Respiration and sleep in Parkinson's disease

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SUMMARY  Sleep and respiration during sleep were studied in patients with idiopathic Parkinson's disease, patients with Parkinsonism with autonomic disturbance, and normal age and sex matched controls. Patients with idiopathic Parkinson's disease showed significantly reduced REM sleep, and more frequent and prolonged waking throughout the night. Hypoventilation and sleep apnoea did not occur in the idiopathic Parkinson's disease or normal groups, but respiration was disorganised with frequent central and obstructive apnoeas in the autonomic disturbance group. Respiratory rate during non rapid eye movement sleep was similar in the idiopathic Parkinson's disease and normal groups, but patients with idiopathic Parkinson's disease showed tachypnoea awake and during REM sleep.

Abnormal sleep has been described in patients with Parkinson's disease. They sleep poorly, frequently waking, and have little stage III/IV or REM sleep.1-3 It has been proposed that these abnormalities are related to a reduction in brain amines,4 and treatment of Parkinson's disease with levodopa or bromocryptine may lead to an improvement in sleep quality.5-6 None of these studies has made any comment on respiration during sleep. McNicholas et al7 studied three patients with idiopathic Parkinson's disease, but reported the data on sleep for their normal subjects and Parkinsonian patients together, only providing respiratory data during sleep separately. Irregular respiration was noted to be a feature of patients with Parkinsonism following encephalitis lethargica, and was said to be worse during sleep.8 More recently there have been several case reports of hypoventilation in patients with post-encephalitic Parkinsonism,9-11 or associated with familial Parkinson's syndrome,12 13 but none of these reports described sleep patterns.

We have studied the patterns of sleep and respiration in patients with idiopathic Parkinson's disease, to document any evidence of hypoventilation or respiratory arrhythmias associated with this condition. This group has been compared with normal controls, and Parkinsonian patients with autonomic disturbance, one of whom had Shy Drager syndrome.

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Methods

We studied 12 patients (3 female, 9 male, mean age 57.9 years SD ± 8.99 years) with idiopathic Parkinson's disease, and compared them with 12 normal control subjects, matched for age and sex (mean age 56.8 years SD ± 8.2 years). In addition two patients with Parkinsonian features and autonomic dysfunction were studied. All of the patients were in a stable clinical state, and were on optimal therapy with anti-Parkinsonian drugs. None of the patients or controls had any history of respiratory or cardiac disease.

The subjects slept on a hospital bed in a quiet darkened room, with the investigator and recording equipment in an adjoining room. The Parkinsonian patients were studied for two nights, and the normal controls, who were all members of staff and accustomed to investigation, for one night. In the Parkinsonian patients the record from the second night of study was taken for analysis. Sleep was assessed by standard methods.14 The electroencephalogram (EEG) was recorded from position C4 A1, and the electro-oculogram was recorded from two electrodes placed obliquely, one above the outer aspect of the left eye, the other placed below the outer aspect of the right eye. The submental electro-myogram (EMG) was recorded from electrodes positioned on the right side of the neck. The electrocardiogram (ECG) was recorded from electrodes on the shoulders in lead I configuration. All of these recordings were obtained using 10 mm silver cup electrodes. Airflow was assessed at the nose and mouth by thermistors mounted on nasal cannulae, or with a laryngeal microphone. Respiratory movements were recorded with inductance coils (Respirtrace) placed around both chest and upper abdomen. Haemoglobin oxygen saturation was measured with a Hewlett Packard ear lobe oximeter attached to the ear throughout the night. All information was recorded onto paper on a 16 channel
Results

IDIOPATHIC PARKINSON'S DISEASE PATIENTS AND NORMAL CONTROLS

The 12 patients with idiopathic Parkinson's disease were compared with an equal number of age and sex matched normal controls. Details of age, sex, duration of disease, severity of disease, and medication for the patients is given in the table.

