Apomorphine in the treatment of Parkinson’s disease: a review

O uso da apomorfina no tratamento da doença de Parkinson: revisão da literatura

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ABSTRACT
Optimizing idiopathic Parkinson’s disease treatment is a challenging, multifaceted and continuous process with direct impact on patients’ quality of life. The basic tenet of this task entails tailored therapy, allowing for optimal motor function with the fewest adverse effects. Apomorphine, a dopamine agonist used as rescue therapy for patients with motor fluctuations, with potential positive effects on nonmotor symptoms, is the only antiparkinsonian agent whose capacity to control motor symptoms is comparable to that of levodopa. Subcutaneous administration, either as an intermittent injection or as continuous infusion, appears to be the most effective and tolerable route. This review summarizes the historical background, structure, mechanism of action, indications, contraindications and side effects, compares apomorphine infusion therapy with other treatments, such as oral therapy, deep brain stimulation and continuous enteral infusion of levodopa/carbidopa gel, and gives practical instructions on how to initiate treatment.

Keywords: Apomorphine; dopaminergic agents; Parkinson’s disease; review.

The management of Parkinson’s disease (PD) aims for adequate control of motor and nonmotor symptoms, minimizing the adverse effects of medications1. A wide range of therapeutic options are available and can be used to tailor treatment to the needs of individual patients. These include dopamine replacement therapy using levodopa; direct stimulation of striatal dopamine receptors by dopamine agonists; and other interventions in dopamine metabolism using monoamine oxidase-B, DOPA decarboxylase or catechol-O-methyltransferase inhibitors2. Despite being used since 1960, and enduring as the most powerful antiparkinsonian drug, levodopa is associated with a high incidence of motor complications3, with 24% to 89% of patients developing dyskinesias after long-term continuous exposure to this drug4. There is debatable data on the role of dopamine agonists in postponing or minimizing these complications when used as monotherapy or in combination with lower doses of levodopa4. In addition, PD presents almost invariably with nonmotor symptoms that include mood and cognitive disorders (anxiety, depression, dementia and psychosis), autonomic dysfunction (urinary incontinence, constipation, dysphagia, gastroparesis, erectile dysfunction, orthostatic...

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hypotension, excessive sweating, drooling), pain, and sleep disturbances, that have a significant impact on quality of life, comparable or occasionally more severe than the motor aspects of the disease.

Given the complexity of PD and its multiple layers of relevant clinical implications, it is quite common that, even with optimal therapy, some patients remain inadequately controlled, requiring invasive treatment, i.e., deep brain stimulation (DBS), enterally-delivered levodopa/carbidopa gel and subcutaneous apomorphine.

The objective of this review is to describe the current role of apomorphine in the treatment of PD.

METHODS

We performed a review of the literature based on review articles, expert opinion manuscripts and clinical trial reports from the PubMed database, using the following descriptors: apomorphine, Parkinson’s disease, motor fluctuations, deep brain stimulation and intestinal levodopa/carbidopa. The search was limited to articles published between 1984 and 2017.

Historical review

Apomorphine was the first dopamine agonist with powerful antiparkinsonian effects used in clinical practice and predated levodopa by ten years. Derived from morphine, it was initially used as an emetic, expectorant, sedative, antipsychotic and anticonvulsant, as well as for managing drug and alcohol addiction. Apomorphine is a highly-lipophilic, short-acting, nonergot dopamine agonist that acts on D1 and D2 dopamine receptors. It was synthesized for the first time in 1869 by Matthiessen and Wright and, although Weill recommended its use for PD in 1884, the first trials started in 1950, according Wenzel, Cotzias, and Lees. In 1951, Schwab et al. observed improvements in rigidity and tremor in PD patients 5-10 minutes after subcutaneous administration of a 0.5 to 1.0 mg dose of apomorphine. These findings were later corroborated by Cotzias et al.

Oral administration of apomorphine required large doses to achieve the desired clinical response, and an exacerbated peripheral response was therefore common. This could include nausea, vomiting, postural hypotension and impaired kidney function, reflected in elevated urea and creatinine. Various administration routes were explored to avoid these adverse effects, the most successful of which proved to be the subcutaneous.

