Drug-Induced Parkinsonism

Hae-Won Shin, a Sun Ju Chung b

aDepartment of Neurology, Chung-Ang University College of Medicine, Seoul, Korea
bParkinson/Alzheimer Center, Department of Neurology, University of Ulsan College of Medicine, Seoul, Korea

Drug-induced parkinsonism (DIP) is the second-most-common etiology of parkinsonism in the elderly after Parkinson’s disease (PD). Many patients with DIP may be misdiagnosed with PD because the clinical features of these two conditions are indistinguishable. Moreover, neurological deficits in patients with DIP may be severe enough to affect daily activities and may persist for long periods of time after the cessation of drug taking. In addition to typical antipsychotics, DIP may be caused by gastrointestinal prokinetics, calcium channel blockers, atypical antipsychotics, and antiepileptic drugs. The clinical manifestations of DIP are classically described as bilateral and symmetric parkinsonism without tremor at rest. However, about half of DIP patients show asymmetrical parkinsonism and tremor at rest, making it difficult to differentiate DIP from PD. The pathophysiology of DIP is related to drug-induced changes in the basal ganglia motor circuit secondary to dopaminergic receptor blockade. Since these effects are limited to postsynaptic dopaminergic receptors, it is expected that presynaptic dopaminergic neurons in the striatum will be intact. Dopamine transporter (DAT) imaging is useful for diagnosing presynaptic parkinsonism. DAT uptake in the striatum is significantly decreased even in the early stage of PD, and this characteristic may help in differentiating PD from DIP. DIP may have a significant and longstanding effect on patients’ daily lives, and so physicians should be cautious when prescribing dopaminergic receptor blockers and should monitor patients’ neurological signs, especially for parkinsonism and other movement disorders.

Key Words Parkinsonism, dopamine receptor blocking agents, clinical manifestations, dopamine transporter imaging, treatment.

Introduction

Drug-induced movement disorders include drug-induced parkinsonism (DIP), tardive dyskinesia (TD), tardive dystonia, akathisia, myoclonus, and tremor. Among these, DIP is the most common movement disorder induced by drugs that affect dopamine receptors. 1-3 Since the clinical manifestations of DIP are very similar to those of Parkinson’s disease (PD), patients with DIP are frequently misdiagnosed as having PD. 1-4 These patients are often prescribed antiparkinsonian drugs unnecessarily for long periods of time, despite recovery being possible simply by discontinuing the offending drugs. Dopamine transporter (DAT) imaging may be used in the differential diagnosis of various etiologies of parkinsonism, including DIP. 5-8 The aim of this review was to provide clinicians with updated information about the clinical characteristics and DAT imaging findings of patients with DIP, and about the correct treatment for DIP.

Epidemiology of Drug-Induced Parkinsonism

The exact prevalence and incidence of DIP are unclear because it is frequently unrecognized or misdiagnosed as PD. The first study of the extrapyramidal side effects (EPS) of the antipsychotic chlorpromazine found that about 40% of these patients exhibited parkinsonism, 7 and several subsequent epidemiological studies found that DIP is the second most common etiology of parkinsonism. 8-10 A community-based survey and a population-based study found DIP prevalence rates of 2.7% and 1.7%, respectively, whereas those of PD were 3.3% and 4.5%, respectively. 8,10 However, 6.8% of the patients diagnosed with PD were later reclassified as having DIP, thus emphasizing the
difficulties in accurately diagnosing DIP and in measuring its prevalence. Age is the most obvious risk factor for DIP, since dopamine concentrations decrease and nigral cells degenerate with age. Another risk factor is female gender, suggesting that estrogen suppresses the expression of dopamine receptors; however, the exact mechanism underlying the gender difference in DIP remains to be clarified. Genetic factors are also thought to be involved in the development of DIP because not all patients taking dopamine receptor blocking agents (DRBAs) develop parkinsonism. Meta-genome-wide screening showed that the genes associated with the gamma-aminobutyric acid (GABA) receptor signaling pathway are involved in neuroleptic-induced TD in schizophrenic patients, suggesting that genetic factors predispose to both DIP and TD.

**Etiology of Drug-Induced Parkinsonism**

Typical antipsychotics, also known as neuroleptics, are the most common causes of DIP. However, atypical antipsychotics, which were thought to be free from EPS, can also induce parkinsonism. In addition to antipsychotics, gastrointestinal (GI) motility drugs, calcium channel blockers (CCBs), and antiepileptic drugs have been found to induce DIP (Table 1).

