


Effective Management of “OFF” Episodes in Parkinson’s Disease: Emerging Treatment Strategies and Unmet Clinical Needs

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Abstract: Motor complications related to the chronic administration of levodopa and failure to prevent the neurodegenerative disease process counterbalance the pivotal discovery of levodopa as the cornerstone of PD treatment. Excellent motor control is offered early during the course of treatment, but this diminishes as pathological changes in the striatum lead to synaptic dopamine levels becoming completely dependent on exogenous dopamine. This non-physiologic stimulation of dopamine receptors eventually manifests as OFF episodes. As no disease modifying therapy exists for PD that can disrupt these pathological changes, most research and treatment focuses on optimization of dopaminergic stimulation of striatal receptors so that they mimic tonic, physiologic stimulation as closely as possible. Strategies focusing on these challenges have included non-pharmacologic approaches, optimizing levodopa pharmacokinetics, using adjunctive treatments including those with non-dopaminergic mechanisms, and implementing rescue therapies. Device aided therapies, including surgery, are also available. In this review, we will focus on effective management of motor symptoms related to OFF periods, including emerging strategies. Unmet clinical needs will be discussed, including non-motor symptoms, targeted molecular therapies and disease modifying therapy.

Keywords: continuous dopaminergic stimulation, advanced Parkinson’s Disease, motor complications, nonmotor symptoms, dyskinesias, chronic levodopa complications

Introduction

Parkinson’s Disease (PD) is the second most common neurodegenerative disorder, and its prevalence has doubled in the last 25 years.¹ It is well established that nigrostriatal dopamine deficiency remains a central factor in the development of the cardinal motor features of PD – tremor, rigidity, bradykinesia, and postural instability. Levodopa, a dopamine precursor, has been the mainstay of PD treatment for the last six decades. While levodopa can substantially ameliorate PD symptoms, it is associated with development of motor complications as the neurodegenerative process continues.¹

Normal function of the nigrostriatal pathway relies on tonic dopaminergic stimulation, accomplished by a group of tonically active dopaminergic neurons which maintain a steady flow of dopamine in the synapse, and a quiescent group which can become tonically activated and compensate if there is injury in the system.^{2,3} There is also a neuronal group which can become phasically activated and release bursts of dopamine, superimposed on background tonic activity.^{3,4} In PD, there is a loss of presynaptic dopaminergic terminals leading to a reduction in dopamine neurotransmission and resultant dysregulation of movement. Early in the disease, when dopaminergic neuronal loss is mild, there is enough stored dopamine which can function to passively stabilize synaptic dopamine levels by way of endogenous tonic input.² Thus, despite intermittent oral dosages of exogenous levodopa, there is constant stimulation of the post-synaptic receptors, and patients are generally unaware of increasing symptoms as plasma levodopa concentrations fall. However, as such doses cross the blood brain barrier, they far exceed the typical amount of endogenous dopamine release, and result in non-physiologic, pulsatile stimulation of dopamine receptors.²⁻⁴ Over the years as neuronal death

becomes more prevalent and dopamine stores are depleted, the endogenous buffering capacity is lost and the striatal dopamine concentrations closely reflect plasma levodopa levels. These plasma levels are prone to large fluctuations, and patients become aware of transitions between periods of good motor response (“ON” state) and periods of poor response/loss of response (“OFF” state).⁵

Motor complications of PD include motor fluctuations (MF) and dyskinesias, but non-motor fluctuations (NMF) can also occur.⁶ OFF symptoms can occur under various conditions (Table 1). NMF are defined as the fluctuating presence non-motor symptoms (NMS, neuropsychiatric, autonomic, sensory) in conjunction with the rise/fall of levodopa concentration, and often in association with MF, though non-dopaminergic neurotransmitter systems may also contribute.⁷ Levodopa-induced dyskinesias (LID) are hyperkinetic movements (eg, chorea, dystonia, ballism or a combination of these) that typically occur during periods of peak levodopa concentrations, and less commonly as a diphasic phenomenon, when levodopa concentrations are either rising or falling.⁸

It is important to note that pathological changes in the central nervous system (CNS) in PD do not fully account for the development of MF and OFF periods. Several pharmacokinetic properties of levodopa contribute to this, including its short plasma half-life, intestinal absorption and transport via amino acid transporters.⁹ Levodopa is readily metabolized by DOPA decarboxylase (DDC), catechol-O-methyltransferase (COMT), and monoamine oxidase B (MAO-B). Plasma half-life of levodopa is about 50 minutes; however, this increases to about 1.5 hours with the addition of a DDC inhibitor (eg, carbidopa or benserazide).¹⁰ Delayed gastric emptying slows the absorption of levodopa in the proximal small intestine.¹¹ Consumption of high protein meals reduces levodopa absorption as dietary proteins compete for amino acid transporters both in the gastrointestinal tract and the blood brain barrier.^{2,11,12} Dysphagia and first pass hepatic metabolism further decrease the bioavailability (F) of orally administered levodopa.² These features cumulatively contribute to fluctuating plasma levodopa levels and result in intermittent dopamine delivery to the CNS. When further combined with loss of the tonic buffer, this creates pulsatile stimulation and desensitization of the postsynaptic dopamine receptors, culminating in motor complications (Figure 1).³

Indeed, the ELLDOPA trial showed that high doses of levodopa/DDC inhibitors (DDCI) are a factor in the development of motor complications, which can emerge as early as 5–6 months after treatment with ≥ 600 mg levodopa per day.^{13,14} As pulsatile stimulation negatively impacts striatal function, the idea of achieving more continuous drug delivery (CDD) in order to better approximate continuous dopamine stimulation (CDS) has become the focus of therapies addressing motor complications (Box 1).

In this review we will discuss strategies which can be employed to manage motor complications, including those in the pipeline, and assess what unmet needs still exist in managing this complex and multifactorial phenomenon. While some non-pharmacologic approaches may be helpful, attempts to achieve CDS/CDD are reflected in efforts to increase plasma half-life of levodopa by developing longer acting formulations and preventing enzymatic degradation. There has also been interest in developing alternate administration routes which bypass gastrointestinal absorption and first-pass

Table 1 Types of off Episodes

Type	Definition
Early morning OFF	Symptoms are experienced upon awakening, prior to the first dose of medication (“morning bradykinesia / akinesia”)
Delayed On	Increased latency for the therapeutic dose of levodopa to alleviate symptoms; usually occurs with the first dose of the day or after eating
Wearing OFF / End-of-dose deterioration	Gradual recurrence of symptoms as patient gets closer to the end of a dosing interval, prior to the next dose
Sudden OFF	Random and unpredictable switch from “On” state to “Off”
Dose failure	Failure of therapeutic dose of levodopa to achieve any motor control

Note: Data from these studies.^{6,125,128}

Box I Strategies for Management of off Episodes

- Non-pharmacologic modifications
 - Dietary changes
 - Assess for underlying HP
- Dose fractionation
- Adjunctive use of longer acting dopamine receptor agonists
- Continuous Dopamine Stimulation (CDS)
 - Prolong levodopa plasma half-life (inhibition of enzymatic degradation)
 - Continuous drug delivery (infusions)
- Adjunctive use of non-dopaminergic medications
- Rescue therapies
- Device Aided therapies

Abbreviations: HP, *Helicobacter pylori*; SIBO, small intestine bacterial overgrowth.

metabolism. Device aided therapies such as levodopa/carbidopa intestinal gel (LCIG) and continuous subcutaneous apomorphine infusion (CSAI) are also of particular interest as they allow for more steady plasma concentrations of medication. Lastly, surgical approaches and non-dopaminergic agents can also be employed.