Apomorphine is superior to other dopamine agonists such as lisuride because it has fewer adverse effects and induces significant improvement in motor and nonmotor symptoms, including hyperhidrosis, nocturia, urge incontinence, fatigue and mood disturbances, in addition to being well tolerated in patients with visual hallucinations, illusions and paranoid ideation. It also improves sleep disorders, such as insomnia and restless legs syndrome, without worsening daytime drowsiness.

In 1988, a group led by Lees, developed a mechanism for continuous subcutaneous apomorphine infusion, which was later routinely recommended for patients with severe, refractory “off” periods. Although continuous apomorphine treatment was first introduced by Stibe et al. in 1988, subcutaneous injections were only approved by the FDA for use in motor “off” periods in 2004. The long-term effectiveness of apomorphine as a rescue medication was also investigated in study APO302, in which 62 patients who had been having rescue injections for at least three months were assessed. In a comparative assessment on the UPDRS scales, there was a significant reduction in motor score after 10 and 20 minutes.

Mechanism of action of apomorphine

Apomorphine binds to pre- and postsynaptic receptors and exerts a therapeutic effect by direct stimulation of postsynaptic striatal dopamine D2 receptors, resulting in activation of the direct pathway and inactivation of the indirect striatopallidal pathways. The motor response occurs after a single dose of subcutaneous apomorphine and is similar to that of levodopa but with faster onset (approximately 4-12 minutes), with a mean effect duration of 45–60 minutes. In light of these therapeutic effects, it became one of the prototypic “rescue medications” in cases of unpredictable “off” periods, as in the case of patients with advanced PD and poorly-controlled motor fluctuations, who experience erratic gastric emptying. Furthermore, it does not share transport mechanisms or metabolic pathways with levodopa and, unlike levodopa, does not require an active transport mechanism to reach the central nervous system. Absorption varies with skin temperature and blood flow, and the best absorption is...
achieved when it is injected into the subcutaneous tissue of the abdominal wall. There is no interaction with cytochrome P450 inhibitors, and the cytochrome P450 system does not interfere with the metabolism of this dopamine agonist. Age, gender, disease duration, levodopa dosage or duration of apomorphine therapy do not appear to play any role in the clearance of the medication.1,11-13,15

When should apomorphine be used?

The first consideration for the use of apomorphine must be the confirmed diagnosis of levodopa-responsive PD.1,11,13,34 Age is typically not a limiting factor, and mild cognitive impairment and axial symptoms are not contraindications.3,12,13,14,2435 Indications for apomorphine include patients with refractory “off” periods, e.g., when there is a delay in the onset of the effects of orally administered medication; patients who have major “off” periods upon waking up; and patients with significant wearing-off periods.7,8,9,10,11,12,13,14,24,35 It can also improve nonmotor symptoms such as urinary disturbances3,12,13 and serve as a diagnostic clue when diagnosis of PD is uncertain.1,12,13

The therapeutic response to maximum levodopa doses does not vary in PD but is less consistent in patients with multiple system atrophy. Some patients with other parkinsonian syndromes are unresponsive or do not respond well to levodopa.1,11-13. A study by the group led by Lees showed that apomorphine can be used as a diagnostic test of response to levodopa and has an accuracy of 90%.1 The study concluded that a response to apomorphine supports a diagnosis of PD, while failure to respond indicates that this diagnosis is extremely unlikely.1