**Antipsychotic drugs**

The history of DIP parallels that of antipsychotics. Parkinsonism as a side effect of chlorpromazine was first reported 3 years after its introduction. It was soon recognized that all typical antipsychotics had the potential to cause EPS, including parkinsonism, acute dystonia, akathisia, and TD. Typical antipsychotics include chlorpromazine, promazine, haloperidol, perphenazine, fluphenazine, and pimozide. About 80% of patients taking typical antipsychotic drugs exhibit more than one kind of EPS. Dopamine receptors are widely distributed in the brain, and typical antipsychotics may act on dopamine receptors in the striatum. Therefore, all patients taking antipsychotics have some risk of developing parkinsonism and other EPS. Parkinsonism usually appears days to weeks after starting antipsychotics, but in rare cases the onset delay may be several months or more. The risk of EPS was thought to be low for atypical antipsychotics. Atypical antipsychotics include clozapine, risperidone, olanzapine, quetiapine, and aripiprazole. It was originally thought that their relatively low frequency of associated EPS was due to them being more strongly antagonistic toward serotonin-2A receptors than toward dopamine receptors. This serotonin-dopamine hypothesis has long been considered a useful model for developing atypical antipsychotics that exhibit superior antipsychotic efficacy with a lower incidence of EPS compared to typical antipsychotics. The recent ‘fast-off’ theory suggested that their rapid dissociation from D2 receptors after they have blocked them can explain their lower risk of EPS.

In 1989, clozapine became the first atypical antipsychotic drug to be approved by the US Food and Drug Administration. It is effective in schizophrenic patients with drug-resistant negative symptoms, with an almost complete absence of EPS. DIP due to clozapine has not been reported, and it was found to improve psychosis without aggravating parkinsonism even in PD patients. However, clozapine has been associated with agranulocytosis in about 1% of patients, making physicians reluctant to prescribe this drug. Other atypical antipsychotics without the risk of agranulocytosis were developed to control psychosis with minimal EPS. Risperidone was expected to have a minimal risk of EPS because it has a high affinity for serotonin receptors. However, it binds D2 receptors in a dose-dependent manner, thus inducing parkinsonism and EPS to a similar extent as high doses of typical antipsychotics. Although the molecular structure of olanzapine is similar to that of clozapine, it carries with it a significant risk of EPS. Quetiapine is an atypical antipsychotic with a low risk of EPS and a low risk of aggravation of parkinsonism when used to treat psychotic symptoms in patients with PD, and is therefore apparently safe for use in elderly patients. Aripiprazole is the

### Table 1. Common offending drugs of drug-induced parkinsonism

| Drug frequently causing parkinsonism | Drug infrequently causing parkinsonism |
|-------------------------------------|----------------------------------------|
| Typical antipsychotics              | Atypical antipsychotics                |
| Phenothiazine, chlorpromazine, prochlorperazine, perphenazine, fluphenazine, promethazine | Atypical antipsychotics                |
| Butyrophenones, haloperidol          | Mood stabilizer                        |
| Diphenylbutylpiperidine, pimozide    | Lithium                                |
| Benzamide substitutes: sulpiride     |                                        |
| Atypical antipsychotics             | Antidepressant                          |
| Risperidone, olanzapine, ziprasidone, aripiprazole | SSRI: citalopram, fluoxetine, prazosin, sertraline |
| Dopamine depleters                  | Antiepileptic drugs                    |
| Reserpine, tetrabenazine             | Valproic acid, phenytoin               |
| Antiemetics                          | Aminetetics                             |
| Metoclopramide, levsulpiride, clebopride | Domperidone, ilotride                  |
| Calcium-channel blocker              |                                        |
| Flunarizine, cinnaeirazine           |                                        |

SSRI: selective serotonin reuptake inhibitor.

16 J Clin Neurol 2012;8:15-21
most recently introduced novel atypical antipsychotic, and has a unique mechanism of action. Although it was expected to have a low risk of EPS, clinical experiences have been disappointing.\textsuperscript{35-36} Thus, to date, only clozapine and quetiapine are associated with low rates of DIP in older patients.