Non-Pharmacologic Considerations

Absorption of levodopa is dependent on the functional integrity of the digestive tract that will allow passage through to the small intestine. Constipation due to autonomic dysfunction is highly prevalent in PD, often preceding clinical Parkinsonian symptoms by years. This causes delayed gastric emptying which disrupts levodopa absorption and can lead to a delay in medication effect (delayed ON), no effect of medication (no ON) or unpredictable MF.¹⁵ *Helicobacter pylori* (HP) infection has been repeatedly shown to have higher prevalence in PD patients compared to the general population and has been thought to contribute to worse motor symptom control.¹⁶ Hashim et al found that at 12 weeks following successful eradication of HP infection in PD patients with motor fluctuations, there was a significant decrease in time to onset of medication effect, as well as overall improvement in daily ON time.¹⁷ More recently, Lolekha et al prospectively evaluated the therapeutic effects of HP eradication on MF. In their cohort of 40 Thai patients with MF, 22 were HP positive and 18 HP negative. At baseline, HP positive patients had longer daily OFF time (4.6 hours vs 3.9 hours). Of the 22 patients who received treatment with triple therapy, 17 had full eradication and they showed a significant decrease in daily OFF time by 0.7 ± 1.3 hours and increase in daily ON time by 0.9 ± 1.2 hours (in contrast to 0.6 hour increase in OFF time and a 1.6 hour decrease in ON time in those who had eradication failure). In total, they found the HP eradication effect to be -1.3 hours of OFF time.¹⁸ Small intestine bacterial overgrowth (SIBO) is relatively common in patients with PD, with prevalence in about half of patients with PD compared to about one-quarter of the general population.¹⁹ It has been postulated that *Enterococcus* species which overpopulate the small intestine in patients with SIBO are responsible for metabolism of levodopa, leading to lower absorption.¹⁹ Fasano et al were one of the first to establish an association between SIBO and MF (particularly delayed ON and no ON) in a small, unblinded, non-randomized trial.^{19,20} Additionally, they found that eradication of SIBO with antibiotic treatment had a meaningful clinical effect in improving MF.^{19,21} Aside from HP and SIBO, there has been growing interest in the gut microbiome of PD patients with ongoing investigation into the role of the microbial environment on levodopa resistance.¹⁵

Absorption of levodopa is facilitated via large neutral amino acid (LNAA) transporters which are also needed for absorption of dietary proteins, and it is often recommended that patients avoid taking their medications in close proximity to protein-rich meals. The protein redistribution diet (PRD) is characterized by limitation of protein intake early in the day, with compensation during dinnertime.²² It is important to note that within PRD, patients still consume the recommended daily intake of protein. Compared with those on a high protein diet, patients on PRD had lower motor disability but more dyskinesias, potentially indicating improved levodopa response.²² In a crossover study comparing PRD (15% daily protein before supper) with normal diet (60% of daily protein before supper), patients were randomized to 2 months of normal diet followed by 2 months of PRD. Results showed significantly improved total daily OFF time

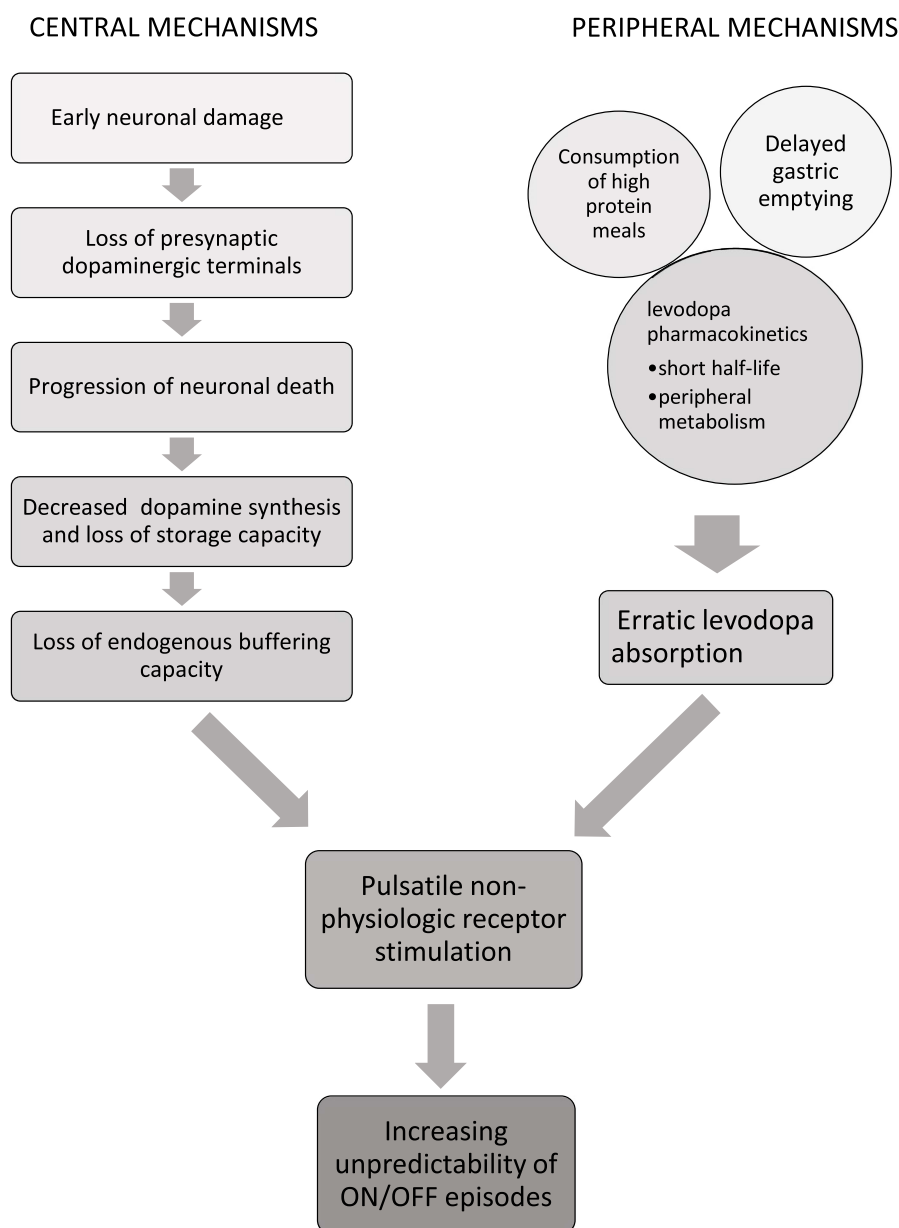


Figure 1 Pathogenesis of motor fluctuations in Parkinson's disease. Both central and peripheral mechanisms contribute to the eventual development of ON/OFF fluctuations by leading to pulsatile non-physiologic dopamine receptor stimulation in the basal ganglia. Successful pharmacologic management of OFF episodes attempts to achieve more continuous drug delivery in order to mitigate the combined effects of these mechanisms.

with PRD (164 minutes) vs normal diet (271 minutes) and significantly improved daily ON time with PRD (852 minutes) vs normal diet (738 minutes).²² While these results are promising, it may be unfeasible for some patients to alter their diets in such a manner. By contrast, low protein diets put patients at risk of weight loss and are not recommended.

