of having the clinical feature(s) for a particular clinicopathological diagnosis, while negative predictive values (NPVs) refer to the probability of not having the clinical feature(s) for a particular clinicopathological diagnosis. As parkinsonism was common to all
groups subjected to statistical assessment, no significant relationships were found for this clinical feature. Separate analyses were conducted using the presence or absence of early features (within 2 years of the first clinical symptom or sign). For pathological correlations, a separate series of ANOVA (feature early/first, late or never) were performed. A similar group analysis was performed for dementia severity correlates with cortical LB densities. Regression analyses were performed to relate the duration of any clinical feature/s to cortical LB densities.

Results

Clinical findings

After applying all exclusion criteria, 63 cases with a mean age of 73 ± 10 years were assessed (47 males, 16 females, age range 42–89 years). A total of 47 cases had a clinical diagnosis of dementia (75% of the sample), and 59 cases (94%) had a clinical diagnosis of parkinsonism. Overall disease duration was significantly shorter for cases with clinical DLB than for those with clinical Parkinson’s disease (Table 1; ANOVA = 11.7, P < 0.0001, protected t-test P values for DLB versus PDD < 0.0001, DLB versus Parkinson’s disease-only = 0.007, PDD versus Parkinson’s disease-only = 0.10). As expected, the average clinical dementia rating (CDR) score was significantly different for demented versus non-demented cases (Table 1; ANOVA = 48.5, P < 0.0001; protected t-test P values for Parkinson’s disease-only versus clinical DLB, P < 0.0001; Parkinson’s disease-only versus PDD, P < 0.0001; clinical DLB versus PDD, P = 0.00003). Cases with DLB had a significantly longer duration of dementia than cases with PDD (Table 1; t = 3.02, P = 0.004). There was no significant difference in the average Hoehn and Yahr score prior to death for any group (Table 1; ANOVA = 0.52, P = 0.60).

Only four cases had no clinical evidence of parkinsonism (DLB-PD group in Table 1). One case had sufficient plaques and tangles to reach NIA–Reagan (National Institute on Aging and Reagan Institute Working Group on Diagnostic Criteria for the Neuropathological Assessment of Alzheimer’s Disease) criteria for the diagnosis of Alzheimer’s disease. The remaining cases in this group all had short disease durations without any evidence of parkinsonism, even though they had significant midbrain cell loss and LB (required for study entry). All remaining cases (n = 59) had a clinical history of parkinsonism. Fifty-eight per cent of dementia cases were classified as DLB (Table 1), 19 with probable DLB and six with possible DLB (McKeith et al., 1996). In terms of dementia severity, six DLB cases had only a mild dementia, six cases had moderate dementia and 13 cases had severe dementia (52% of DLB group; Table 1). In contrast, the majority of PDD cases had only a mild dementia (67%; Table 1). 72% of DLB cases (n = 18) and 89% of PDD cases (n = 16) had early parkinsonism, severe in 52% and 67%, respectively (Hoehn and Yahr scores of 4–5; Table 1).