**GI motility drugs**

GI prokinetic drugs, including metoclopramide, levosulpiride, clebopride, itopride, and domperidone, have also been associated with DIP. These drugs have been used clinically to manage motor disorders of the upper GI tract, including functional dyspepsia and emesis. The prokinetic effect of these drugs is mediated through their blockade of enteric inhibitory D\textsubscript{2} receptors.\textsuperscript{37} Besides binding to receptors in the peripheral organs, thus inducing antiemetic effects via D\textsubscript{2} receptor blockade in the area postrema, they also antagonize central D\textsubscript{2} receptors, leading to adverse effects including hyperprolactinemia and EPS. All prokinetics with D\textsubscript{2}-receptor-antagonizing properties have been found to induce EPS, although the extent of symptoms varies. Among the GI prokinetics, metoclopramide is the most-well-known cause of drug-induced movement disorders.\textsuperscript{17,38-40} Furthermore, levosulpiride is used widely in several Asian and European countries to treat nausea, vomiting, and functional dyspepsia. Until recently, the drug-induced movement disorders related to levosulpiride were under-recognized, but it has now been shown that levosulpiride frequently causes parkinsonism.\textsuperscript{41} Whereas metoclopramide usually induces TD, levosulpiride causes parkinsonism more frequently than TD or other EPS. Although metoclopramide and levosulpiride have the same mechanism of action, they show different patterns of adverse effects, the reason for which remains to be clarified. In general, domperidone is considered to be safe for the management of GI discomfort, even in patients with PD, because it does not cross the blood-brain barrier.\textsuperscript{37} However, although rare, acute dystonic reactions to this drug may occur.\textsuperscript{42}

**Other drugs**

Parkinsonism may also be caused by CCBs, antiepileptic drugs, and lithium, although their exact pathophysiological mechanisms are not yet known. CCBs including flunarizine and cinnarizine, which are used to control dizziness and headache,\textsuperscript{43} may cause DIP by reducing dopamine neurotransmission or acting directly on dopamine receptors.\textsuperscript{44-47} The long-term use of valproic acid was found to induce parkinsonism in 5\% of patients,\textsuperscript{48-50} while the exact underlying pathophysiology has not been clarified, one suggestion is that it is due to oxidative stress and mitochondrial dysfunction.\textsuperscript{50} Lithium is an infrequent cause of DIP and is thought to act by decreasing dopamine in the striatum, or as an anticholinesterase that increases central cholinergic activity.\textsuperscript{51,52}

### Clinical Characteristics of Drug-Induced Parkinsonism

DIP is generally characterized clinically as bilateral and symmetric parkinsonism, with more prominent bradykinesia and rigidity than in patients with PD.\textsuperscript{13,53} However, it has been suggested in many studies that clinical manifestations alone cannot be used to differentiate DIP from PD (Table 2).

Typically 30-50\% of patients with DIP show asymmetric parkinsonism and tremor at rest; these characteristics are considered to support a PD diagnosis.\textsuperscript{16,41,44-46,54} Interestingly, DIP patients with typical tremor at rest tremors usually also have postural tremors. The similar clinical manifestations of DIP and PD indicate that patients with DIP may have been in a preclinical stage of PD and that their parkinsonism may have been unmasked by the offending drugs. This is supported by findings that parkinsonism persists or even progresses after cessation of the drug in many DIP patients.\textsuperscript{17,30,40,41,45} In a long-term follow-up study, parkinsonism reappeared several months after the complete resolution from DIP in 7\% of patients;\textsuperscript{41} DIP might have antecedent PD in those patients.

The mean age at symptom onset is greater for DIP than for PD patients. In addition, DIP is more common in females, whereas males are more likely to have PD. While the duration of symptoms is similar, neurological deficits are more severe in

### Table 2. Clinical characteristics of drug-induced parkinsonism

| Drug                        | Mean age | Gender | Asymmetric parkinsonism (percentage of number of patients) | Tremor (percentage of number of patients) | Combined TD (percentage of number of patients) |
|-----------------------------|----------|--------|----------------------------------------------------------|-------------------------------------------|-----------------------------------------------|
| Hardie et al.\textsuperscript{14} | 61       | F>M    | 44                                                       | Not described                             | 42                                            |
| Hassin-Baer et al.\textsuperscript{14} | 46       | F<M    | 41                                                       | Rest tremor (44\%)                        | 45                                            |
| Garcia-Ruiz et al.\textsuperscript{44} | 71       | P>M    | 36                                                       | Postural tremor                           | Rest tremor                                   |
| Martí-Massó et al.\textsuperscript{45} | 75       | P>M    | 13                                                       | Postural tremor (51\%)                    | 7                                             |
| Shin et al.\textsuperscript{41} | 70       | P>M    | 38                                                       | Rest tremor (45\%)                       | Postural tremor (53\%)                       |

F: female, M: male, TD: tardive dyskinesia.
Drug-Induced Parkinsonism

DIP than in PD. Orolingual dyskinesia in drug-naïve PD patients is rare, whereas it is frequently combined with parkinsonism in DIP patients. The symmetry and type of tremor are not useful in distinguishing DIP from PD.