Dietary fiber has shown some positive results in regards to optimizing levodopa therapy. Astarola et al found that in those patients with constipation, a diet rich in soluble fiber (28g fiber/day) for 3 months significantly reduced constipation, reduced the levodopa metabolite 3-OMD, and significantly increased levodopa plasma concentration.²² Another trial assessing the water soluble fiber *Plantago ovata* husk found no significant effect on the pharmacokinetics of levodopa, but did find a lower number of levodopa concentration peaks, potentially suggesting more smoother plasma profile that may be able to avoid peaks and troughs that contribute to MF.²²

Strategies to Consider Early After Onset of Motor Fluctuations

Alteration of the levodopa dosing regimen is one of the earliest approaches to consider once MF arise. Doses can either be increased to prolong ON time, or broken down into smaller doses (“fractionating”) which are more frequently administered, to reduce OFF time.²³ In 2021, the US Food and Drug Administration (FDA) approved the first functionally scored carbidopa-levodopa tablet that has deep grooves and a drug-free bottom layer that allows for more precise fractionation.²⁴ While dose fractionation may temporarily improve symptoms, it does not prolong levodopa half-life in the plasma and so repeated medication adjustments are often required, which can further contribute to erratic pharmacokinetics.²³ Increasing frequency of doses may also indirectly worsen motor symptoms by reducing compliance in some patients.^{23,25} Other limitations include risk of suboptimal control with smaller doses or conversely increased risk of dyskinesia with higher doses.²⁶ Flexilev® (Sensidose Pharmaceuticals, Uppsala, Sweden) is a rapidly soluble microtablet formulation that has been approved in Sweden since 2014.²⁷ It is unique in that it is delivered via an automatic dose dispenser, My Flexible Individual Dosing (MyFID™, Sensidose Pharmaceuticals, Uppsala, Sweden), allowing for more precise and individualized drug delivery.²⁷ Dosing regimens are programmed by a physician and the device is able to collect information on symptoms and medication adherence among other things. It is theorized that this may provide the most optimal mode of oral delivery for achieving CDD.²⁷

Dopamine agonists (DAs) are used in monotherapy as a levodopa-sparing option or as an adjunct to levodopa in order to reduce OFF time (Table 2). Levodopa has a short plasma half-life which increases to approximately 1.5 hours with the addition of a DDCl,¹⁰ while DAs have half-lives ranging from 5 to 8 hours depending on the medication.^{28–30} This allows for a more prolonged motor response due to prolonged stimulation of dopamine receptors, possibly reducing pulsatile stimulation for more tonic stimulation.³¹ Their use in early PD monotherapy has fallen out of favor due to their

Table 2 Effect of Dopamine Agonists When Used as Adjuncts to Levodopa

Drug Name (Generic)	Brand Name (Manufacturer)	Route of Administration / Dosing Frequency	Effect on Off Time	Effect on Levodopa Dose	Side Effects
Ropinirole Ropinirole ER	Requip® Requip XL® (GSK)	Both oral. TID for standard formulation, once daily for ER	Reduce (–1.5 hr/day for standard, –2.1 hr/day for ER)	Reduction possible (19.4% mean reduction with standard, 34% reduction with ER)	Nausea Dizziness Somnolence Fatigue Hallucinations Impulse control disorders
Pramipexole Pramipexole ER	Mirapex® Mirapex ER® (Boehringer Ingelheim)	Both oral. TID for standard formulation, once daily for ER	Reduce (–1.1hr change from baseline standard, –1hr for ER).	Reduction possible (27% reduction in levodopa dose for standard; Pramipexole ER increases levodopa Cmax by 40%)	Nausea Dizziness Somnolence Hallucinations Constipation
Rotigotine	Neupro® (UCB)	Transdermal patch, applied to skin every 24 hours	Reduce (PREFER study: –2.7hr 8mg, –2.1hr 12 mg; CLEOPATRA: study –2.5hr for up to 16mg)	Reduction is possible, amount not specified	Nausea Vomiting Somnolence Dizziness Application site reactions

Notes: All information adapted from FDA drug labels and trials mentioned within.

Abbreviations: TID, three times daily; Cmax, maximum plasma concentration.

less favorable side effect profile and overall lower clinical efficacy than levodopa.³² Pramipexole and ropinirole are DAs available in both conventional immediate release (IR) and once daily extended release (ER) formulations. Addition of either formulation of pramipexole or ropinirole to treatment with levodopa results in reduced OFF time (approximately 1–2 hours) and may allow reduction in levodopa dosing (Table 2). In a study comparing efficacy of pramipexole IR vs pramipexole ER as adjuncts to CD/LD, both were found to be comparable in safety and efficacy, with the advantage of once daily dosing with the ER formulation compared to three times daily dosing.³³ Similarly, the PREPARED study compared ropinirole IR and prolonged release (PR, same as ER) as adjuncts to CD/LD in patients experiencing MF and found results comparable, but generally favoring ropinirole PR in regards to reduction in OFF time.³⁴

Another DA, rotigotine, is the only transdermal patch currently authorized for use as monotherapy and as an adjunct to levodopa in PD. The patch is composed of a thin, matrix-type transdermal system composed of three layers and is available in six doses, which can be combined using the different patches.²⁹ Approximately 45% of rotigotine is released from the patch over 24 hours and this does not differ significantly depending on application sites. Once applied, there is an initial lag of 3 hours (range 1–8 hours) until it can be detected in the serum and time to maximal plasma concentration (Tmax) typically occurs between 15 and 18 hours post-dose, however there is no classic Cmax.²⁹ Steady-state plasma concentrations are achieved within 2–3 days.²⁹

Rotigotine is unique in that unlike the other non-ergoline DA's, pramipexole and ropinirole, it has activity at D3/D2/D1 receptors, such that it more closely resembles dopamine.³⁵ The efficacy of adjunctive rotigotine in addressing MF with reduction in OFF time is well established in placebo-controlled clinical trials, all of which show a 1–2 hour advantage over placebo in OFF time reduction.²⁹ Several NMS have also been shown to improve with rotigotine, however most lack high level class I evidence.³⁵ Nocturnal sleep disturbances have been shown to improve as a whole with rotigotine and this was demonstrated in 3 randomized controlled trials (RCTs), thus the Movement Disorder Society Evidence Based Medicine Committee (MDS-EBM) considers it “possibly useful” for the treatment of insomnia and “likely efficacious” in improving sleep.³⁵ The DOLORES study evaluated efficacy of rotigotine for MF-related pain and found insufficient evidence to define it, therefore MDS-EBM considers it as “investigational” for pain related to PD motor complications.³⁵ Given the close resemblance to levodopa and dopamine as well as the potential benefit in NMS, rotigotine offers potential as a strong therapeutic for advanced PD. Other formulations for rotigotine delivery are being investigated including rotigotine loaded nanoparticles for nose-to-brain delivery.³⁶ Alternatively, a ropinirole transdermal patch is being developed in Japan and was found to be non-inferior to ropinirole IR tablets, with maintained efficacy and plasma concentrations for up to 52 weeks.^{37,38}

Prolonging Levodopa Plasma Half-Life

Increasing levodopa plasma half-life can be accomplished by changing the levodopa formulation or inhibiting enzymes which metabolize it. Sustained-release carbidopa-levodopa (CD/LD SR) was the first longer-acting formulation introduced. It is supplied in tablets composed of a polymeric-based drug delivery system that releases the drug in a controlled manner.³⁹ However, clinical experience has revealed increased latency in time to “ON” and dose failures due to greater time to Tmax compared with IR (2 hours vs 0.5 hours for IR).³⁹ Additionally, there is high risk of dyskinesias as the (F) of CD/LD SR is approximately 70% that of IR, and so higher doses are needed, which may accumulate throughout the day and cause worsening of dyskinesias especially in the evening.⁴⁰ This was improved upon with carbidopa-levodopa extended release (CD/LD ER) capsules (Rytary®, Amneal Pharmaceuticals, Bridgewater, NJ; Numient®, Amneal Pharma Europe Ltd, Dublin, Ireland), a multiparticulate capsule of carbidopa-levodopa composed of both immediate release beads and sustained release beads combined in a specific ratio.⁴¹ The idea behind this was for rapid initial increase in plasma concentration with the IR beads, followed by a more sustained plasma concentration from the delayed release beads.^{3,41,42} As with CD/LD SR, the (F) of levodopa from CD/LD ER capsules relative to CD/LD IR is approximately 70%, but the Tmax is the same as CD/LD IR (1 hour).⁴² Following the peak, plasma concentrations are maintained for 4–5 hours before declining.⁴² The quick Tmax is intended to prevent delayed ON or dose failures that are seen with CD/LD SR, while also giving the advantage of a smoother plasma profile that eliminates the steep peaks and troughs which clinically manifest as MF. However, in the practical management of individuals with PD, it has been noted

that there was less reduction in OFF time and less reduction of dyskinesia when converting patients from CD/LD IR to CD/LD CR capsules when baseline greater amounts of troublesome dyskinesia are present.⁴³