Sleep

Analysis of sleep revealed that the mean time taken for Parkinsonian patients to go to sleep was longer than that for the normals, but that this was not statistically significant (mean ± standard deviation. Normals 16 ± 10 min, patients with idiopathic Parkinsonism 22.5 ± 17 min). During the night there were more frequent wakes in the patients with idiopathic Parkinson's disease (normals 6-1 ± 2.0; patients 8.9 ± 3.1; p < 0.05), and more time was spent awake in this group (normals 81 ± 50 min.; patients 169 ± 77 min; p < 0.01). This was associated with a significantly shorter duration of REM sleep in the Parkinsonian patients (normals 43 ± 19 min; patients 19.5 ± 23 min; p < 0.05), six of whom showed evidence of REM sleep for less than 1 minute. There was no difference in the time spent in any of the stages of non REM sleep between the two groups. Total sleep time was shorter in the patient group than in the normal group, owing to a reduction in the duration of REM sleep in the patient group, but this did not reach statistical significance (fig 1). REM latency was similar in the two groups (normals 157 ± 93 min; patients 148 ± 50 min). On going to sleep the submental EMG decreased in both groups, and decreased further in both groups equally on going into REM sleep.
Haemoglobin oxygen saturation

Haemoglobin oxygen saturation fell in all subjects during sleep, to the same extent in both groups (fig 2). Haemoglobin oxygen saturation was similar when awake (normals 96·5 ± 1·3%; patients 96·3 ± 1·1%), in stage II sleep (normals 95·8 ± 1·8%; patients 95·4 ± 0·9%), and in REM sleep (normals 95·3 ± 1·9%; patients 95·5 ± 0·9%). Minimum saturation was similar in the two groups (normals 91·3 ± 4·8%; patients 95·5 ± 2·0%), and a similar proportion of time was spent with a haemoglobin oxygen saturation of <95%, (normals 16·5% of the night; patients 17% of the night), with very little time spent with a saturation of <90% (normals 3 min; patients 1 min).

Respiratory rate and rhythm

Respiratory rate was significantly faster in the patient group when awake, than in the normal controls (normals 13·9 ± 1·1 breaths/min; patients 17·9 ± 1·6: \( p < 0·006 \)), but on going to sleep the respiratory rate decreased in the patient group so that in stage II sleep the respiratory rate was no longer significantly faster than the normal group. (normals 14·0 ± 0·57 breaths/min; patients 15·0 ± 0·69: NS). During REM sleep the mean respiratory rate was greater in the patient group, with no significant change in the normal group (normals 14·6 ± 1·6/min; patients 17·0 ± 1·4/min: \( p < 0·05 \)) (see fig 3).

In only one subject was there any evidence of any significant number of apnoeic episodes. This was a 68-year-old normal control subject, who gave no history of snoring or disturbance of sleep. During the night he had 30 central apnoeas, and 10 obstructive apnoeic episodes, all occurring in Stage I NREM, or in REM sleep. None of the apnoeas were longer than 25 seconds, mean apnoea length was 20 seconds; there were no cardiac dysrhythmias, and although haemoglobin oxygen saturation fell during each apnoea, only 1 minute was spent during the whole night with a saturation of <90%. In five other normal subjects there were central apnoeas (1, 1, 2, 2, 1, apnoeas each), and in five of the patient group (1, 1, 2, 3, 4, apnoeas each), but none of these apnoeas lasted longer than 20 s, and all apnoeas occurred during stage I NREM, or in REM sleep.

PATIENTS WITH AUTONOMIC DYSFUNCTION

Two patients with autonomic disturbance and Parkinsonian features were studied.

**Patient 1** This 57-year-old lady had a 5 year history of Parkinsonism, severe postural hypotension, snoring and constipation. She showed severe lability of blood pressure, and had evidence of autonomic failure, with no change in pulse rate during a Valsalva manoeuvre. It was felt that this patient had orthostatic hypotension and idiopathic Parkinson's disease. There was no stridor when awake, and vocal cord movement was normal. The sleep study revealed a low (85%) haemoglobin oxygen saturation when awake, which fell further on going to sleep (mean asleep, 82%, minimum 80%). The respiratory rhythm whilst asleep was irregular both in rate and volume, and respiration was noisy. There were occasional episodes of obstructive apnoea, in which there was no airflow despite continuing respiratory movement, and paradoxical movement of the chest inwards during attempted inspiration. There was little sleep, but frequent wakings after episodes of apnoea.