Pathophysiology of Drug-Induced Parkinsonism

Dopamine receptors in the brain consist of those of the D₁ family, comprising D₁ and D₂ receptors, and the D₂ family, comprising D₂a, D₂b, and D₃ receptors. The central dopaminergic system consists of the mesolimbic, mesocortical, tuberoinfundibular, and nigrostriatal pathways. All antipsychotic drugs have potent D₂ receptor blocking capacity and the therapeutic effects of these drugs on psychosis are related to their action on the limbic system, where they reduce dopamine transmission. The blockage of D₂ receptors by antipsychotic drugs in the striatum leads to disinhibition of GABA- and encephalin-containing striatal neurons at the origin of the indirect pathway with resultant alteration of the direct pathway, followed by disinhibition of the subthalamic nucleus. This leads to increased GABAergic inhibition of the thalamocortical projection by facilitation of the inhibitory projection from the globus pallidus/substantia nigra pars reticulata (Fig. 1A). This pathway resembles the model of disturbance of the basal ganglia-motor loop in PD. More than 80% of D₂ receptors were found to be occupied in patients with EPS who were taking neuroleptics, in agreement with results showing that clinical symptoms of PD began when over 80% of nigral neurons had degenerated.

TD, defined as hyperkinetic movement in the orolingual or oromandibular area, is caused by long-term use of dopaminergic blocking agents, and frequently accompanies DIP. The occurrence of TD with parkinsonism may be due to dopaminergic receptor supersensitivity resulting from long-term D₂ receptor blockage. Chronic administration of these drugs increases dopamine D₂ receptor density in the striatum. Moreover, withdrawal from neuroleptics was found to aggravate dyskinetic symptoms, whereas increased doses of neuroleptics transiently suppressed dyskinesia. D₂ receptors may also be involved in the development of orolingual dyskinesia when D₂ receptors are chronically blocked. Chronic administration of D₂ receptor blockers also induces changes in the direct pathway of the basal ganglia-motor loop to activate the striatopallidal pathway and increase the inhibition of the striatopallidal pathway (Fig. 1B). This imbalance between direct (D₁) and indirect (D₂) motor pathways and the resulting alterations in the globus pallidus/substantia nigra pars reticulata complex may lead to hyperkinetic orolingual movements, thus explaining the coexistent and sequential development of parkinsonism and dyskinesia.

Diagnosis of DIP and the Role of DAT Imaging

The clinical diagnostic criteria for DIP are defined as 1) the presence of parkinsonism, 2) no history of parkinsonism before the use of the offending drug, and 3) onset of parkinsonian symptoms during use of the offending drug. Since asymmetrical rest tremors are common in many DIP patients and symptoms persist or progress after cessation of the offending drug, patients clinically diagnosed with DIP may include individuals in the preclinical stage of PD whose symptoms were unmasked.

![Fig. 1. Changes in basal ganglia-thalamocortical motor loop due to blockade of D₂ receptors by DRBAs. The blockage of D₂ receptors by DRBAs in the striatum leads to disinhibition of GABA- and encephalin-containing striatal neurons at the origin of the indirect pathway, followed by a disinhibition of the subthalamic nucleus. This leads to increased GABAergic inhibition of the thalamocortical projection by facilitation of the inhibitory projection from the GPi/SNr (A). Chronic D₂ receptor blockage also induces changes in the direct pathways of the basal ganglia-motor loop to cause orolingual dyskinesia (B). DA: dopamine, DRBAs: dopamine receptor blocking agents, GABA: gamma-aminobutyric acid, GPe: globus pallidus pars externa, GPi: globus pallidus pars internal, SNr: substantia nigra pars compacta, SNc: substantia nigra pars reticulata, STN: subthalamic nucleus, TD: tardive dyskinesia.](image)
Dopamine receptor blocking agents, PD: Parkinson’s disease. DAT: dopamine transporter, DIP: drug-induced parkinsonism, DRBA: dopamine receptor blocking agents (B). A patient who was diagnosed with PD unmasked by DRBAs (B). Whereas it decreased severely in the right striatum in a patient (A), whereas it was normal and symmetric in the bilateral striatum in a pure DIP patient (A). Therefore, most DAT imaging studies, including those in patients with DIP, have utilized SPECT.

