Psychosis is common in Parkinson’s disease (PD), particularly in its later stages. The symptoms range from comparatively minor illusions, vivid dreams, and occasional, non-disturbing visual hallucinations to frank psychosis. The pathogenesis of psychosis in PD is not fully known. Management of psychosis in PD requires a multidisciplinary approach. Some of the newer atypical antipsychotics are effective against psychosis with no significant worsening of PD. Psychosis in PD is associated with poor quality of life for patients and the carers.

Psychosis affects nearly one third of patients with Parkinson’s disease (PD). Its manifestations range from comparatively minor symptoms of mild illusions, vivid dreams, and occasional, non-disturbing visual hallucinations in a clear sensorium to a frank psychosis with disturbing visual (and rarely, auditory and tactile) hallucinations, paranoid delusions, and confusional psychosis. Although it was a recognised complication even before L-dopa was introduced, psychosis has been increasingly reported in patients treated with dopaminergic drugs. Psychosis in PD significantly worsen or even precipitate psychosis in PD. The pathogenesis of psychosis in PD is not fully known. Management of psychosis in PD requires a multidisciplinary approach. Some of the newer atypical antipsychotics are effective against psychosis with no significant worsening of PD. Psychosis in PD is associated with poor quality of life for patients and the carers.

Psychosis in PD generally occurs late in the course of presumed PD or when they occur shortly after the introduction of dopaminergic medicines, other diagnoses besides PD, for example, dementia with Lewy bodies or Alzheimer’s disease should be considered. As hallucinations become more vivid, insight may be lost and the patient may start acting upon hallucinations. Auditory and tactile hallucinations may follow causing an organic confusional psychosis. A simple scale to assess the severity of hallucinations (table 1) can be used. Paranoid delusions, often directed towards the spouse or other family members, may be very distressing and often precipitate institutionalisation.

PATHOPHYSIOLOGY

Despite advances in neurochemistry, the pathophysiological basis of psychosis in PD remains poorly understood. Both endogenous (for example, drugs) and endogenous (related to the disease process itself) factors contribute to the development of psychosis in PD. The interplay between various neurotransmitters including dopamine, acetylcholine, and serotonin (5-hydroxytryptamine, 5-HT) seems to be of importance. Dopamine has long been recognised as an important neurotransmitter in the disorders of perception. Dopamine agonists (for example, amphetamine) are known to produce psychosis and the commonly used antipsychotic agents are blockers of dopamine receptors. It is suggested that the dopaminergic overactivity in the pathways to limbic system and cerebral cortex is involved in the production of psychosis.

Degeneration of some of the pathways using 5-HT as a neurotransmitter may play an important part. The improvement of psychosis with ondansetron (a 5-HT3 antagonist) and atypical

emannine (REM) sleep disturbances, and vivid dreaming may precede hallucinations by weeks to months. Hallucinations are mostly visual and often non-threatening, at least initially. The visual hallucinations are mostly complex, usually containing animals, objects, or persons although simple hallucinations (characterised by the absence of form, for example, flashes of light or colour) also occur. They tend to occur in poor lighting conditions (“sundowning”) or on waking from sleep. Patients may “see” crawling bugs on the patterned carpets or moving insects on the furniture.

Hallucinations usually occur after patients have taken PD drugs for several months or years. When hallucinations occur early in the course of presumed PD or when they occur shortly after the introduction of dopaminergic medicines, other diagnoses besides PD, for example, dementia with Lewy bodies or Alzheimer’s disease should be considered. As hallucinations become more vivid, insight may be lost and the patient may start acting upon hallucinations. Auditory and tactile hallucinations may follow causing an organic confusional psychosis. A simple scale to assess the severity of hallucinations (table 1) can be used. Paranoid delusions, often directed towards the spouse or other family members, may be very distressing and often precipitate institutionalisation.

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Degeneration of some of the pathways using 5-HT as a neurotransmitter may play an important part. The improvement of psychosis with ondansetron (a 5-HT3 antagonist) and atypical
neuroleptics (with their blocking effects on serotonin receptors and dopamine receptors) lends some support to this concept.

Acetylcholine is another neurotransmitter with a potential role in psychosis of PD.

Pathological studies have shown significant degeneration of cholinergic neurons in PD. Goetz et al postulated that acetylcholine block could induce PD related hallucinations.