In order to further improve the pharmacodynamic profile of ER levodopa, IPX203 has been developed (Amneal Pharmaceuticals, Bridgewater, NJ) with the goals of rapidly delivered therapeutic levodopa plasma concentrations, longer duration of such concentrations, minimal peak to trough fluctuations, and greater (F).⁴⁴ It is designed to be taken every 8 hours. Early data show that it has a rapid initial increase in plasma concentration comparable to CD/LD IR and CD/LD ER capsules, but it is able to sustain the plasma concentration longer (2.6 hours more than CD/LD IR, 0.9 hours more than CD/LD SR).⁴⁵ In regards to reduction in OFF time, recent data from a phase 3 RCT comparing IPX203 to CD/LD IR revealed a statistically significantly 0.53 hour/day increase in “good on time” as well as significantly less OFF time (−0.48 hour/day) from baseline to the end of study.⁴⁶ A post-hoc analysis of the data with a modified intent-to-treat population again showed results favoring IPX203 over optimized CD/LD IR regimen in regards to increased “good on time”.⁴⁷

Enzyme inhibition to prevent metabolism of levodopa has been in use since the early 1970s when carbidopa, a DDCI, was shown to increase the plasma half-life of levodopa from 50 minutes to approximately 1.5 hours.^{39,48} Benserazide is an alternative DDCI, used more commonly in Europe. Both DDCIs not only increase plasma half-life of levodopa, but also have the added benefit of increasing tolerability of levodopa by allowing lower dosages. COMT and MAO-B are two other major enzymes in the dopamine metabolic pathway. MAO-B inhibitors include selegiline, rasagiline, and safinamide (Table 3). Rasagiline is approved by the FDA for use as monotherapy in early PD while all three are approved for use as adjuncts to levodopa. Selegiline is unique in that it is metabolized into 1-methamphetamine and eventually 1-amphetamine, which may potentially play a role in improving alertness in the elderly, however the concentrations are negligible.⁴⁹ Rasagiline is more bioavailable than selegiline and may have neuroprotective qualities in vitro.^{49,50} It has been shown to improve morning OFF as well as wearing OFF, and post-hoc analyses have suggested a possible benefit in measures of depression, cognition and fatigue.⁵¹ Safinamide is unique in its mechanism of action by also working at voltage-gated sodium and N-type calcium channels to inhibit release of glutamate.⁵² This modulation of glutamate is a potentially important feature as excessive glutamate has been implicated in loss of nigral dopaminergic cells, in the development of levodopa-induced dyskinesias and in NMS such as pain.^{53–55} Furthermore, it has been suggested that it may also modulate metabolism of other neurotransmitters, which could provide benefit for NMS including depression and cognition.⁵² All three MAO-B inhibitors offer reduced OFF time with the associated possibility of reducing the total levodopa daily dose, and possibly improving NMS (Table 3).

Entacapone, tolcapone and opicapone are three approved COMT inhibitors (Table 3). Entacapone inhibits peripheral COMT and is available both on its own or in a formulation in combination with CD/LD at set dosage strengths (carbidopa-levodopa-entacapone, CLE, Stalevo[®], Orion Pharma, Espoo, Finland). Strong evidence to support the use of entacapone comes from four phase 3 RCTs which showed that the CLE improved pharmacokinetic profile translates clinically to greater efficacy in controlling motor symptoms compared to CD/LD alone, even with a 12% reduction in mean daily levodopa dose.^{23,56–60} Tolcapone is the most potent of the three COMT inhibitors as it acts both peripherally and centrally, however it is associated with potential for severe hepatic injury and carries a boxed warning from the FDA.⁶¹ Opicapone is the newest in the class and contains a pyridine N-oxide residue which gives it high affinity to COMT inhibition.⁶² Similar to entacapone, it does not cross the blood brain barrier, can only work to inhibit peripheral COMT and the effect on COMT inhibition is dose dependent.⁶² It has also been found to have the longest half-life, with COMT inhibition detected at 72-hours post-dose.⁶² This is in contrast to the 8 hour and 18 hour duration of effect found with entacapone and tolcapone, respectively.⁶² A network meta-analysis comparing the three found that tolcapone was the most efficacious but at the expense of a poor safety profile while entacapone was the least efficacious; opicapone had the best profile in regards to efficacy and safety.⁶³ It should be noted, however, that opicapone had the highest rate of dyskinesias and so may not be suitable for certain patients.⁶³ The addition of all three COMT inhibitors to treatment with carbidopa/levodopa can reduce OFF times from 1.2 to 3 hours, with a variable reduction in the levodopa dose possible (Table 3).

Table 3 Effect of Enzyme Inhibitors When Used as Adjuncts to Levodopa

COMT INHIBITORS					
Drug Name	Brand Name (Manufacturer)	Route of Administration/ Dosing Frequency	Effect on OFF Time	Effect on Levodopa Dose	Side Effects
Entacapone	Comtan ® (Orion Pharma)	Oral, with each levodopa dose up to 8 times/day	Reduce (–1.2 to –1.3hr reduction)	Reduce (average reduction about 25% in trials)	Dyskinesias Nausea Diarrhea Urine discoloration
Levodopa-Carbidopa-Entacapone	Stalevo ® (Orion Pharma)	Oral, up to 8 times daily with 200mg pill	Reduce (–1.2 to –1.3hr reduction)	Reduce	Dyskinesias Nausea Diarrhea Urine Discoloration
Tolcapone	Tasmar ® (Bausch, Valeant)	Oral, TID	Reduce (–1.4 to –2hr for 100mg; –1.6 to –3hr for 200mg)	Reduce (–32 to –166mg for 100mg dose; –21 to –207mg for 200mg dose)	BLACK BOX WARNING – severe hepatic injury Dyskinesias, nausea, sleep disorder, dystonia
Opicapone	Ongentys ® (Neurocrine Biosciences)	Oral, once daily	Reduce (–1.95 to –1.98hr)	Reduce (peak Cmax for levodopa increased 43–44% and AUC 62–94%)	Dyskinesia Constipation Hypotension Weight loss
MAO-B INHIBITORS					
Drug Name	Brand Name (Manufacturer)	Route of Administration/ Dosing Frequency	Effect on OFF Time	Effect on Levodopa Dose	Side Effects
Rasagiline	Azilect ® (Teva Neuroscience)	Oral, once daily	Reduce (PRESTO study: –0.49hr for 0.5mg and –0.91hr for 1mg) (LARGO study: –1.18hr for 1mg)	Minimal (LARGO trial: –24mg/day decrease in levodopa dose)	Orthostatic hypotension Dyskinesias Nausea Constipation
Selegiline Selegiline ODT	Eldepryl ® (Apotex) Zelapar ® (Valeant)	Oral, BID Oral disintegrating, once daily	Reduce (ODT reduced by –2.2hr)	Reduce (10–30% for oral; 24% for ODT)	Nausea Dizziness Syncope Confusion
Safinamide	Xadago ® (US WorldMeds)	Oral, once daily	Reduce (–1.3hr to –1.56hr)	No change	Dyskinesia Fall Nausea Insomnia

Notes: All information adapted from^{129–131}.

Abbreviations: COMT, catechol-O-methyltransferase; MAOB, monoamine oxidase B; ODT, orally disintegrating tablets; hr, hour.