Routes of administration for apomorphine

Various routes of administration have been investigated.3,13,14,22,27,40 Oral administration was first tried by Cotzias, who increased the dose gradually to 1,500 mg a day.2,15 Tolerance of 150 to 1,440 mg daily doses was generally good, but higher doses caused azotemia.7,8,10,11,12,14 Bioavailability of apomorphine by this route is less than 4% and this route was therefore considered unfeasible, as it required very high doses to achieve the desired effect, leading to significant side effects such as nephrotoxicity, reflected in elevated creatinine and urea.7,8,10,11,12,14 Bioavailability of sublingual apomorphine is also low (10–22%), and a 3 mg subcutaneous dose and 30 mg sublingual dose have similar pharmacological profiles and clinical responses.1,7,27. Using this route, Ondo et al.27 reported an effect duration of between 60 and 130 minutes, while Hughes et al.29 reported a latency of up to 25 minutes and effect duration of 118 minutes. Apomorphine can cause nausea, orthostatic hypotension, an unpleasant taste and severe stomatitis, the latter being reported in about 50% of patients.7,8,10,11,12,14,15,16,17,18,19,20,21 The intranasal spray has a pharmacokinetic profile similar to that of subcutaneously administered apomorphine but requires pretreatment with domperidone.1,2,7,27,40,41 Latency to effect has been shown to be between 5 and 15 minutes, and the effect duration between 30 and 60 minutes.1,2,7,27,40 In a study by Obering et al., the “off” period was significantly reduced from 5.3 hours/day to 3.8 hours/day. In another study on the use of apomorphine in PD, no statistically significant difference was observed between levodopa and apomorphine in terms of the UPDRS score.18

Intermittent apomorphine injection

Ideally, the first dose should be administered in a hospital setting so that the clinical response can be observed and a tailored therapeutic dose identified.1,14,15

Peripheral dopaminergic adverse events can occur as a complication of injections of apomorphine, particularly nausea. Oral domperidone (10–20 mg three times a day) should be started one to three days before apomorphine therapy (Box 1).

Currently, penject (intermittent injection) and portable minipumps (continuous infusion) have been approved in most European countries (Figures 2 and 3), where they play an important role in advanced PD treatment and have yielded good results.11,14,15,31,35,36 Hughes et al.1 published a study of 71 patients treated with intermittent injections (10/day) or continuous infusion (when more than 10 injections were needed). In 49 patients treated with intermittent injections, there was a 50% reduction in “off” time. After one year of treatment, 80% of the patients reported that the therapy was still effective.1

Intermittent infusion of apomorphine (penject) is performed with an insulin syringe mounted in an injector pen with premarked doses for ease of administration.11,14,15. Injections sites can be administered in the abdominal region, arms and thighs.1,12,14 The number of injections can vary between 1 and 30, and the dose can vary between 1 mg

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Box 1. Apomorphine test.

1) Pretreatment with domperidone. (10–20 mg three times/day, started one to three days before apomorphine therapy)

2) Performed when the patient has been off dopamine medication for 12-24 hours. Levodopa should be suspended the night before the test.

3) Measure blood pressure with the patient lying down, and seated, before the test.

4) Start with 1 mg to 1.5 mg subcutaneous apomorphine. Record the UPDRS-III score (pre- and post-apomorphine – OFF, ON, OFF) and any adverse effects (nausea, orthostatic hypotension, drowsiness, and dyskinesias).

5) Repeat administration of apomorphine at intervals of 1 to 1.5 hours, increasing the dose by 1 mg until a good or acceptable clinical effect is observed.