As in many organic conditions associated with visual hallucinations, abnormalities of visual processing have also been implicated in the generation of hallucinations in PD. Parkinsonian patients with hallucinations had difficulty distinguishing between images they saw and images created in their mind. It has been shown that PD patients without hallucinations process visual information in the occipital, temporal and parietal areas, whereas those with hallucinations do it in the frontal cortex.

Sleep disturbances commonly precede psychosis in PD. Acetylcholine is an important neurotransmitter in the pathways controlling REM sleep. Some authors have suggested a link between hallucinations and perturbations of REM sleep.

Two different mechanisms have been proposed for the two subgroups susceptible to hallucinosis in PD. In patients with disease duration of five years or less, postsynaptic dopamine receptor denervation supersensitivity in the mesolimbic/mesocortical system may underlie hallucinosis. In patients with disease duration greater than five years, hallucinosis may be mediated either by changes outside the basal ganglia or by the serotoninergic system.

It is probable that the psychosis in PD is multifactorial and is caused by abnormalities in several neurochemical transmitters and neural structures.

**MANAGEMENT**

Psychosis in PD poses great management challenge because of potentially catastrophic consequences for the patient and the carers, lack of universally effective and safe drugs, and the often progressive nature of this complication.

**General principles of management**

- It is vital to exclude delirium in all patients who present with psychosis. This is important as the treatment of the precipitating illness can potentially “cure” psychosis. A thorough search should be undertaken to exclude an intercurrent physical illness, drug effect, electrolyte imbalance, and systemic infection.

- A multidisciplinary team approach, involving patient, carer, physician, liaison psychiatrist, PD specialist nurse, and social worker is required for the effective management. Proper explanation of the nature and course of psychosis to the patient and carer helps them to cope with the situation.

- Simple measures should be tried first, for example, improving sleep hygiene, avoiding excessively patterned furniture, and reducing sensory overload and sensory deprivation.

- Psychiatric comorbidities of non-psychotic nature, for example, depression, anxiety, and apathy are common in PD patients with psychosis and may add to the functional impairment. Treatment of these conditions may reduce morbidity and distress to carers.

- Vivid dreams alone usually do not warrant medical treatment. Similarly, occasional visual hallucinations with insight retained may not require any action beyond reassurance.

- If simple measures fail, reduction in drug dose or elimination of one or more antiparkinsonian agents should be considered. The agent introduced most recently or one with the lowest antiparkinson efficacy (anticholinergics, amantadine, selegiline) should be stopped first. Next in order are dopamine agonists, catecholamine-o-methyl transferase inhibitors, and L-dopa. It should be emphasised that dopamine agonists are much more likely to produce psychosis than L-dopa. One might have to accept worsening of parkinsonism.

**Drug treatment**

When the above measures fail, drug treatment becomes necessary. Drugs used to treat psychosis in PD include atypical antipsychotics (AA), 5-HT3 receptor antagonists, and acetylcholinesterase inhibitors (AChI).

(1) Atypical antipsychotics (AA): conventional antipsychotics (for example, haloperidol) cause intolerable worsening of motor function and should be avoided. Newer AA are associated with fewer extrapyramidal side effects because of their non-selective antagonism at both serotonergic and dopaminergic receptors. Examples of AA include clozapine, quetiapine, olanzapine, risperidone, and aripiprazole. The following factors should be carefully considered while using AA:

- With few exceptions, all AA have comparable efficacy against psychosis and the choice is mainly based on their ease of use and the side effect profile. Risperidone and olanzapine are associated with sedation and can cause considerable worsening of parkinsonism. Additionally, olanzapine can worsen cognition and hyperglycaemia in patients with diabetes. A recent CSM warning suggests an increased risk of strokes associated with the use of risperidone and olanzapine in old people. Clozapine is probably the most effective agent and does not cause worsening of parkinsonism. However, it is associated with the risk of potentially fatal, idiosyncratic agranulocytosis necessitating regular white blood cell counts. It has a restricted licence in the UK. Quetiapine is favoured by many psychiatrists because of its better side effect profile. Interestingly, a recent study showed quetiapine to be as effective as clozapine. Preliminary experience with the latest AA, aripiprazole, has not been very encouraging, as it caused worsening of PD symptoms.