Non-Dopaminergic Adjuncts

Limitations of dopaminergic replacement therapy have led to exploration of alternative drug targets. Medications such as amantadine, zonisamide and istradefylline may still have some indirect dopamine modulating activity, but their primary mechanisms are non-dopaminergic.

Amantadine

Amantadine is the only drug currently indicated for both reduction in OFF time and treatment of dyskinesias. This unique property is likely due to its effect on multiple neurotransmitter systems. Experimental models show that amantadine enhances dopamine transmission, and increases dopamine biosynthesis, turnover, uptake, and release in rodent models.⁶⁴ In human models, it is shown to increase striatal L-amino acid decarboxylase activity, which increases dopamine synthesis.⁶⁴ Antagonism at NMDA receptors is thought to account for its anti-dyskinetic effect.⁶⁴ It has also been seen to interact with serotonergic, noradrenergic, and cholinergic systems.⁶⁴ While it is available in both IR and ER formulations, only amantadine ER, Gocovri® (Adamas Pharmaceuticals, Emeryville, California) is indicated for treatment of dyskinesias and reduction in OFF time. Amantadine ER has a long half-life with a median Tmax around 12 hours, which means with bedtime dosing, patients wake up with higher plasma concentrations during the day when benefits are needed.⁶⁵ In a pivotal phase 3 trial, amantadine was shown to reduce OFF time by -1.2 hour/day.⁶⁶ The unique duality in improving both dyskinesias and OFF time is an important feature of amantadine ER, as one of the biggest obstacles in treatment of motor complications is the successful management of OFF time without worsening dyskinesias. Aside from motor benefits, amantadine ER was also found to provide statistically significant and clinically meaningful reductions in Movement Disorder Society-Sponsored Revision of the Unified Parkinson's Disease Rating Scale part II (MDS-UPDRS II) motor experiences of daily living.⁶⁷

Zonisamide

Zonisamide is an anticonvulsant drug with additional evidence for motor benefit in PD patients. It has been approved as an adjunct to levodopa in Japan since 2003.⁶⁸ The mechanism of action as an adjunct in PD is unknown, but it has both dopaminergic and non-dopaminergic mechanisms including reversible MAO-B inhibition and modulation of levodopa receptor expression and metabolism.⁶⁸⁻⁷² Zonisamide is taken once daily, therefore improving compliance. The recommended 25mg-50mg dose in PD is markedly lower than that used in epilepsy (200-400mg), which may be more tolerable. Otherwise, the most common side effects of zonisamide amongst PD patients include decreased appetite, weight loss, somnolence, apathy and dizziness.⁶⁸ Pivotal Phase 2 and 3 trials done in Japan revealed zonisamide 25mg and 50mg to be more effective than placebo in improving UPDRS III (motor) scores, while a post-hoc analysis found that zonisamide 50mg was superior to placebo in improving OFF time (zonisamide 50mg reduced mean daily OFF time by 0.98 hours compared to placebo).^{68,73-75} An important finding in the post-hoc analysis was that zonisamide 50mg may additionally benefit dyskinesias when compared to placebo, since it was associated with a decrease in disability of dyskinesia score (UPDRS part 4, item 33) from baseline to week 12 by 0.20 points without causing an increase in dyskinesia duration (UPDRS part 4, item 32).⁷⁵ In regards to NMS, there is conflicting evidence on zonisamide's role in development of ICD as earlier studies suggested some association between zonisamide and development of these behaviors, however a more recent open label trial found the opposite results. Furthermore, it is suggested that zonisamide may reduce cognitive decline more than the other PD therapies via potential neuroprotective effects such as protection against oxidative stress and potentially against progressive dopaminergic cell loss.⁷⁶

Istradefylline

Istradefylline is the first adenosine A2A receptor antagonist that is labeled for use in PD.⁷⁷ Adenosine is a neuromodulator that exhibits action via multiple receptor subtypes in response organ stress or tissue damage and in the CNS is involved in memory, mood, and motor function.⁷⁸ Adenosine A2A receptors are highly expressed in basal ganglia nuclei in close relation to dopamine D2 and D3 receptors such that activation of A2A receptors inhibits the indirect pathway.⁷⁸ While istradefylline has not been approved as monotherapy, it exhibits effectiveness in OFF time reduction as an adjunct to conventional PD therapy.⁷⁸ Four pivotal trials found an overall reduction in OFF time of approximately 1-2 hours, and the mean reduction in daily OFF time relative to placebo was -0.75hr (20mg group) and -0.82hr (40mg group). Open label studies also demonstrated enduring OFF time reduction at 52 weeks and overall positive safety profile.^{77,79,80} The most common side effects were dyskinesia, dizziness, constipation, decreased weight,

hallucinations, and insomnia.^{77,79} The overall lack of serious side effects associated with istradefylline has been attributed to its lack of direct dopaminergic activity.^{77,80}

In an indirect comparison of istradefylline to other adjunct therapies including dopamine agonists, COMT inhibitors, and MAO-B inhibitors, the mean reduction in OFF time relative to placebo from the 4 pivotal trials of istradefylline [−0.75 hr (20mg dose) and −0.82hr (40mg dose)] was compared to that of other agents. The mean reduction in OFF time relative to placebo was −1.54hr for dopamine agonists, −0.93hr for MAO-B inhibitors, and −0.83hr for COMT inhibitors.⁷⁷ Taking all this information into account, while istradefylline may be comparable in efficacy to MAO-B and COMT inhibitors, it is less efficacious than dopamine agonists in OFF time reduction.⁷⁷

Rescue Therapies

Parcopa

An orally disintegrating form of CD/LD (Parcopa[®], Schwarz Pharma, Mequon, WI) was FDA approved in 2004 and later by the EMA in 2011. This formulation uses RapiTab[™] technology (Schwarz Pharma, Monheim, Germany), which allows the tablet to disintegrate on the tongue within seconds, without the need for water.^{3,81} An early open label study comparing this orally disintegrating pill (ODT) to conventional CD/LD IR found no statistically significant differences in motor scores in both ON and OFF states between the two groups.^{82,83} A small pilot study further corroborated this, finding no statistically significant difference between ODT vs conventional CD/LD IR in onset to clinical benefit or overall motor performance, though it should be noted that all trends favored ODT.⁸⁴ This may be attributed to the fact that while, theoretically, absorption may be quicker due to quick oral disintegration, this formulation still relies on gastrointestinal absorption in the same way as CD/LD IR, while providing no benefit for sustained plasma levels as is seen with CD/LD CR, CD/LD ER, and CLE. While there was no significant difference between UPDRS motor scores in patients receiving ODT vs conventional oral therapy, Nausieda et al found a patient preference for ODT, citing accessibility of the medication to treat OFF times, reduced concerns about swallowing, its use for nighttime dosing and the fact that it is less noticeable thus less stigmatizing when taking the medication in front of others.⁸² CD/LD ODT most commonly addresses early morning OFF and delayed ON features of PD (Table 1).

Inbrija

Levodopa inhalation powder, Inbrija[®] (Acorda Therapeutics, New York, NY) is delivered via Arcus[®] technology (Acorda Therapeutics, New York, NY) to the pulmonary system. It consists of capsules which are broken, releasing the powder which is then orally inhaled by the patient. Single dose pharmacokinetics show a median Tmax of 30 minutes which clinically translates to a quick onset of action.⁸⁵ Clinical improvement is seen within 5–10 minutes, which makes it a faster rescue therapy than the alternative apomorphine subcutaneous injection. The dose-normalized Cmax of levodopa from Inbrija is approximately 50% of that from IR.⁸⁵ Placebo-controlled studies show superiority in reduction of UPDRS part III (motor scores) from pre-dose to 30 minutes post-dose, with benefit evident up to 60 minutes after inhalation.⁸⁶ While respiratory side effects are the most common (eg, mild cough), no significant differences were seen in spirometry measures between Inbrija and placebo at initiation of treatment, at 4 weeks, 12 weeks, and even at 12 months.^{87–89} Some patients may prefer administering medication via an inhaler compared to a more invasive injection. CD/LD inhalation powder may also be particularly useful in patients with difficulty swallowing pills and those with severely delayed gastric emptying due to its unique property in bypassing gastrointestinal absorption.