Fig. 1 Photomicrographs (A and B) and graphs of Lewy body density data (C–H). Mean ± Standard error of the mean. * indicates group(s) significantly different from Parkinson’s disease without dementia (PD-only) group. (A and B) Photomicrographs showing the typical appearance of cortical Lewy bodies (LB) visualized in serial sections of the parahippocampal gyrus of a demented case using (α-synuclein (A) or ubiquitin (B) immunohistochemistry following formic acid pre-treatment. A similar pattern of LB pathology is seen (equal numbers of Lewy bodies were identified using either antibody, P > 0.3). Scale in B is equivalent to that for A. (C and D) Bar graphs of the total LB burden (C) and the individual regions with high LB densities (D) for the three clinicopathological groups, dementia with Lewy bodies (DLB), Parkinson’s disease with later-onset dementia (PDD) and Parkinson’s disease cases without dementia (PD-only). The total LB burden is determined by adding the maximum LB density per microscopic field for each of the anterior cingulate cortex, frontal association cortex, parahippocampal cortex, inferior temporal cortex and amygdala. LB densities were greatest in the amygdala. Both the DLB and PDD groups have greater LB densities than the PD-only group, although the PDD group was only significantly greater than the PD-only group in the parahippocampal and amygdala. The DLB group differed from the PDD group in the total sum of LB and in the LB density in the inferior temporal cortex. ANOVA (all regions, C) = 13.7, P < 0.0001; post hoc tests—DLB versus PDD, P = 0.04; DLB versus PD-only, P = 0.0001; PDD versus PD-only, P = 0.04. ANOVA (cingulate, D) = 4.6, P = 0.01; post hoc tests—DLB versus PDD, P = 0.20; DLB versus PD-only, P = 0.004; PDD versus PD-only, P = 0.10. ANOVA (parahippo, D) = 10.3, P = 0.0002; post hoc tests—DLB versus PDD, P = 0.06; DLB versus PD-only, P = 0.0001; PDD versus PD-only, P = 0.02. ANOVA (amygdala, D) = 7.4, P = 0.0014; post hoc tests—DLB versus PDD, P = 0.49; DLB versus PD-only, P = 0.0004; PDD versus PD-only, P = 0.006. ANOVA (temporal, D) = 9.6, P = 0.0003; post hoc tests—DLB versus PDD, P = 0.017; DLB versus PD-only, P < 0.0001; PDD versus PD-only, P = 0.08. (E and F) Bar graph comparing the total LB burden (E) and the individual regions with high LB densities (F) for cases who had hallucinations compared with those without them. Cases with hallucinations had greater LB densities in the parahippocampal cortex (F, t = 5.1, P = 0.03), and amygdala (F, t = 4.2, P = 0.05), but no significant differences were found for the overall LB burden (E, t = 2.8, P = 0.10), or for the other regions examined (F, anterior cingulate cortex (t = 0.1, P = 0.7), inferior temporal cortex (t = 1.4, P = 0.25), frontal association cortex (t = 0.2, P = 0.9)). (G and H) Bar graphs comparing the total LB burden (G) and the individual regions with high LB densities (H) for the cases who experienced hallucinations first or early in their disease (early), for those with hallucinations only later in their disease process (late), or those who never experienced hallucinations during their disease course (never). Only cases with early hallucinations had a greater LB burden compared with non-hallucinators (G). These cases had higher LB densities in the parahippocampal and inferior temporal cortices (H). ANOVA (all regions, G) = 3.9, P = 0.03; post hoc tests—early hallucinations versus late hallucinations, P = 0.03; early hallucinations versus no hallucinations, P = 0.007; late hallucinations versus no hallucinations, P = 0.5. ANOVA (cingulate, H) = 2.6, P = 0.08. ANOVA (parahippo, H) = 5.2, P = 0.009; post hoc tests—early hallucinations versus late hallucinations, P = 0.03; early hallucinations versus no hallucinations, P = 0.002; late hallucinations versus no hallucinations, P = 0.2. ANOVA (amygdala, H) = 2.8, P = 0.07. ANOVA (temporal, H) = 3.8, P = 0.03; post hoc tests—early hallucinations versus late hallucinations, P = 0.02; early hallucinations versus no hallucinations, P = 0.01; late hallucinations versus no hallucinations, P = 0.9.
Eleven cases had parkinsonism as their only feature (69% of Parkinson’s disease-only group). As most cases with parkinsonism were at end-stage, parkinsonism was severe in most (Table 1). Fifty cases (85%) had parkinsonism early in the disease course. Of the two remaining DLB core symptoms, recurrent visual hallucinations were present in 35 cases (31 with dementia), an early symptom in 11 cases (31% of hallucinating cases, all with dementia). There were four Parkinson’s disease-only cases that had visual hallucinations during their disease—although this was a feature only late in the disease course. Two of these four cases also had fluctuating cognition, including one case where fluctuating cognition was an early sign. Fluctuating cognition was recorded in 25 cases overall (22 with dementia) and was an early feature in 13 (52% of cases with this feature; Table 1).

**Frequency of core DLB features in the clinicopathological groups**

As dementia and parkinsonism were used for clinical grouping, these features will not be considered further. The presence of the other core clinical features of DLB (i.e. fluctuating cognition and visual hallucinations) were examined within each group to determine diagnostic profiles. The majority of DLB (56%) and PDD (78%) cases had visual hallucinations, an early sign in 43% and 21%, respectively. In contrast, only 25% of the Parkinson’s disease-only group had hallucinations, and none was early. The majority of DLB cases had cognitive fluctuations (52%), an early sign in 69%. In contrast, 39% of PDD and 19% of Parkinson’s disease-only cases had recorded cognitive fluctuations, an early sign in only a minority (Table 1).