No more than 7–8 mg per hour of apomorphine should be administered.
and 10 mg per injection\textsuperscript{11,23}. Apomorphine solution is supplied at a concentration of 1 mg/0.1 mL in 3 mL (30 mg) glass cartridges with a manual reusable multidose injector pen that releases doses from 0.02 mL to 1 mL (10 mg)\textsuperscript{11,15,23}. To minimize mistakes, the apomorphine dose should be prescribed in milliliters (rather than milligrams) as the injector pen uses this unit. An optimal therapeutic dose is between 0.3 mL and 0.5 mL, or 3 mg and 5 mg\textsuperscript{11,23}. Once a suitable dose has been identified, it rarely needs to be adjusted. The half-life of apomorphine is 45 minutes, and the minimum recommended time between injections is 60 minutes\textsuperscript{11,23}. The drug is absorbed quickly after subcutaneous injection, reaching maximum serum levels in 20 minutes, and the clinical effect can be observed between 5 and 15 minutes after administration\textsuperscript{11,13,23}. Apomorphine is generally indicated for short periods. The mean daily dose is 3–30 mg/day, and it is important to determine the dose required to reverse the “off” period\textsuperscript{11,23}. Intermittent apomorphine infusion is a good rescue therapy in cases of motor fluctuations, such as wearing-off and on-off fluctuations, because of its fast action\textsuperscript{11,12,23}. When required, apomorphine should be used as a rescue medication during “off” periods, without changing the levodopa schedule. It can reverse predictable and unpredictable “off” periods and is indicated when the “on” period is delayed\textsuperscript{11,15,23}. It helps with morning or nocturnal akinesia as well as painful dystonias, and is a suitable choice when absorption of orally administered levodopa is hampered by delayed gastric emptying\textsuperscript{11,12,23}. Apomorphine can also improve psychiatric symptoms such as depression and panic attacks\textsuperscript{24}. Some surgical centers use apomorphine administered by penject as a rescue medication in preoperative patients when oral medication cannot be administered\textsuperscript{11,25}.

**Continuous apomorphine infusion**

An infusion pump is recommended when “off” periods are poorly controlled by oral treatment or when apomorphine injections are effective but required more frequently (more than 4–6 times a day)\textsuperscript{12,23,14,6}. The patient can be kept in a continuous “on” state with an improvement in dyskinesias, and the levodopa dose can be reduced\textsuperscript{11,21,36}. Like intermittent infusion, continuous infusion of apomorphine helps nonmotor symptoms such as pain and mood swings. This form of administration is recommended for patients in whom duodenal levodopa infusion and DBS are contraindicated. Of the options available for advanced stages of the disease, it is the least invasive\textsuperscript{36}. Unlike with DBS, age and neuropsychiatric changes are not absolute contraindications for apomorphine infusion\textsuperscript{11,46,47}. Continuous apomorphine infusion is also an alternative to oral treatment, which is complex and may have limited adherence or may not result in adequate absorption, as it minimizes drug interactions when the patient is taking several medications\textsuperscript{6,11,23}. Various studies have shown that the levodopa dose can be reduced by between 16% and 18% after apomorphine is started\textsuperscript{11,12,23}. Apomorphine can also reduce the “off” period by 50% to 80% and guarantees patient mobility during the day even in the absence of levodopa\textsuperscript{6,11,23}.

The initial assessment should include an electrocardiogram to exclude the presence of a long QT interval,
tachycardia, bradyarrhythmias, atrial fibrillation and premature ventricular contractions, and should exclude pre-existing hemolytic anemia. Box 2 shows the main contraindications for use of apomorphine.

One day before starting apomorphine treatment, patients should be pretreated with 10 mg of domperidone, which can be discontinued as soon as possible, when the adverse effects of apomorphine have been controlled. The teflon delivery needle is inserted in the subcutaneous tissue of the abdominal wall, and the site is changed at least every day. Continuous infusion is usually started in a hospital setting, with a flow rate of 0.5 or 1 mg/hour during the first day. The infusion cycle is generally 12–24 hours a day (typically 16 hours), and the usual dose is 4-7 mg/hour. The hourly flow rate is adjusted depending on its effectiveness and patient tolerability. Generally, the rate is increased by 0.5-1.0 mg/hour every day during the initial period and more slowly after that at weekly intervals. Some patients are encouraged to use a booster dose in anticipation of “off” periods.

In patients being treated with continuous subcutaneous infusion of apomorphine, oral dopamine agonists can gradually be discontinued during the apomorphine titration period. Sudden interruption of these agonists can lead to dopamine agonist withdrawal syndrome. Other antiparkinsonian medications (monoamine oxidase-B inhibitors, catechol-O-methyltransferase inhibitors, amantadine, anticholinergics) should then be discontinued gradually, usually in the first seven days. Levodopa is generally reduced when the desired therapeutic dose of apomorphine is reached but can be reduced when apomorphine is started if dyskinesia is present.