INP-107 (Impel Neuropharma, Seattle, WA) is an inhalation powder aimed for intranasal delivery via POD[®] devices (Impel Neuropharma, Seattle, WA). Similar to the already available CD/LD inhalation powder, it bypasses gastrointestinal absorption and allows for rapid relief of symptoms.⁹⁰ Unlike the oral inhalation route implicated with use of Inbrija, intranasal delivery is more natural and convenient as it avoids side effects such as dry mouth.⁹¹ INP107 is currently in development with phase 2 trials complete.⁹⁰

Apomorphine – Sublingual and Subcutaneous

Apomorphine is an alkaloid derived from morphine which has structural similarity to dopamine and functions as a DA but with key differences in its dopamine receptor affinity.⁹² Similar to rotigotine, apomorphine has a high affinity for D1 receptors, thus making it closer in action to dopamine.⁹² Apomorphine is available in the form of a sublingual film (Kynmobi[®], Sunovion Pharmaceuticals, Marlborough, MA), a subcutaneous injection (Apokyn[®], Supernus Pharmaceuticals, Rockville, MD; APO-Go[®], Britannia Pharmaceuticals, Reading, UK), and a subcutaneous infusion. The sublingual film and subcutaneous injection are indicated as rescue therapies; apomorphine subcutaneous infusion will be discussed separately. Both rescue formulations bypass first pass metabolism and are efficacious in reducing OFF time more rapidly than orally ingested pills. In a direct comparison of the two, apomorphine injection had a higher maximum plasma concentration (C_{max}), a shorter T_{max} but overall similar systemic exposure (AUC).⁹³ So while C_{max} was rapidly achieved (<1 hour) for both, the injection was about 15 minutes faster than the sublingual film.⁹³ While it may be important for some patients to achieve a quicker response, patient preference surveys have found them to generally prefer oral disintegrating formulations as opposed to injection or inhalation.⁹³ Regardless of formulation, a disadvantage of apomorphine is the severe nausea which requires additional treatment with trimethobenzamide or domperidone, neither of which medications are available in the US and so alternative treatments may be required.⁹⁴ Clinically, because of their short duration of onset, apomorphine sublingual and subcutaneous formulations can be used to address any of the OFF types that are commonly encountered in PD (Table 1). However, attempts to further improve the latency to ON with apomorphine have led to the development of AZ-009, an inhaled formulation that is rapidly systemically absorbed with a median T_{max} of 1–2 minutes, and with early evidence for clinical efficacy in successfully treating morning OFF amongst PD patients.^{95,96}

Infusion Therapies

Levodopa-Carbidopa Intestinal Gel (LCIG)

Levodopa-carbidopa intestinal gel (LCIG, trade name Duodopa[®] / Duopa[®], AbbVie LTD, Maidenhead, UK) is a stable gel suspension designed for continuous delivery of levodopa, developed in Sweden in the 1990s and approved in Europe in 2004 and US in 2015.⁹⁷ The levodopa containing gel is delivered straight to the proximal small intestine via a percutaneous endoscopic gastrostomy with jejunal (PEG-J) tube that is connected to a portable infusion pump, which is carried by the patient.⁹⁸ The gel is infused for 16 hours/day during waking hours, though some patients may use it for longer.⁹⁷

LCIG allows for a two-pronged approach in addressing OFF episodes in PD. Firstly, continuous infusion allows for a steady stream of levodopa circulating to the CNS that is a contrast from the peaks and troughs with repeated dosing with conventional oral therapy. Second, the PEG-J tube allows for direct delivery to the primary area of levodopa absorption, thereby bypassing gastric absorption which can be delayed and lead to erratic absorption in advanced PD.⁹⁹ A pivotal RCT comparing LCIG to CD/LD CR found lower mean plasma concentration variation, lower average change of variation of plasma concentration and lower mean levodopa exposure with LCIG as compared to CR.¹⁰⁰ These findings underlie the clinical effects of increased ON time (associated with reduced OFF time) and reduced dyskinesia. Other trials after this have confirmed reduction in time spent in the OFF state in patients receiving LCIG vs other conventional therapies including optimizing CD/LD IR and CD/LD ER.^{100,101} In a placebo controlled, double blinded study comparing LCIG with optimized medical therapy, LCIG provided a –4.04 hour/day decrease in OFF time vs –2.14 hour/day difference seen with optimized medical therapy.¹⁰¹ A qualitative review of long-term effects of LCIG found that in a majority of the studies reviewed, there was a statistically significant reduction in mean OFF time by at least 2 hours at a minimum of 12 months, with the longest follow-up being 120 months.⁹⁹

With LCIG, there is a unique set of adverse effects that is attributable to the invasive nature of the delivery system. The most frequent adverse effects are related to the surgical procedure including wound infection, tube dislocation, and abdominal pain.⁹⁹ Peripheral neuropathy as a result of levodopa-based therapy has been observed more frequently in patients on LCIG versus oral formulations.¹⁰² While the exact mechanism of the development of neuropathy is unclear, vitamin B deficiency and hyperhomocysteinemia have been cited, and thus vitamin B supplementation has been

employed to manage this.⁹⁹ Diphasic dyskinesias may develop in up to 15% of patients treated with LCIG and can be addressed by increasing the infusion time to 24 hours.¹⁰³

LCIG is considered one of the advanced options in the treatment of PD, alongside deep brain stimulation (DBS) and continuous apomorphine infusion (CSAI). While there have been trials that extensively compare levodopa infusion to conventional oral therapies, there are less head-to-head data thus far regarding surgery vs infusion options. A review comparing available data on LCIG and subthalamic nucleus (STN) DBS found that while the two were comparable in the motor control domain, LCIG offers some cognitive benefit.¹⁰⁴ This can be helpful when considering advanced therapeutics as those patients with dementia may potentially be better suited to LCIG vs DBS.¹⁰⁴ Other important aspects to consider include economic (LCIG is the more costly than DBS and CSAI in many places), patients' willingness to carry a harness with the pump, and other co-morbid conditions.¹⁰³

As a result of its design, LCIG is the closest to physiological delivery of levodopa to the CNS.³ Other pros of this include the ability to individualize treatment by adjusting flow rates and administering boluses, which can be more precisely targeted than by splitting conventional pills.⁹⁹ This is an effective treatment option for patients with severe motor complications in which OFF episodes are not treated by conventional oral therapy, however its costs and efficacy in comparison to other advanced therapeutics needs to be considered closely.¹⁰³

A newer device aided therapy with a lighter, more practical pump system is Levodopa-entacapone-carbidopa intestinal gel (LECIG) infusion (Lecigon®/Lecigimon®; Lobsor Pharmaceuticals AB, Uppsala, Sweden).¹⁰⁵ This was developed based on data showing that the addition of oral entacapone to LCIG allowed for a dose reduction in levodopa whilst maintaining clinical efficacy.¹⁰⁵ Senek et al conducted a randomized, open-label, cross-over study to measure systemic levodopa exposure with LECIG infusion using a 20% reduced dose vs patients' usual dose of LCIG.¹⁰⁶ Despite the reduced dosage with LECIG, systemic levodopa exposure did not differ between the two treatments, the dose-adjusted levodopa exposure was significantly higher during LECIG administration, and the treatment response scale (TRS) did not differ between the two.^{105,106} Patients on LECIG also had lower plasma concentrations of the levodopa metabolite 3-OMD, the formation of which is hypothesized to play a role in development of neuropathy.¹⁰⁶ It is therefore speculated that rates of neuropathy may be lower with LECIG than with LCIG.¹⁰⁶ All adverse effects were mild, including headache and nausea.¹⁰⁶ While these results are promising, larger comparative studies are needed to assess safety, efficacy and long-term outcomes.¹⁰⁵