To analyse whether visual hallucinations and/or cognitive fluctuations were better predictors for either DLB or PDD, positive and negative predictive values were calculated using the proportion of cases with or without these features in each dementia group. Of these two symptoms, cognitive fluctuations had the best overall predictive power for DLB (PPV = 52%, NPV = 61% compared with PPV = 56%, NPV = 22% for hallucinations). The combination of these two features did not improve this predictive power (PPV = 32%, NPV = 67%). When the onset of the clinical feature was early, NPV improved for PDD (89% for fluctuations, 83% for hallucinations, 94% for both features occurring early) at the expense of PPV for DLB (36% for fluctuations, 24% for hallucinations, 4% for both features occurring early). These data show that, although neither of these features are good predictors of DLB in pathologically proven patients, the absence of these features early in the disease course is highly suggestive of PDD.

**Distribution and densities of cerebral pathology**

Across all groups, only six DLB cases had rare parietal lobe LB and fewer had LB in occipital cortex. LB in these regions were not considered further. Frontal association cortex had relatively low LB densities (2.1 ± 0.5 LB/field) with only 30% of cases having a maximum density of two or more LB/field. The greatest density distribution was in the frontal operculum.


**Hallucinations in Lewy body disease**

**Relationship between clinical features and cerebral Lewy bodies**

**Dementia**

The sum of the combined maximum LB densities across all regions did not differentiate cases with mild dementia from those with more severe dementia ($P > 0.3$), although cases with DLB had more LB overall compared with both Parkinson’s disease groups (Fig. 1C). There was no correlation between overall cortical LB burden and the duration of dementia ($P = 0.96$). As described above, DLB cases had significantly higher LB densities in the inferior temporal cortex (Fig. 1D), although there was no correlation between LB density and the duration of dementia ($P = 0.2$). This shows that high LB densities in the inferior temporal neocortex relate to the presence of early dementia and a diagnosis of DLB. In the PDD group, cases with moderate to severe dementia had higher densities of LB in the frontal cortex than those with mild dementia ($t = 5.6$, $P = 0.04$), as previously shown (Samuel et al., 1996; Mattila et al., 1998, 2000). However, this relationship did not hold true for cases with DLB ($t = 0.9$, $P = 0.3$) and was not related to the duration of dementia ($P = 0.7$).

**Hallucinations**

The majority of cases with hallucinations (28 out of 32) had dementia (either DLB or PDD). Cases with hallucinations had significantly more LB/field on average than those without hallucinations in the parahippocampus ($P = 0.028$) and amygdala ($P = 0.046$), but not in the frontal ($P = 0.88$), anterior cingulate ($P = 0.74$) or inferior temporal ($P = 0.25$) cortices (Fig. 1F). The overall cortical LB burden was significantly greater in those cases hallucinating initially or within the first years of disease onset regardless of diagnostic group (Fig. 1E and G; $P = 0.03$). This association was due to significantly more LB within the temporal cortices ($P = 0.009$ for parahippocampus, $P = 0.03$ for inferior temporal cortex). This indicates that LB pathology in medial temporal regions predisposes to hallucinations with the increasing involvement of more inferolateral temporal cortices ensuring the presence of hallucinations in cases with LB.

**Fluctuating cognition**

No significant statistical relationships were found between the summed or regional LB densities and fluctuations in cognition.

**Discussion**

The present study compares the clinical and pathological features of patients with parkinsonism and either early (DLB) or late (PDD) dementia. Our recent work revealed marked overlap between these clinical groups in the concentration and distribution of cortical LB (Harding and Halliday, 2001).

densities of LB were found in the amygdala ($6.8 \pm 0.6$ per field), anterior cingulate ($4.4 \pm 0.5$ per field), inferior temporal ($3.9 \pm 0.6$ per field) and parahippocampal ($3.8 \pm 0.6$ per field) regions. Over 60% of cases had two or more LB per field in these cortical regions, with approximately half of these cases having seven or more LB per field. With the exception of the amygdala, where high densities of LB were present in the vast majority of cases, high densities of LB in one region usually predicted high densities in other regions (multiple regression analyses, $P < 0.001$, $r^2$ range = 0.64 to 0.81). These densities and distributions are similar to those reported by Mattila et al. (1998).