DAT uptake in the striatum is significantly decreased in PD patients, even during the early stages of the disease, because the motor symptoms of PD do not appear until 60-80% of dopaminergic neurons degenerate. In addition, drugs causing parkinsonism, such as DRBAs, have negligible affinity for DAT. DAT scans may show symmetric uptake of radiotracer in the bilateral striatum in patients with pure DIP, even if they have significant parkinsonism (Fig. 2A). PD can be diagnosed in DIP patients whose DAT uptake decreases asymmetrically in the striatum (Fig. 2B). Therefore, DAT scans may be useful for differentiating PD unmasked by drugs from pure DIP. Follow-up DAT scans of DIP patients with initially normal DAT scans exhibited normal uptake in the striatum, whereas scans of patients with initially decreased DAT scans in the striatum exhibited progressive reduction of striatal tracer uptake.

These findings suggest that DAT imaging provides useful information for diagnosing DIP and may help in the design of a therapeutic plan. Further studies involving larger numbers of patients are needed to determine the sensitivity and specificity of DAT scans in DIP diagnosis.

Treatments and Outcomes of DIP

DIP is generally treated by cessation of the offending drugs. Patients who cannot stop taking antipsychotic drugs because of their psychiatric diseases, such as those with schizophrenia or major depressive disorders, may be switched to atypical antipsychotics that have a lower risk of EPS. People who are prescribed dopamine antagonists due to simple GI disturbance, headache, dizziness, or insomnia should stop taking the offending drugs as soon as possible. Anticholinergics including trihexyphenidyl, benztropine, amantadine, and levodopa have been empirically tested for their ability to relieve symptoms of DIP, but this has produced no clear evidence of their effects in DIP patients.

DIP usually resolves within weeks to months after stopping the offending drug; however, parkinsonism may persist or progress in 10-50% of patients. The prognosis of patients with DIP can be classified into the following types: 1) full and long-lasting recovery from DIP with no subsequent development of parkinsonism, 2) persistence but not progression of parkinsonism, 3) persistence and eventual worsening of parkinsonism, and 4) full remission of parkinsonism but later reappearance after discontinuation of the offending drug. Only patients classified as type 1 can be defined as having ‘pure DIP’, whereas those classified as type 3 or 4 may be in the preclinical stages of PD. The finding of Lewy bodies in two DIP patients who had completely recovered after stopping the offending drugs suggests that clinical course alone cannot distinguish ‘pure DIP’ among patients clinically diagnosed with DIP. In DIP patients classified as type 2, the persistence of parkinsonism may be due to permanent DRBA-induced damage to dopamine receptors. DAT imaging may be useful in diagnosing DIP patients and predicting their clinical course. A large prospective study may provide more information on the exact prognosis and factors related to recovery in DIP patients.

Conclusions

DIP is important because it is a common etiology of parkinsonism and is frequently either unrecognized or misdiagnosed as PD. In addition, parkinsonism in DIP patients is sufficiently severe to affect daily activities and may persist for long periods of time even after cessation of the offending drug. DAT imaging may be useful for accurately diagnosing patients with DIP.
Drug-Induced Parkinsonism

and may help to identify the clinical characteristics and exact prognosis of this disorder. About 50% of patients with DIP and other movement disorders are treated with DRBAs for conditions unrelated to psychosis, including depression, GI disturbance, anxiety, and insomnia. Physicians should avoid prescribing DRBAs and CCBs for inappropriate reasons such as anxiety, insomnia, dizziness or dyspepsia in elderly patients and should monitor these patients’ neurological signs, especially parkinsonism and other movement disorders, when prescribing these drugs.

Conflicts of Interest

The authors have no financial conflicts of interest.