Continuous Subcutaneous Apomorphine Infusion (CSAI)

CSAI is approved in Europe and under review by the FDA.¹⁰⁷ The infusion is delivered via a battery-driven, portable pump system that delivers a continuous dose of drug, although bolusing is an option.⁹² Similar to LCIG, the infusion is typically 12–16 hours during the waking day, but can be extended to 24 hours in certain cases.⁹² Its efficacy was established in the TOLEDO trial which found that from baseline to 12 weeks, the CSAI arm had an absolute difference of reduction in OFF time of –1.9 hours when compared to placebo and a 2 hour increase in ON time without troublesome dyskinesias.¹⁰⁸ CSAI offers a smooth continuous pharmacokinetic profile that is comparable to LCIG without the need for a surgical procedure. The most common adverse events are skin nodules at the injection site. A special consideration for both the apomorphine subcutaneous injection and apomorphine infusion is that a challenge test needs to be done in a supervised setting by a trained health-care professional to avoid the risk of hypotension.⁹⁴ A disadvantage in some patients may be the severe nausea caused by apomorphine (all forms), thus patients are routinely administered anti-emetics.¹⁰⁹

Under Investigation

Several levodopa subcutaneous infusions are under development to address persistent challenges in levodopa administration in order to better achieve CDD. These include ND0612 (NeuroDerm, Rehovot, Israel), ABBV-951 (Abbvie Inc, Chicago, IL), and Infudopa® (Dizlin Pharmaceuticals, Gothenburg, Sweden). These aim to address problems associated with levodopa solutions, including high instability at room temperature and limited solubility in water requiring large volumes.¹¹⁰ Successful subcutaneous solutions require high aqueous solubility and excellent chemical stability.¹¹¹ While ND0612 achieves a highly concentrated CD/LD solution, its alkaline pH causes a high rate of skin nodules – around 95% of people receiving ND0612.¹¹² ABBV-951 is a phosphate prodrug of CD/LD that appears more stable and is at a near

neutral pH, thus avoiding excessive skin irritation.¹¹³ Lastly, Dizlin pharmaceuticals is currently developing DZ102, which consists of a stock solution (CD/LD dissolved at a low pH) and a buffer that brings the solution to a physiologically favorable pH when mixed.¹¹¹ The stock solution is stable at room temperature for >3 months and the buffered mixture allows for better skin tolerability.¹¹¹

In a phase 2 place-controlled study, patients receiving levodopa standard of care (SoC) and ND0612 had a –2.1 hour/day reduction in OFF time vs a –1.4 hour/day reduction for patients receiving levodopa SoC and placebo.¹¹² A new drug application has been submitted to the FDA for ABBV-951 based on results of an open-label phase 3 study which demonstrated superiority of ABBV-951 vs placebo in reduction in off time (–2.75 hours/day ABBV-951 arm, –0.96 placebo arm) and increase in on time without troublesome dyskinesias (+2.72 hours/day ABBV-951 arm vs +0.97 hours/day in placebo arm).¹¹⁴ DIZ102 was found to be non-inferior to LCIG in regards to fluctuations of levodopa plasma levels, a property which is essential for infusions.¹¹¹ If approved, these investigational products may provide a less invasive and better tolerated alternative to infusion therapies, especially in older patients in whom DBS is not an option and a PEG-J procedure may be less desired.

Deep Brain Stimulation (DBS)

The current FDA-approved indication for DBS in PD includes patients who have had PD for at least 4 years, with at least 4 months of motor complications. DBS is the most frequently performed surgical procedure for PD, with two main target sites – subthalamic nucleus (STN) and globus pallidus interna (GPi).¹¹⁵ In a seminal trial, Deuschl et al demonstrated that STN stimulation was superior to medical management alone as evidenced by greater improvements from baseline to 6 months in UPDRS III (motor) and PDQ-39 (mobility, ADLs, emotional wellbeing, bodily discomfort).¹¹⁶ Numerous trials have supported this, such as the Veterans Affairs NIH trial (VA-NIH) that showed bilateral DBS (either STN or GPi) was superior to best medical therapy (BMT) as seen by the 4.6 hour increase in ON time without troublesome dyskinesia in the group treated with DBS (pooled for both targets) vs 0 hour/day improvement in the group assigned to continued BMT at 6 months, based on patient motor diaries.¹¹⁷ There was an associated 2.4 hour/day decrease in OFF time from baseline to 6 months in patients with DBS.¹¹⁷ Another study found a statistically significant increase in the percent of patients experiencing no OFF time during waking hours (from 3% at baseline to 29% at 1 year in the DBS + BMT group vs 2% increase in BMT alone group) as well as an increase in the percent of patients experiencing no dyskinesias during waking hours (37% increase in the DBS+BMT group vs –1% in the BMT alone group).¹¹⁸ It is important to note that while results from the DBS + BMT arm of the study were pooled for both STN and GPi, the majority of DBS patients in the study underwent bilateral STN electrode placement.¹¹⁸ In further randomized studies, one reporting effects from a multiple independent constant current-controlled device found bilateral STN stimulation provided a 3.74 hour increase in good ON time without troublesome dyskinesias at 3 months, while another one evaluating bilateral STN with constant-current stimulation found it to improve ON time without troublesome dyskinesias by 4.27 hours/day at 3 months.^{119,120} These differences likely relate to the baseline characteristics of the enrolled cohorts and methods of assessment. Regardless, multiple studies have confirmed the major benefits of DBS in PD, regardless of target, to be reduced OFF time, increased ON time, and reduced dyskinesias (Table 4).

One of the challenges in the device-aided treatment of PD with MF is choosing the best device (LCIG, CSAI, or DBS) for an individual patient. Currently, there are no trials directly comparing all three; however, recent meta-analyses have provided some guidance. A network meta-analysis done by Antonini et al demonstrated that DBS and LCIG had greater improvements in OFF time than CSAI with DBS resulting in greatest OFF time reduction.¹²¹ The change in OFF time was –1.45 hours/day for DBS and –1.35 hours/day for LCIG.¹²¹ However, LCIG had the highest ranking in QoL improvement.¹²¹ Marsilli et al also found DBS and LCIG to be superior to CSAI and BMT in improving motor function, however at the expense of higher lifetime costs (CSAI had the lowest lifetime costs and LCIG had the highest).¹²² An advantage of CSAI is that no surgical procedure is required, however it is not available in all countries. LCIG may be an option for patients with dementia or cognitive impairment in whom DBS is not an option.¹²²

Unmet Needs / Future Perspectives

Motor OFF times in PD are responsible for a substantial portion of the social and economic impact of PD on individuals, their caregivers and on health-care systems. One study assessing data from the “Financial and Social Impact of PD Survey”

Table 4 Comparison of DBS Target Sites

Authors (Year)	Study Design (Study Length)	Outcome Measures	Results
Follet et al (2010) ¹³²	Randomized, blinded (24 months)	Primary: change in motor function on UPDRS III with “stimulation on” and “no medication” Secondary: Self reported function, QoL, level of depression, serious adverse events	Both targets reached the primary outcome, however there was no statistically significant difference between the two No significant difference in self-reported function or QoL. Level of depression worsened after STN and improved after GPi. No significant intra-group differences between STN and GPi in regards to adverse events at 24 months. <u>Effect on motor complications:</u> <ul style="list-style-type: none"> change in ON time WITH troublesome dyskinesias from baseline to 6 months GPi -3.23hr, STN -2.1hr change in ON time WITHOUT troublesome dyskinesias from baseline to 6 months GPi +4.9hr, STN +4hr <u>LEDD reduction from baseline → 24 months (mg)</u> GPi: 1631 ± 545 → 1118 ± 562 (-513) STN: 1295 ± 585 → 887 ± 545 (-408)
Odekerken et al (2013) ¹³³	Randomized (12 months)	Primary endpoint: functional health as measured by ADLs as well as a composite score for cognitive, mood, and behavioral effects for up to 12 months after surgery Secondary endpoint: symptom scales, QoL, Adverse events, drug use	Both reached significance for the primary outcomes. There was no statistically significant difference between the two targets. Larger improvements in the “off drug” phase were seen in the STN group in regards to mean change in UPDRS III score, ALDS score, and medication reduction. All other secondary endpoints had no statistically significant difference between the two sites. <u>Effect on motor complications:</u> <ul style="list-style-type: none"> change in ON time WITH troublesome dyskinesias from baseline to 12 months GPi -2hr, STN -2.1hr change in ON time WITHOUT troublesome dyskinesias from baseline to 12 months GPi +3hr, STN +3.1hr change in OFF time per day from baseline to 12 months GPi -1.2hr, STN -1.7hr <u>LEDD reduction from baseline → 12 months:</u> GPi: 1331 → 1112 (-208) STN: 1254 → 708 (-546)

Notes: Trials are referenced with each author name.