Comparing between the groups, Parkinson’s disease-only cases were screened to exclude cases with significant cortical LB (Harding and Halliday, 2001) and all demented cases had significant densities of cortical LB. As described for both DLB and PDD (Samuel et al., 1996; Haroutunian et al., 2000; Mattila et al., 2000), the combined sum of the maximum LB density across all regions examined was significantly greater in cases with DLB compared with PDD, and significantly greater in PDD compared with the Parkinson’s disease-only cases (Fig. 1C; $P < 0.0001$). However, further analysis of the contribution of each individual region to this result shows that there were significantly higher LB densities only in the inferior temporal cortex of DLB cases compared with those with PDD (Fig. 1D; $P = 0.017$). Surprisingly, LB densities in all other regions examined were not significantly different between the dementia groups (Fig. 1D).

Twenty-seven cases (43%) had neuritic plaque densities sufficient for a CERAD diagnosis of probable or definite Alzheimer’s disease. These criteria for Alzheimer’s disease were reached by 72% of DLB cases and 33% of PDD cases. The cases with dementia and CERAD Alzheimer’s disease had more cortical LB only in the parahippocampus (LB/field) compared with the demented cases without Alzheimer’s disease ($6.4 \pm 5.1$ LB/field versus $2.8 \pm 2.2$ LB/field, $P = 0.011$). Six cases had significant numbers of neurofibrillary tangles (Braak stages 4–6, all demented). One DLB case had significant tangles but rare plaques, whereas the remaining five cases reached NIA–Reagan criteria for the diagnosis of Alzheimer’s disease (National Institute on Aging and Reagan Institute Working Group on Diagnostic Criteria for the Neuropathological Assessment of Alzheimer’s Disease, 1997). LB concentrations within the amygdala (12 LB/field) were significantly higher in these cases with Braak stages 4–6 ($P = 0.0002$), even though the overall LB burden did not differ ($P = 0.96$). Together these data show that cases with DLB are more likely to have plaque pathology and higher temporal lobe LB compared with cases with PDD. Neurofibrillary tangles are rare in both groups using the exclusion criteria outlined. However, those cases with tangles that achieve all criteria for Alzheimer’s disease (likely to contribute to or cause the dementia itself) concentrate LB in the amygdala.
This prompted us to question the correlation between cortical LB and dementia. In order to address this issue, the analysis of any associations between other core clinical features and the degree of LB pathology is required. A direct comparison of the presence or absence of well-formed visual hallucinations between patient groups revealed that significant numbers of patients with either DLB or PDD had visual hallucinations. There is some suggestion in the literature that there is some suggestion in the literature that fluctuations in cognition give greater accuracy of diagnosis for DLB (McKeith et al., 2000a; Walker et al., 2000). While our results suggest that this feature concentrates in DLB, particularly as an early feature, our statistics show that this feature is also a poor isolated predictor in patients with parkinsonism.

**Histopathological differences**

Apart from our recent work (Harding and Halliday, 2001), few studies have compared the neuropathology of patients with DLB versus those with PDD. For such a comparison, we standardized our methods for analysing LB, plaques and tangles comparing the maximal densities for all lesions—as now widely adopted (Mirra et al., 1991; Harding and Halliday, 1998; Newell et al., 1999). Our data show that cases with DLB are more likely to have cortical plaque densities similar to those seen in Alzheimer’s disease. This association is well-known (Hansen et al., 1993; McKenzie et al., 1996; Armstrong et al., 1997, 1998; Samuel et al., 1997; Brown et al., 1998; Mattila et al., 1998, 2000; Haroutianian et al., 2000), although few studies have compared DLB with PDD cases. Our results suggest that the vast majority of cases with PDD do not have sufficient densities of cortical plaques to suggest Alzheimer’s disease. Our previous analysis showed that, in DLB patients, both LB and plaques independently contributed to the dementia process (Harding and Halliday, 2001). We speculated that the combinations of these pathologies may account for the rapidity of the disease process in DLB (Harding and Halliday, 2001).