- In older patients, AA can cause sedation and postural hypotension. Sedation can be helpful in patients whose behavioural problems occur at night when they are awake.

- AA should be started at very low doses with gradual increments if needed, as hypersensitivity to these agents can induce delirium or malignant neuroleptic syndrome. The doses required to treat psychosis are much lower than those used in schizophrenia.

- All AA cause prolongation of QT interval on ECG, particularly in older patients. Therefore, baseline ECG and periodic ECG monitoring are advisable.

- Optimal duration of treatment with AA in PD associated psychosis is not known. Because of the risk of serious side effects, long term use of AA should not be taken lightly.

**Table 1 Hallucination scale in Parkinson’s disease**

<table>
<thead>
<tr>
<th>Symptom</th>
<th>Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>0</td>
</tr>
<tr>
<td>Vivid dreams, illusions, sense of presence</td>
<td>1</td>
</tr>
<tr>
<td>Dreams encroaching on waking hours, occasional tolerable hallucinations</td>
<td>2</td>
</tr>
<tr>
<td>Regular evening and night time intrusive visual hallucinations</td>
<td>3</td>
</tr>
</tbody>
</table>
However, stopping AA in patients in remission is fraught with the possibility of recurrences and even ‘rebooms’ of psychosis. Fernandez et al. conducted a prospective study on PD patients receiving successful treatment with quetiapine and clozapine as these drugs were withdrawn. After the antipsychotic agent was stopped, psychosis recurred in five of six patients. In three patients the ‘rebound psychosis’ was worse than the original psychotic episode requiring higher antipsychotic drug doses.

- **AA treatment improves prognosis in patients with psychosis.** In a study of 59 patients with psychosis treated with clazapine, the death rate was lower at 28% within two years compared with 100% in a study done before the availability of this treatment. Of all AA, clozapine is the most well studied agent and is considered as the “gold standard” treatment in psychosis of PD. In a randomised, placebo controlled trial, clozapine at a mean dose lower than 50 mg/day improved drug induced psychosis in PD without significant worsening of motor function. The detailed account of individual AA in psychosis is beyond the scope of this article and interested readers are referred to a recent review.

(2) Acetylcholinesterase inhibitors (AChI): these agents increase concentrations of acetylcholine in the brain and have been in use for treating Alzheimer dementia. Their use in psychosis of PD has been advocated on the basis of reports of beneficial effects in dementia with Lewy bodies. There is a significant loss of cholinergic neurons in PD and AChI may prove beneficial by increasing acetylcholine concentrations. In an open label study, rivastigmine led to improvement in cognitive and functional abilities, as well as the resolution of behavioural problems and visual hallucinations. Donepezil, another commonly used AChI, has also shown some promise in open label studies.

(3) Ondansetron: it is a 5-HT3 antagonist widely used to treat vomiting attributable to anticancer agents. In an open label, short term (four to eight weeks) study, it was used to treat psychosis in PD at a dose of 12 to 24 mg daily. There was pronounced to moderate improvement in measures of visual hallucinations, paranoid delusions, confusion, and the associated global functional impairment. It did not cause worsening in PD symptoms and was well tolerated. However, there have not been any large sized, controlled studies using ondansetron in PD associated psychosis. Ondansetron is an expensive drug.

Electroconvulsive therapy (ECT): ECT has been used in treating drug resistant depression in PD. There have also been reports of its success in improving motor symptoms of PD. However, it has not been systematically tested in treating psychosis of PD. There have been a few reports of its benefit in drug refractory psychosis of PD. Because of its cost and associated stigma, ECT is unlikely to be a primary treatment, except possibly for the drug resistant or drug intolerant cases or when psychosis is associated with severe depression.

**CONCLUSIONS**

Psychosis commonly complicates PD, especially in its later stages. It is a strong predictor of institutionalisation. Several risk factors have been identified including, old age, sleep disturbances, long duration of PD, history of depression, cognitive impairment, and visual disorders. Pathophysiology of psychosis remains unknown, although dopamine, serotonin, and t acetylcholine are the major neurotransmitters of interest. Effective treatment of psychosis requires multidisciplinary team approach and attention to the intercurrent illnesses, optimising drug treatment of PD, and judicious use of AA. Further studies are needed to evaluate the use of 5-HT3 antagonists, acetylcholinesterase inhibitors, and ECT in psychosis of PD.

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