Abbreviations: STN, subthalamic nucleus; GPi, globus pallidus interna; UPDRS III, Unified Parkinson Disease Rating Scale motor subscore; ADL, activities of daily living; QoL, quality of life; hr, hour.

(commissioned by MJ Fox Foundation and Parkinson’s Foundation) found that compared to those patients experiencing no OFF, patients experiencing OFF episodes were less likely to work full-time, more likely to have days with lower productivity, and more likely to miss work due to PD.¹²³ Additionally, they found that those providing care for patients with OFF episodes were more likely to stop working their other jobs due to PD and were more likely to have annual income losses when compared to caregivers of patients who were not experiencing OFF episodes.¹²³ Patients experiencing OFF were also more likely to have more neurologist consultations, more hospital and ICU admissions, and more ER visits as a result of their PD.¹²⁴

Several factors play into this, including the heterogeneity of these episodes between patients with varying frequencies and severities of each fluctuation even within the same patient. This is further complicated by misunderstanding of

symptoms by patients which results in underreporting and thus under-recognition by clinicians.¹²⁵ Some patients may also see these complications as an inevitable part of their disease and thus do not mention them.¹²⁵ Better tools for clinicians to assess motor complications are needed, since clinical scales that are commonly used in practice do not adequately assess these symptoms.¹²⁶ A survey done by Armstrong et al, in which PD patients and their caregivers were administered the Wearing off Questionnaire 19 (WOQ 19), found that while most fluctuating motor symptoms were assessed in some way with the WOQ 19, there were some motor, but numerous NMS which were reported but not included in WOQ 19, including fatigue and irritability.¹²⁵ Furthermore, a cross-sectional study focusing on the frequency and severity of NMS in patients with advanced PD who were experiencing MF showed that all patients experienced at least 2 NMS and all reported worsening of at least 2 of their NMS during their motor OFF episodes.¹²⁵ The Neuropsychiatric Fluctuations Scale (NFS) was recently developed but does not address sensory or autonomic fluctuations.¹²⁶ The Nonmotor Fluctuation Assessment Scale (NoMoFa) is a scale that was recently developed to address neuropsychiatric, autonomic, and sensory fluctuations and represents an important step in the ability to quantify NMF alongside MF and any improvements that may occur with intervention. Thus, both the recognition and treatment of NMF still represent a major unmet area in the management of PD.

Aside from scales to assess symptoms, physicians rely on home diaries and patient self-report to assess for MF, NMS and MMF, but it remains imperative that patients are educated on the nature of these symptoms, why they occur, the relation to their medications, and the availability of treatment options. Wearables and sensors may be better able to identify symptom dynamics (for both MF and NMF), and importantly may be able to provide objective data that can not only signal the need for treatment of OFF episodes, but also support the development of more targeted therapies or facilitate a holistic approach to managing PD symptomatology, beyond motor manifestations.

Effective management of OFF episodes in PD involves the manipulation of levodopa pharmacokinetics using non-pharmacologic approaches, adjunctive medications or alternate routes of delivery, the use of rescue therapies, or device-assisted therapies. While some of them are efficacious alone, a combination of these approaches is generally needed.¹²⁵ Efficacy is generally defined as reduction in absolute time spent in the OFF state (as defined by motor symptoms) without worsening of dyskinesia, and with evidence of improved QoL. None of these strategies, however, can completely ameliorate MF and dyskinesias. More notable effects can be seen with device aided therapies such as infusion therapies and DBS, which have been shown to deliver sustained reduction in OFF times due to their mechanisms, offer better control of dyskinesias, and have greater amounts of data exploring their effects on NMS and NMF. However, adequate guidelines for more timely intervention against MF and NMF with device-assisted therapies are needed.

Finally, due to the progressive nature of neurodegenerative disease, the risk of developing MF only increases as the disease advances and another major unmet need is for medications which can prevent disease progression. This approach will require better understanding of PD pathophysiology. While some therapies for monogenetic forms of PD are in development, it is not yet known whether these approaches will impact the evolution of MF and/or NMS/NMF. Alternative therapeutic targets will also need to be considered, as is currently being done with different neurotransmitter systems and identifying potential biomarkers.¹²⁷

Conclusion

While levodopa has been the cornerstone of PD treatment, adding or switching to different therapeutic strategies is required to mitigate the effects of MF. A working knowledge of the differences between these therapeutic strategies, their expected benefits, and risk profiles is required in order to make effective recommendations for management. Early on, it may be possible to adjust doses of medications patients are already taking, such as with dose fractionation or with non-medical approaches such as dietary changes to restrict eating within a certain time frame of taking medication. These benefits are likely short-lived and eventually other approaches are needed to try and achieve CDS/CDD in an effort to minimize OFF times.

As we have reviewed here, these approaches include the use of longer acting levodopa formulations, the use of enzyme inhibitors to prevent metabolic degradation of levodopa, and the use of non-dopaminergic medications, often in combination. The selection of specific strategies for an individual patient may be dictated by cost, patient preference, impact on associated levodopa dosing, risk of worsening dyskinesia, or other side effects. For example, a patient with mainly MF and no dyskinesia may benefit from a dosing schedule change, or adjunctive dopaminergic or non-

dopaminergic medications, while a patient with both MF and LID may benefit more from a rescue therapy, adjunctive non-dopaminergic medication or device assisted therapy. In fact, device aided therapies such as infusions and DBS are highly effective options that may provide more sustained improvement in MF, however more definitive guidelines regarding which patients to refer for these advanced therapies and when are needed. The resulting under-utilization of these options is further complicated by the fact that they are not universally available in all countries or may be subject to regulatory policies.

Emerging therapeutic targets for management of OFF episodes take advantage of the pitfalls of currently approved treatments, optimizing aspects such as absorption, drug half-life, and ease of administration. Newer agents such as IPX203 and new inhaled levodopa aim to improve upon pre-existing formulations, and will likely result in clinically relevant, albeit incremental, improvements in patient's motor control. Newer therapies also aim to increase ease of administration, eg, adapting once daily formulations, which may help improve compliance. Levodopa infusion therapies currently in development may advance our ability to achieve CDS/CDD, which could potentially be seen as "disease modifying" if physiologic stimulation can be achieved for a greater portion of the disease course.

While all these improvements are essential in the effort to improve the daily lives of PD patients, they still represent tools which clinicians must learn to use in combination, and strategically to mitigate risk of current and future functional impairments amongst patients with PD. Patients must receive adequate education about these treatments, their expected effects, and any adverse effects or consequences of their use. Clinician knowledge balanced with consideration of patient preferences as well as psychosocial and economic factors are therefore important requisites of true shared decision-making in the effective management of OFF periods in PD.

Disclosure

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