**Patient 2** This 43-year-old man had first presented with urinary retention three years previously and been found to have a neuropathic bladder. Subsequently he developed a right extensor plantar response, pyramidal weakness of the right arm, minimal cerebellar signs, and mild extrapyramidal signs. Initially there was no postural hypotension, but this developed later, with an absent heart rate response to the Valsalva manoeuvre. This patient had many features of the Shy-Drager syndrome. Sleep study revealed grossly disturbed sleep, with frequent obs-
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tructive apnoeas leading to waking. The haemoglobin oxygen saturation was normal when awake, but fell on going to sleep, falling further during apnoeic episodes. During sleep the respiratory rate and rhythm were irregular. When awake there was evidence of upper airway obstruction and stridor, and indirect laryngoscopy revealed paralysis of abduction of the vocal cords.

Discussion

Sleep studies in patients with Parkinson's disease have shown a wide variety of abnormalities. An increase in sleep latency, frequency of waking during the night, and increased time spent awake, have all been described. The authors have reported reduced stage III/IV sleep, and a reduction in the duration of REM sleep. Other workers have reported normal stage III/IV sleep, or even completely normal sleep. These discrepancies have been explained in terms of there being two groups of patients with Parkinson's syndrome, one group with reduced REM sleep, and increased submental muscle activity during REM sleep, and the other with normal REM sleep, and repetitive blinking at the beginning of the night.

Few of these studies have attempted, however, to differentiate patients according to aetiology, age, or severity of disease. Friedman has shown for patients with Parkinson's disease that impairment of sleep may be directly related to the severity of disease. In addition, some of the abnormalities described in association with Parkinson's disease are present in a normal ageing population which makes the assessment of more elderly patients difficult.

We have studied a relatively young group of patients with idiopathic Parkinson's disease, and compared them with age and sex matched controls. Our results show frequent and prolonged waking in the Parkinsonian patients. There was a reduction in the duration of REM sleep, but no abnormalities of NREM sleep. These results are similar to those of Friedman, who found reduced stage III/IV sleep only in patients with severe disease (Hoehn and Yahr grade IV/V, and only one of our patients was in this group). We did not find either increased submental activity, or repetitive blinking in any of our patients, so it was not possible to analyse our data as did Mouret. In our patients there was no correlation between the severity of disease and limitation of REM sleep.

Several reports of an improvement with dopaminergic therapy, with prolongation of REM sleep duration, have suggested that dopamine deficiency is important in causing the sleep abnormalities seen in these patients. All of our patients were receiving either Sinemet, Madopar, levodopa, amantidine, or bromocryptine, which increase dopamine activity centrally. It is possible that the abnormalities of sleep seen in our patients would have been more marked had they not been receiving such therapy, and this may explain why our results show less abnormality of sleep than some of the earlier studies. It is unlikely that the drug therapy itself caused the abnormalities seen, as previous studies have demonstrated similar abnormalities prior to therapy, with improvement on therapy.

In contrast to the patients with idiopathic Parkinson's disease, who showed only mild abnormalities of sleep, the two patients with Parkinsonism and autonomic dysfunction showed grossly disturbed sleep. In neither patient was there much organised sleep; apnoeas and obstructive episodes caused frequent arousals. McNicholas et al and Guilleminault have reported similar findings in patients with the Shy-Drager syndrome, although Parkinsonism was not a major feature in all their patients. Most of the abnormalities of sleep appear to be related to waking in response to arousal after central or obstructive apnoeas, which are rare in other Parkinsonian patients. It is probable that these abnormalities are more related to the autonomic failure in these patients, and are not related to the Parkinsonism.

McNicholas et al have reported on respiration during sleep in three patients with idiopathic Parkinson's disease. They showed no evidence of sleep apnoea or desaturation during sleep, and noted a regular respiratory rhythm awake and asleep; they suggested that automatic control of respiration was normal in their patients with this condition. Our patients showed similar results, with no evidence of hypventilation or significant sleep apnoeas. All case reports of hypoventilation in Parkinsonian patients have appeared in patients with postencephalitic Parkinsonism, or familial Parkinsonism.

In Parkinsonism following encephalitis lethargica there is evidence of more widespread brainstem damage than in idiopathic Parkinson's disease. Even after encephalitis lethargica only a small proportion of patients showed respiratory complications. Perhaps only with the widespread brainstem damage present in a few cases of encephalitis lethargica is there failure of automatic control of respiration, which may then lead to the hypoventilation and respiratory dysrhythmias which are characteristic of such failure of respiratory control.