REFERENCES

1. Miller LG, Jankovic J. Neurologic approach to drug-induced movement disorders: a study of 125 patients. South Med J 1990;83:525-532.
2. Montastruc JL, Llau ME, Rascol O, Senard JM. Drug-induced parkinsonism: a review. Fundam Clin Pharmacol 1994;8:293-306.
3. Sethi KD. Movement disorders induced by dopamine blocking agents. Semin Neurol 2001;21:59-68.
4. Esper CD, Factor SA. Failure of recognition of drug-induced parkinsonism in the elderly. Mov Disord 2008;23:401-404.
5. Kägi G, Bhatia KP, Tolosa E. The role of DAT-SPECT in movement disorders. J Neurol Neurosurg Psychiatry 2010;81:5-12.
6. Scherfler C, Schwarz J, Antonini A, Grosset D, Valdiciola F, Marek K, et al. Role of DAT-SPECT in the diagnostic work up of parkinsonism. Mov Disord 2007;22:1229-1238.
7. Hall RA, Jackson RB, Swain JM. Neurotoxic reactions resulting from chlorpromazine administration. J Am Med Assoc 1956;161:214-218.
8. Benito-León J, Bermejo-Pareja F, Rodríguez J, Molina JA, Gabriel R, Morales JM. Neurological Disorders in Central Spain (NEDICES) Study Group. Prevalence of PD and other types of parkinsonism in three elderly populations of central Spain. Mov Disord 2003;18:267-274.
9. Wenning GK, Kiechl S, Seppi K, Müller J, Högl B, Saletu M, et al. Prevalence of movement disorders in men and women aged 50-89 years (Bruneck Study cohort): a population-based study. Lancet Neurol 2005;4:815-820.
10. Barbosa MT, Caramelli P, Maia DP, Cunningham MC, Guerra HL, Lima-Costa MF, et al. Parkinsonism and Parkinson’s disease in the elderly: a community-based survey in Brazil (the Bambuí study). Mov Disord 2006;21:800-808.
11. Avorn J, Bohn RL, Mogun H, Gurwitz JH, Monane M, Evenit D, et al. Neuroleptic drug exposure and treatment of parkinsonism in the elderly: a case-control study. Am J Med 1995;99:48-52.
12. Susatia F, Fernandez HH. Drug-induced parkinsonism. Curr Treat Options Neurol 2009;11:162-169.
13. Tharwat B, Treadwell S. Drug induced parkinsonism: a common cause of parkinsonism in older people. Postgrad Med J 2009;85:322-326.
14. Volkow ND, Ding YS, Fowler JS, Wang GJ, Logan J, Gatley SJ, et al. Dopamine transporters decrease with age. J Nucl Med 1996;37:554-559.
15. Stephen PJ, Williamson J. Drug-induced parkinsonism in the elderly. Lancet 1984;2:1082-1083.
16. Hardie RJ, Lees AJ. Neuroleptic-induced Parkinson’s syndrome: clinical features and results of treatment with levodopa. J Neurol Neurosurg Psychiatry 1988;51:850-854.
17. Miller LG, Jankovic J. Metoclopramide-induced movement disorders, Clinical findings with a review of the literature. Arch Intern Med 1989;149:2486-2492.
18. Bedard P, Langelier P, Villeneuve A. Oestrogens and extrapyramidal system. Lancet 1977;2:1367-1368.
19. Metzer WS, Newton JE, Steele RW, Claybrook M, Paige SR, McMillan DE, et al. HLA antigens in drug-induced parkinsonism. Mov Disord 1989;4:121-128.
20. Liao YJ, Wang YC, Chen YJ, Bai YM, Lin CC, Liao DL, et al. Association analysis of polymorphisms in the N-methyl-D-aspartate (NMDA) receptor subunit 2B (GRIN2B) gene and tardive dyskinesia in schizophrenia. Psychiatry Res 2007;153:271-275.
21. Inada T, Koga M, Ishiguro H, Horiuchi Y, Suyu A, Yoshio T, et al. Pathway-based association analysis of genome-wide screening data suggest that genes associated with the gamma-aminobutyric acid receptor signaling pathway are involved in neuroleptic-induced, treatment-resistant tardive dyskinesia. Pharmacogenet Genomics 2008;18:317-323.
22. Sharon J, Kaplan SM, Pierce CM, Ross WD. An interesting reaction to a tranquilizer: tonic seizures with perphenazine (trilafon). Am J Psychiatry 1957;114:556.
23. Christian CD, Paulson G. Severe motility disturbance after small doses of prochlorperazine. N Engl J Med 1958;259:828-830.
24. Janno S, Holm M, Tuiska K, Wahlbeck K. Prevalence of neuroleptic-induced movement disorders in chronic schizophrenia inpatients. Am Psychiatry 2004;161:160-163.