The present study also shows that patients with DLB have higher densities of LB in temporal neocortex compared with cases with PDD. This suggests that abnormal protein deposition in temporal neocortex relates to onset of cognitive decline and dementia in DLB rather than the stage or duration of dementia. A recent study suggests otherwise, reporting that the only clinicopathological relationship in cases with DLB is an increase in global LB burden with the duration of dementia (Gomez-Tortosa et al., 1999). However, this study did not evaluate many of the temporal lobe structures analysed in the present study and included dementia cases who had parkinsonism for >2 years (PDD in the present study). Increasing pathology within the temporal neocortex relating to dementia onset may have been predicted based on multiple studies in cases with Alzheimer’s disease (e.g. Braak and Braak, 1991) where increasing temporal neocortical pathology relates to dementia onset, stage and disease duration (Gertz et al., 1996; Berg et al., 1998; Delacourte et al., 1999). Our results show that high densities of LB in the inferior temporal neocortex differentiate cases with DLB, and that from other dementia syndromes (Hirono et al., 1999).
cases of PDD have increasing frontal LB densities with increasing dementia severity. We have previously shown that increasing densities of frontal LB are associated with significant frontal lobe atrophy (Cordato et al., 2000), and others have shown increasing frontal LB burden relates to increasing cognitive impairment (Samuel et al., 1996; Mattila et al., 1998, 2000). This dichotomy of regional pathology suggests that the disease processes driving DLB and PDD differ.

In the present study, neurofibrillary tangles were rarely seen in either DLB or PDD. However, in those cases with sufficient neuritic pathology to reach all criteria for Alzheimer’s disease (likely to contribute to or cause the dementia itself), LB concentrated in the amygdala. It has been noted that LB occur in the amygdala in the majority of cases with diverse neurodegenerative disorders, including familial and sporadic Alzheimer’s disease (Lippa et al., 1998; Hamilton, 2000; Marui et al., 2000; Mukaetova-Ladinska et al., 2000), Down’s syndrome (Lippa et al., 1999), Hallervorden–Spatz syndrome (Saito et al., 2000) and the parkinsonian–dementia complex of Guam (Yamazaki et al., 2000). These diseases have tau deposition in the amygdala in common with LB formation, suggesting that the formation of neurofibrillary tangles appears to precipitate LB within the amygdala independently of Parkinson’s disease and/or DLB.

**Clinicopathological correlates**

We found no associations between cortical LB pathology and fluctuations in cognition, consistent with previous reports (Gómez-Tortosa et al., 1999). In the present study, transient reductions or loss of consciousness were required for this feature to be positively identified. Reliable identification of this clinical feature is an acknowledged problem (McKeith et al., 1999). Previous studies describe excessive daytime drowsiness and other milder features as consistent with cognitive fluctuations (McKeith et al., 1996), and we may have chosen an arbitrarily severe cut-off for case inclusion. The prospective use of operational criteria for less severe fluctuations in cognition (Walker et al., 2000) may reveal relationships not identified in the present study. Additionally, the measurement of subcortical pathology may relate better to this clinical feature than the cortical pathology we analysed.

High LB densities in the amygdala and parahippocampal cortex were associated with visual hallucinations. Furthermore, increasing numbers of temporal lobe LB were associated with the earlier onset of this clinical feature. Although not significant, higher average LB densities have been previously observed in paralimbic cortices in patients with hallucinations compared with those without (Gómez-Tortosa et al., 1999). The larger numbers of cases analysed in the present study (35 out of 63 compared with 12 out of 25) are likely to have contributed to identifying this relationship as the numerical densities of cortical LB are relatively low. It should be noted that four of the 35 cases with recurrent visual hallucinations did not have dementia, but still had higher densities of temporal lobe LB. This frequency of visual hallucinations in patients with Parkinson’s disease is consistent with recent reports (Aarsland et al., 1999a, b; Fénelon et al., 2000). The amygdala and parahippocampus are regions commonly affected by pathology in dementing disorders, being the initiating sites for the pathology of Alzheimer’s disease (Braak and Braak, 1991), which possibly precipitates amygdala LB formation (Lippa et al., 1998; Hamilton, 2000; Marui et al., 2000; Mukaetova-Ladinska et al., 2000). Prevalently about 30% of cases with Alzheimer’s disease were thought to experience similar well-formed visual hallucinations (Ballard et al., 1997). However, recent studies suggest significantly fewer cases of Alzheimer’s disease have this clinical feature (<5%) compared with the majority of cases with DLB (Hirono et al., 1999). Parkinsonism and exaggerated cognitive decline are significant predictors of visual hallucinations in patients meeting criteria for probable Alzheimer’s disease (Paulsen et al., 2000), with prospective studies showing cortical LB in such cases (McShane et al., 1995). This suggests that a similar distribution of cortical LB in cases with Alzheimer’s disease may contribute to the clinical manifestation of visual hallucinations.