Adverse events

The most common long-term side effect, which occurs in up to 70% of individuals, is nodules at the injection site. Some patients may develop itching, bruising or pain. These local adverse effects are related to drug concentration, infusion time or injection depth. In some patients (10–20%) reactions may be more severe, and necrotic nodular ulcerations or panniculitis may occur. This can be solved by rotating the injection site every day, ensuring asepsis, applying silicone gel patches or, in some cases, it may be necessary to use ultrasound treatment. García Ruiz et al. found that sedation occurred in 29% of patients using an apomorphine pump. Orthostatic hypotension can be a manifestation of dysautonomia or due to dopamine stimulation and can be improved with domperidone. In some cases, patients may need to wear compression stockings, keep their legs raised and take salt tablets or even fludrocortisone together with midodrine. Hematological tests at regular intervals are recommended to avoid the risk of hemolytic anemia.

Electrocardiographic changes (QT prolongation) can occur with doses of 6 mg or more. Prophylactic treatment with dopamine antagonists (e.g., metoclopramide and prochlorperazine) and serotonin receptor antagonists (e.g., granisetron and ondansetron) should be avoided because these agents cross the blood-brain barrier and can interact with apomorphine. When used with ondansetron, apomorphine can cause severe hypotension and loss of consciousness. Concomitant use of 5-HT3 antagonists is contraindicated. Doses of more than 6 mg do not lead to additional benefits and are not recommended.

Table 1 summarizes the main adverse effects of subcutaneous apomorphine.

### Conventional oral treatment vs. apomorphine

The quality of the response to oral levodopa is indistinguishable from the quality of the response to apomorphine. Apomorphine, however, produces a shorter motor response, supporting the idea that the integrity of the postsynaptic receptors is the key factor that determines the dopamine response in Parkinson’s disease treatment. In other words, the clinical responses to the drugs are the same although they have different mechanisms of action.

Apomorphine has various advantages over levodopa, such as the fact that it is a monotherapy and increases the “on” period by maintaining a continuous dopamine stimulus, reducing the need for levodopa and, in turn, reducing dyskinesias and motor fluctuations.

As it is administered parenterally, apomorphine improves treatment adherence in patients who cannot tolerate oral medicine or in whom absorption is erratic. However, it requires help from relatives or caregivers to handle the pump. Furthermore, apomorphine crosses the blood-brain barrier quickly without depending on an active transport system and without competing with proteins in the circulation.

### Other device-aided therapy strategies vs. apomorphine

Deep brain stimulation is currently widely accepted as an alternative in stages of PD when motor complications

<table>
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<th>Table 1. Adverse effects associated with apomorphine.</th>
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<tr>
<td><strong>Local</strong></td>
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<tr>
<td><strong>Systemic</strong></td>
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<tr>
<td><strong>Hematological</strong></td>
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<td><strong>Psychiatric</strong></td>
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are significant, or the symptoms of the disease are not well controlled. Deep brain stimulation of the subthalamic nucleus and apomorphine infusion produces significant improvements in parkinsonian symptoms and motor fluctuations through different mechanisms. The choice of therapy depends on the profiles of individual patients. Although the surgical procedure is considered relatively safe, it obviously is not risk-free and potential adverse effects of DBS include hypophonia and other bulbar symptoms, behavioral and cognitive changes, infections, transient confusion, seizures, intracranial bleeding, and various forms of hardware dysfunction. For example, a Spanish study that compared DBS and treatment with apomorphine showed that neuropsychological testing for the apomorphine group did not change, while for the patients who underwent DBS surgery, verbal phonemic fluency and word-naming speed were affected negatively. Additionally, apomorphine infusion therapies can be used in patients with severe cognitive impairment and psychiatric disorders. Borgemeester et al. published a retrospective, long-term follow-up study about continuous subcutaneous apomorphine infusion in PD patients with cognitive dysfunction. The study demonstrated that apomorphine infusion is an effective treatment in advanced PD patients with cognitive dysfunction, including visual hallucinations and orthostatic hypotension. Combined use of DBS and an apomorphine pump can be considered in patients with adverse effects on axial functions, such as altered gait or dysarthria.

