Key information for hospital pharmacists
Introduction

The number of people diagnosed with Parkinson’s in the UK is around 145,000\(^1\). There’s no cure yet for Parkinson’s, but medication can greatly improve symptoms.

Medication routines can be complex and because Parkinson’s is a progressive condition, people need regular, specialist review of their medication. This means that if necessary, it can be adjusted to maintain its benefits.

If an appropriate review of Parkinson’s medication is not completed, and a person’s routine is not properly maintained, their symptoms can become poorly controlled. This can mean their Parkinson’s symptoms get worse, even within a short period of time.

In England, people over 65 with Parkinson’s are three times more likely to have an unplanned admission to hospital than over 65s without the condition\(^2\).

Because of these factors it is likely that you will see a person with Parkinson’s in your hospital. It could be a planned admission but it could also be an emergency.

When a person with Parkinson’s is admitted to hospital, day-to-day life changes. They are more likely than the general population to experience disruption to their medication, including wrong doses or timing. They may also miss doses completely. These factors mean that a stay in hospital can have negative effects on someone’s condition and could make it worse\(^3\).
At Parkinson's UK, we want to work with pharmacists like you to highlight the importance of effective medicines management in easing these problems in hospitals. You can play a vital role in making sure that people with Parkinson's take the correct medication, on time. This will achieve good symptom control for people with the condition and they will have the best possible outcome from a hospital stay.

Understanding Parkinson’s for a better hospital stay

What is Parkinson’s?

Parkinson’s is a progressive neurological condition. It’s more common in older people, but younger people can also develop symptoms.

Parkinson’s develops when cells in the substantia nigra area of the brain stop working properly and are lost over time. It means these cells can no longer produce the chemical dopamine and the symptoms of Parkinson’s start to appear. Why this happens is unclear but it is thought to be a mix of genetic and environmental factors.

Motor symptoms

The main symptoms of Parkinson’s are slowness of movement (bradykinesia), rigidity and tremor. Most drug treatments aim to ease these motor symptoms.

Non-motor symptoms

People with Parkinson’s can experience a wide range of non-motor symptoms. This is because dopamine plays a role in neural pathways involved in cognition, arousal, motivation and reward. People with Parkinson’s often say non-motor symptoms can cause more distress in everyday life.

Common symptoms include pain, fatigue, mental health issues (such as depression and anxiety), sleep problems, and bladder and bowel problems. Constipation is particularly problematic in Parkinson’s as it can affect the absorption of medications, which in turn can worsen symptoms.

It is important to treat these symptoms appropriately. You should take care to use medications that are compatible with those for Parkinson’s, but which will not worsen motor symptoms. (See the section ‘Make sure other medications do not make Parkinson’s symptoms worse’ further on in this information).

Communication difficulties

It is important to remember that Parkinson’s can also lead to difficulties in communication. This can include

• quietening of the voice
• slurring of speech
• reduced facial expressions and body language.

This means it is important to make sure that your hospital environment supports your patient to communicate as well as they can.

Medication considerations when managing a person with Parkinson’s

Different types of medication

There are a wide range of Parkinson’s treatments. The most commonly used medicines are listed at parkinsons.org.uk/drug-treatments

They compensate for the loss of the dopamine-producing neurons.

• The levodopa group increases the levels of dopamine in the brain.
• The dopamine agonist group stimulates post-synaptic receptors that would normally be activated by dopamine.
• Other drugs block the action of enzymes and neurotransmitters that break down dopamine.

Levodopa

People using levodopa to manage their Parkinson’s may notice that over time, it becomes less effective. When symptoms are well controlled with levodopa, people with Parkinson’s are experiencing an ‘on’ period. As the body metabolises the drug, symptoms become poorly controlled and the person may have an ‘off’ period.
When this happens, movements become stiffer and the person may become completely unable to move. These motor fluctuations can be managed by taking medication on time.

As the condition progresses, ‘on’ and ‘off’ periods may increase so medication is needed more frequently. But increasing the amount of levodopa may also increase the chance of developing dyskinesia. These are involuntary, jerky movements that can affect any part of the body.

Because of these complexities, a specialist should oversee any significant changes in Parkinson’s medication routines.

When a person with Parkinson’s experiences issues linked to levodopa in your hospital, there are a number of strategies you can try. These include:

- using smaller doses more frequently to reduce the ‘peaks and troughs’ of levodopa and the incidence of dyskinesia. But any reduced dose should still achieve symptom control
- advising a person with Parkinson’s to wait for 30 – 60 minutes before eating a meal after taking their medication, as protein may inhibit levodopa absorption. A low protein snack such as crackers can be eaten with medication to avoid any nausea
- managing any underlying issues that might affect levodopa absorption, such as constipation or timings of any prescribed iron supplements. This is because in some cases chelates can form in the gastrointestinal tract. It is advised that 2–3 hours is left between iron and levodopa preparations

**Side effects**

Parkinson’s medications have potential side effects you should be aware of. These may include nausea, vomiting, drowsiness and low blood pressure, which can lead to dizziness and fainting.

**Impulsive and compulsive behaviour**

With some Parkinson’s drugs, particularly dopamine agonists and in a small number of cases, levodopa, some people experience problems with impulsive and compulsive behaviour. Examples of this behaviour may include gambling, compulsive spending, binge eating or hypersexuality. These behaviours can have a huge impact on people’s lives.