25. Ossowska K. Neuronal basis of neuroleptic-induced extrapyramidal side effects. Pol J Pharmacol 2002;54:299-312.
26. Tarsy D, Baldessarini RJ, Tarnai FI. Effects of newer antipsychotics on extrapyramidal function. CNS Drugs 2002;16:23-45.
27. Kuruki T, Nagao N, Nakahara T. Neuropharmacology of second-generation antipsychotic drugs: a validity of the serotonin-dopamine hypothesis. Prog Brain Res 2008;172:199-212.
28. Kapur S, Seeman P. Does fast dissociation from the dopamine d(2) receptor explain the action of atypical antipsychotics?: a new hypothesis. Am J Psychiatry 2001;158:360-369.
29. Stahl SM. “Hit-and-Run” actions at dopamine receptors, part 2: illustrating fast dissociation from dopamine receptors that typifies atypical antipsychotics. J Clin Psychiatry 2001;62:747-748.
30. Low-dose clozapine for the treatment of drug-induced psychosis in Parkinson’s disease. The Parkinson Study Group. N Engl J Med 1999;340:757-763.
31. Schotte A, Janssen PF, Gommersen W, Luyten WH, Van Gompel PJ, Lesage AS, et al. Risperidone compared with new and reference antipsychotics: in vitro and in vivo receptor binding. Psychopharmacology (Berl) 1996;124:57-73.
32. Gómez JC, Sacristán JA, Hernández J, Breier A, Ruiz Carrasco P, Antón Saiz C, et al. The safety of olanzapine compared with other antipsychotic drugs: results of an observational prospective study in patients with schizophrenia (EFESO Study). Pharmacopsychiatriological Study of Olanzapine in Schizophrenia. J Clin Psychiatry 2000;61:335-343.
33. Rabey JM, Prokhorov T, Miniovitz A, Dobronevsky E, Klein C. Effect of quetiapine in psychotic Parkinson’s disease patients: a double-blind labeled study of 3 months’ duration. Mov Disord 2007;22:313-318.
34. Juncos JL, Roberts VJ, Evatt ML, Jertwood RD, Wood CD, Potter LS, et al. Quetiapine improves psychotic symptoms and cognition in Parkinson’s disease. Mov Disord 2004;19:29-35.
35. Wickremaratnchi M, Morris HR, Ali JM. Aripiprazole associated with severe exacerbation of Parkinson’s disease. Mov Disord 2006;21:1538-1539.
36. Sharma A, Sorrell JH. Aripiprazole-induced parkinsonism. Int Clin Psychopharmacol 2006;21:127-129.
37. Tonini M, Cipollina L, Poluzzi E, Crema F, Corazza GR, De Ponti F. Review article: clinical implications of enteric and central D2 receptor blockade by antipsychomimetic gastrointestinal prokinetics. Aliment Pharmacol Ther 2004;19:379-390.
38. Kerney C, Hunter C, Davidson A, Jankovic J. Metoclopramide, an increasingly recognized cause of tardive dyskinesia. J Clin Pharmacol 2008;48:379-384.
39. Parkinsonism: an under-recognised complication of metoclopramide use. *Ceylon Med J* 1996;41:125.
40. Sethi KD, Patel B, Meador KJ. Metoclopramide-induced parkinsonism. *South Med J* 1989;82:1581-1582.
41. Shin HW, Kim MJ, Kim JS, Lee MC, Chung SJ. Levosulpiride-induced movement disorders. *Mov Disord* 2009;24:2249-2253.
42. Madej TH. Domperidone—an acute dystonic reaction. *Anaesthesia* 1985;40:202.
43. Sørensen PS, Hansen K, Olesen J. A placebo-controlled, double-blind, cross-over trial of flunarizine in common migraine. *Cephalalgia* 1986;6:7-14.
44. Garcı́a-Ruiz PJ, Javier Jiménez-Jiménez F, Garcı́a de Yébenes J. Calcium channel blocker-induced parkinsonism: clinical features and comparisons with Parkinson’s disease. *Parkinsonism Relat Disord* 1998;4:211-214.
45. Martí-Massó JF, Poza JJ. Cinnarizine-induced parkinsonism: ten years later. *Mov Disord* 1998;13:453-456.
46. Negrotti A, Calzetti S. A long-term follow-up study of cinnarizine- and flunarizine-induced parkinsonism. *Mov Disord* 1997;12:107-110.
47. Takada M, Kono T, Kitai ST. Flunarizine induces a transient loss of tyrosine hydroxylase immunoreactivity in nigrostriatal neurons. *Brain Res* 1992;590:311-315.