We observed a tachypnoea in our patients with idiopathic Parkinson’s disease, which decreased on
going to sleep. This tachypnoea might be related to central factors, either to the disease itself, or the effect of drugs, to the presence of respiratory disease, or to chest wall rigidity and stiffness as a result of the disease. Tachypnoea was a frequent finding during the encephalitic epidemic of the 1920s, and has been reported since in patients with Parkinson's syndrome prior to the introduction of levodopa therapy. Tachypnoea, breathlessness, and respiratory dysrhythmias have been reported following levodopa therapy. Similar irregularity of respiration in patients with tardive dyskinesia, in whom excess dopamine activity has been implicated, have been reported. All these observations suggest that both dopamine deficiency and dopamine excess may lead to tachypnoea and affect the respiratory rate and rhythm. In our patients there were no respiratory dysrhythmias or very fast tachypnoea characteristic of dopamine overactivity, but a slightly faster rate than normals, similar to that reported by Kim. It is possible that this abnormality could be caused by dopamine underactivity from the disease, as despite therapy, the patients still had signs of Parkinsonism. Since we did not study the effect of alteration of drug dosage, it is not possible to say that the abnormality was not related to drug therapy.

An alternative explanation for our results would be that the patients had evidence of respiratory disease, but his was denied by our patients, and all patients had a normal awake haemoglobin oxygen saturation, which suggests that there was no respiratory disease sufficient to cause hypoxia. A further possible explanation for our results might be that the tachypnoea was caused by increased stiffness of the chest wall. Awake patients with Parkinson's syndrome show increased intercostal muscle activity throughout the respiratory cycle, and this leads to increased stiffness of the chest wall. This might be expected to lead to increases in respiratory rate in compensation for the increased work of breathing.

On going to sleep patients with Parkinson's disease lose their rigidity and tremor, and in our patients this was observed. Similarly on going to sleep the respiratory rate fell in the Parkinsonian patients but not in the normal controls. This may be explained in terms of alterations in the central control of breathing, or may be due to relaxation of the chest wall muscles, resulting in reduced work for breathing, hence leading to a reduction in respiratory rate. Either of these explanations fits our results. During REM sleep there is characteristically even more hypotonia than in non REM sleep, but respiration is less automatic, and depends more upon voluntary control. In our normal subjects there was a small increase in the respiratory rate; in the Parkinsonian patients there was no evidence of a further fall in respiratory rate, as might be expected if peripheral chest wall muscle tone was the sole determinant of rate. The respiratory rate increased slightly in those patients who had enough REM sleep for analysis, suggesting that central factors were more important in determination of the respiratory rate than the chest wall tone.

These minor abnormalities of respiratory rate seen in the patients with idiopathic Parkinson's disease are in complete contrast to the changes seen in patients with evidence of autonomic dysfunction. Both of these patients showed frequent apnoeic episodes when asleep, with falls in haemoglobin oxygen saturation, and a disorganised respiratory rhythm both awake and sleeping. It has been suggested that the disordered respiratory pattern seen in these patients is associated with a defect in the automatic respiratory rhythm generator in the brain stem. In addition, the obstructive episodes seen in our patients, and in those cases previously reported, are due to a failure of abduction of the vocal cords, causing obstruction of the larynx during inspiration. This vocal cord palsy was also present in one of our two patients when awake. As none of these abnormalities was seen in any of our patients with idiopathic Parkinson's disease, it is probable that their occurrence is due to the disease which results in the central autonomic dysfunction in these patients and not the Parkinson's disease itself.

Patients with idiopathic Parkinson's disease show abnormal sleep with frequent waking and reduced REM sleep. They also show a tachypnoea when awake, that disappears when they go to sleep, and have a normal haemoglobin oxygen saturation both awake and asleep, with no evidence of sleep apnoea. Patients with autonomic dysfunction and Parkinsonism, in contrast, show grossly disturbed respiration, with hypoventilation and disorganised respiratory rhythm, and wake frequently, showing very little normal sleep. Both Parkinson's disease, and the drugs used in its treatment appear to affect both respiration and sleep. These respiratory effects may be due to either direct central effects on the respiratory rhythm generation, to effects on the respiratory rate due to changing chest wall tone, or to a combination of the two.

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