Continuous enteral infusion of levodopa/carbidopa gel (LCIG) was tested in small studies published in 1986 and 1988, and then in a long-term study developed at the University of Uppsala, Sweden, as a therapeutic alternative for the advanced stage of the disease. It is a combination of levodopa (20 mg/mL) and carbidopa (5 mg/mL) in a pseudoplastic gel and is delivered directly into the proximal jejunum by means of a portable infusion pump and duodenal catheter. It has the advantage that it ensures a stable flow of dopamine into the striatum and, consequently, an increase in "on" time without dyskinesias. It also improves nonmotor symptoms such as drowsiness, fatigue, impaired attention, memory loss, and gastrointestinal, urinary and cardiovascular problems. Ricciardi et al. evaluated 24-hour infusion of LCIG in eight PD patients to address severe nocturnal dyskinesia unresponsive to oral therapies. They found significant improvements in fatigue and sleep quality, mood/cognition, hallucinations, and urinary function, and there was no change in motor severity or motor complications. The most common adverse effect is dyskinesia, although this is less common than with oral treatment. Long-term use of high doses of levodopa leads to increased homocysteine levels and reduced cobalamin metabolism. The risk of severe infections such as peritonitis is low. Nevertheless, such infections, if they occur,

Table 2. Comparison of the different treatment options.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Indication</th>
<th>Advantages of the procedure</th>
<th>Disadvantages of the procedure</th>
<th>Adverse effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deep brain stimulation</td>
<td>The best option in cases of dyskinesia</td>
<td>Less need for dopamine medication</td>
<td>Invasive therapy with surgical risks such as hemorrhage and infection</td>
<td>Worsening of neuropsychiatric function, cognitive changes Can lead to worsening of speech, postural instability and freezing of gait</td>
</tr>
<tr>
<td>Apomorphine</td>
<td>Rescue from “off” episodes</td>
<td>Mildly invasive</td>
<td>Parenteral administration (important for patients who cannot tolerate oral medication or have poor absorption)</td>
<td>Frequent blood tests and ECG are required ReQUIRES relatives or caregivers to handle the pump</td>
</tr>
<tr>
<td>Jejunal levodopa-carbidopa</td>
<td>Frequent &quot;off&quot; periods or severe dyskinesia</td>
<td>More physiological release of dopamine and less pulsatile stimulus</td>
<td>Crosses the blood-brain barrier quickly without any need for active transport and without competing with proteins in the circulation.</td>
<td>Can worsen dyskinesia Can cause anxiety, depression, hallucination and confusion</td>
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No age limit; mild and moderate dementia are not contraindications
can lead to death\textsuperscript{35-35}. Other adverse effects may include skin problems at the surgical site, weight loss and peripheral axonal neuropathy, the mechanism of which is poorly understood\textsuperscript{35}. There are small trials comparing apomorphine infusion and LCIG, showing that the apomorphine pump is easier for caregivers and patients to use, and is less expensive\textsuperscript{56-58}. However, if apomorphine infusion does not yield satisfactory results, jejunal levodopa may be indicated\textsuperscript{11,12,58}. The choice of apomorphine or LCIG depends on the individual patient and should take into account the adverse effects and technical aspects of each therapy\textsuperscript{11,12,58,59}. The expert consensus groups recognize the rapid and consistent relief from the symptoms of PD provided by apomorphine. Its mode of delivery is less invasive than DBS or LCIG, other therapies also considered for the treatment of this stage of the disease. Also, apomorphine infusion can be easily and immediately reversed, either when adverse effects occur or at the patient's request\textsuperscript{60}. Table 2 summarizes the main therapies discussed here.

CONCLUSION

Apomorphine is a treatment option for advanced Parkinson’s disease that is well tolerated and optimizes motor fluctuations and nonmotor symptoms frequently found in the condition. It is another treatment option that can be used to improve the patient’s quality of life.

References