48. Onofri M, Thomas A, Pacci C. Reversible parkinsonism induced by prolonged treatment with valproate. *J Neurol* 1998;245:794-796.
49. Easterford K, Clough P, Kellett M, Fallon K, Duncan S. Reversible parkinsonism with normal beta-CTF-SPECT in patients exposed to sodium valproate. *Neurology* 2004;62:1435-1437.
50. Jamora D, Lim SH, Pan A, Tan L, Tan EK. Valproate-induced Parkinsonism in epilepsy patients. *Mov Disord* 2007;22:130-133.
51. Reches A, Tietger J, Lavy S. Parkinsonism due to lithium carbonate poisoning. *Arch Neurol* 1981;38:471.
52. Holroyd S, Smith D. Disabling parkinsonism due to lithium: a case report. *J Geriatr Psychiatry Neurol* 1995;8:118-119.
53. Tinazzi M, Antonini A, Bovi T, Pasquin I, Steinmayr M, Mampetti R. Dopamine transporter ligand 18F-FECNT. *J Nucl Med* 2010;51:1331-1334.
54. Diaz-Corralles FJ, Sanz-Viedma S, Garcia-Solis D, Escobar-Delgado T, Mir P. Clinical features and 123I-FP-CIT SPECT imaging in drug-induced parkinsonism and Parkinson’s disease. *Eur J Nucl Med Mol Imaging* 2010;37:556-564.
55. Reches A, Tietger J, Lavy S. Parkinsonism due to lithium carbonate poisoning. *Arch Neurol* 1981;38:471.
56. Holroyd S, Smith D. Disabling parkinsonism due to lithium: a case report. *J Geriatr Psychiatry Neurol* 1995;8:118-119.
57. Tinazzi M, Antonini A, Bovi T, Pasquin I, Steinmayr M, Mampetti R. Dopamine transporter ligand 18F-FECNT. *J Nucl Med* 2010;51:1331-1334.
58. Reches A, Tietger J, Lavy S. Parkinsonism due to lithium carbonate poisoning. *Arch Neurol* 1981;38:471.
59. Holroyd S, Smith D. Disabling parkinsonism due to lithium: a case report. *J Geriatr Psychiatry Neurol* 1995;8:118-119.
60. Tinazzi M, Antonini A, Bovi T, Pasquin I, Steinmayr M, Mampetti R. Dopamine transporter ligand 18F-FECNT. *J Nucl Med* 2010;51:1331-1334.
61. Davis MR, Votaw JR, Brenner JD, Byas-Smith MG, Faber TL, Voll RJ, et al. Initial human PET imaging studies with the dopamine transporter ligand 18F-FECNT. *J Nucl Med* 2003;44:855-861.
62. Varrone A, Halldin C. Molecular imaging of the dopamine transporter. *J Nucl Med* 2010;51:1331-1334.
63. Diaz-Corralles FJ, Sanz-Viedma S, Garcia-Solis D, Escobar-Delgado T, Mir P. Clinical features and 123I-FP-CIT SPECT imaging in drug-induced parkinsonism and Parkinson’s disease. *Eur J Nucl Med Mol Imaging* 2010;37:556-564.
64. Tinazzi M, Ottaviani S, Isaias IU, Pasquin I, Steinmayr M, Vampini C, et al. [123I]FP-CIT SPECT imaging in drug-induced Parkinsonism. *Mov Disord* 2008;23:1825-1829.
65. Poewe W, Scherfler C. Role of dopamine transporter imaging in investigation of parkinsonian syndromes in routine clinical practice. *Mov Disord* 2003;18 Suppl 7:S16-S21.
66. Lavalaye J, Linszen DH, Booji J, Dingemans PM, Reneman L, Habraken JB, et al. Dopamine transporter density in young patients with schizophrenia assessed with [123I]FP-CIT SPECT. *Schizophr Res* 2001;47:59-67.
67. Tolosa E, Coelho M, Gallardo M. DAT imaging in drug-induced and psychogenic parkinsonism. *Mov Disord* 2003;18 Suppl 7:S28-S33.
68. Fann WE, Lake CR. Amantadine versus trihexyphenidyl in the treatment of neuroleptic-induced parkinsonism. *Am J Psychiatry* 1976;133:940-943.
69. Jankovic J. Tardive syndromes and other drug-induced movement disorders. *Clin Neuropharmacol* 1995;18:197-214.
70. Rajput AH, Zoldijsky B, Hornykiewicz O, Shannak K, Lee T, Seeman P. Reversible drug-induced parkinsonism. Clinico-pathologic study of two cases. *Arch Neurol* 1982;39:644-646.
71. Bishnoi M, Chopra K, Kulkarni SK. Protective effect of adenosine reuptake inhibitors in haloperidol-induced orofacial dyskinesia and associated behavioural, biochemical and neurochemical changes. *Pharmacology* 2007;79:171-183.
72. Tsai G, Goff DC, Chang RW, Flood J, Baer L, Coyle JT. Markers of glutamatergic neurotransmission and oxidative stress associated with tardive dyskinesia. *Am J Psychiatry* 1998;155:1207-1213.