Few other neuropathological changes have been associated with well-formed visual hallucinations. Ocular pathology can precipitate visual hallucinations possibly via abnormal cortical release phenomena (Manford and Andermann, 1998; Santhouse et al., 2000). Brainstem vascular lesions in the midbrain reticular formation can also cause complex visual hallucinations (Manford and Andermann, 1998). Although speculative, this is possibly due to the involvement of ascending noradrenergic, cholinergic and serotonergic pathways which are also associated with sleep disturbances (Manford and Andermann, 1998; Arnulf et al., 2000; Turner et al., 2000). To our knowledge, we have described the first consistent association between the presence of visual hallucinations and a structural pathology in the cortex. As isolated lesions of the temporal lobe do not cause such visual hallucinations, the participation of other brain pathways is highly likely. In this regard, greater reductions in choline acetyltransferase activity in the temporal lobe have been associated with a higher prevalence of hallucinations in Parkinson’s disease, PDD and DLB (Perry et al., 1991; Tiraboschi et al., 2000). Thus, visual hallucinations are associated with two biological changes in the temporal cortex in LB disease—LB formation and decreased choline acetyltransferase activity. It will be important to determine the time course of the loss of presynaptic acetyl choline to reveal whether this pathology is a consequence of LB formation in the postsynaptic cortical neurones or a contributor to such intracellular pathology. This has important implications for long-term treatment benefit with cholinesterase inhibitors currently reported in cases with DLB (McKeith et al., 2000b; Rojas-Fernandez, 2001).

The underlying brain activation patterns of people undergoing complex visual hallucinations have recently been described. In particular, patients with Charles Bonnet...
syndrome, who report similar visual hallucinations to patients with LB, activate the anterior temporal projection of the ventral visual pathway when hallucinating about landscapes, figures, and vehicles with appropriate emotional context (ffytche et al., 1998; fftyche and Howard, 1999; Santhouse et al., 2000). These hallucinations are thought to occur because of a lack of occipital stimulation due to ocular pathology (ffytche et al., 1998; fftyche and Howard, 1999; Santhouse et al., 2000). Downstream ventral association cortices increase their activity as a result of cortical disinhibition with this abnormal activity in visual processing regions activating complex visual hallucinations. It would appear more than coincidental that the same brain regions concentrating LB in our patients with visual hallucinations were those activated by visual hallucinations in patients with the Charles Bonnet syndrome (Santhouse et al., 2000), particularly as decreased occipital glucose metabolism also occurs in patients with LB (Imamura et al., 1999, 2001; Lobotesis et al., 2001). The reduction in occipital metabolism occurs without underlying pathological inclusions, with a recent study suggesting that white matter abnormalities contribute to this metabolic deficit (Higuchi et al., 2000). In addition, DLB patients have preserved glucose metabolism in ventral temporal lobe regions, particularly those cases with visual hallucinations (Imamura et al., 1999; Higuchi et al., 2000; Lobotesis et al., 2001). These data suggest that activity in ventral visual pathways in association with underactivity of the primary visual cortices contributes to the well-formed visual hallucinations reported in cases with LB.

While it is generally believed that intracytoplasmic inclusions are a sign of neurodegeneration, several studies quantifying cortical neuronal loss in patients with DLB have shown a remarkable neuronal preservation (Gómez-Isla et al., 1999; Broe et al., 2001), suggesting that cortical LB may not signify neurodegeneration. If these cortical inclusions do not disrupt neuronal metabolism sufficiently to cause degeneration, they may disrupt cell mechanisms sufficiently to cause an increase in metabolic demand for neuronal survival. This may contribute to the association between the concentration of temporal lobe LB and well-formed visual hallucinations found in the present study. In fact, the overall pattern of temporal lobe LB formation appears to be more relevant to the onset of visual hallucinations than to the onset of dementia in cases with cortical LB disease.

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