If, during an admission, impulsive and compulsive behaviour is observed or reported, it is important that medications are not suddenly stopped as this may make other symptoms worse. Instead, people with Parkinson’s and their family or carers should speak to their specialists about getting their medication reviewed as soon as possible.

Helpful information for patients about impulsive and compulsive behaviour can be found at parkinsons.org.uk/impulsivecompulsive

When a person with Parkinson’s is admitted to hospital there are other possible complications or ‘red flag’ signs that a specialist should review. These are discussed in Appendix 3.

Remember that people with Parkinson’s may not adhere to medication routines if the side effects of taking their medication outweigh the benefits. This means that asking your patient how their medication routine is helping them to function is key to understanding if it is improving their quality of life.

**Helping a person with Parkinson’s to take their medication**

Investigate whether a person with Parkinson’s is having difficulty taking their usual oral medication and manage it accordingly.
Below are some common issues. A person with Parkinson's may experience swallowing problems. There are different ways to manage this.

- Consider the best posture for an effective swallow. Sitting upright with the chin neutral can help.
- Review by a speech and language therapist for specialist advice.
- Ask patients if they are having any trouble taking their medication, as swallowing can make this more difficult. Thickened fluids or soft food can help.
- Use liquid or dispersible versions of drug preparations may help with swallowing.

**NEVER** crush or split modified release preparations, labelled CR, MR, XL or PR.

A patient may also experience nausea/vomiting or altered levels of consciousness, confusion, agitation or hallucinations. In these cases, it is important to:

- check for any previous history or underlying cause, including infection or dehydration, and treat it accordingly
- only use preparations of medication that do not worsen Parkinson's symptoms (See ‘Make sure other medications do not make Parkinson’s symptoms worse’ in the next section, below)

If a person is still not able to take their next prescribed oral dose, it may be necessary to consider administration via a naso-gastric, naso-jejunal or PEG tube, or via rotigotine patches

See Appendix 1.

Some people with Parkinson's may also be using non-oral medications or have had surgery to help them control their symptoms. See Appendix 2.

**Make sure other medications do not make Parkinson’s symptoms worse**

People may speak to you about problems that may or may not be associated with their Parkinson’s. While managing these effectively is clearly a priority, it is also important to make sure that medication given to treat these problems does not make their Parkinson’s worse.

**Treating nausea and vomiting**

Usually, Oral/PR domperidone (Motilium) is the antiemetic used to prevent and treat nausea and vomiting for people with Parkinson’s.

A European review in 2014 found a small increased risk of serious cardiac side effects with domperidone and advised that it is contraindicated in people with known cardiac conditions. But the advice also highlighted the need to consider the safety of domperidone with the clinical need to use it with other Parkinson's medication.
Drugs to avoid

Any drug that blocks dopamine receptors could make the symptoms of Parkinson’s worse or even mimic Parkinson’s symptoms without the condition being present.

When introducing any new medications for patients with Parkinson’s, it’s important that you think through the mechanisms of action.

Some drugs should be used with caution and the person with Parkinson’s should be closely monitored in case their symptoms get worse. In other cases, some drugs should be avoided.

For nausea and vomiting

avoid:

• metoclopramide (Maxalon)
• prochlorperazine (Stemetil)

If an injectable or alternate antiemetic is required, cyclizine (Valoid) may be considered post-operatively. Ondansetron can also be used although a common side effect is constipation. In addition, it is contraindicated if the person is using apomorphine because of the risk of hypotension.

For hallucinations/confusion

avoid:

• chlorpromazine (Largactil)
• fluphenazine (Modecate)
• perphenazine (Fentazin)
• trifluoperazine (Stelazine)
• flupenthixol (Fluanxol/Depixol)
• haloperidol (Serenace/Haldol)

Refer to a specialist for review and management.

For coughs and colds

avoid:

• preparations containing sympathomimetics (such as pseudoephedrine and ephedrine) with MAO-B inhibitors

Vigilance is particularly required with the use of:

• antihistamines, especially cinnarizine (Stugeron/Arlevert), which if used long-term, can mimic Parkinson’s symptoms
• antidepressants
• antipsychotics
• antihypertensives, such as diltiazem (Adizem/Angitil/Calcicard/Dilcardia/Dilzem/Solzem/Tildiem/Viazem/Zemtard) and other calcium channel blockers. Note that these should also be monitored by the GP clinical system.

More information about potential interactions with Parkinson’s medications can be found in Appendix 1 of the British National Formulary. The NICE Clinical Knowledge Summaries service also has information about which drugs to avoid with Parkinson’s.
Support the maintenance of prescribed medication routines

Parkinson’s affects everyone differently in terms of how symptoms present and the speed and nature of its progression. Because of this, medication routines may vary significantly between different people with Parkinson’s in terms of preparation, dosage and timing.

Routines will also need adjusting as the person’s Parkinson’s symptoms change over time and the condition progresses. Therefore it is important that people with Parkinson’s are regularly reviewed by a specialist to ensure they are getting the right treatment for them. This specialist may be a consultant neurologist, geriatrician, Parkinson’s nurse or specialist pharmacist.

It often takes time for someone with Parkinson’s to establish a medication routine that works for them and it is vital this is maintained so symptoms are controlled effectively. But this can be difficult in hospitals because:

• information about medication routines might not be easily accessible (particularly with an emergency admission or if the person is transferred between departments)
• the medications that a person with Parkinson’s is taking might not be easily available
• the medication routines may not fit easily with hospital procedures, such as timings of traditional drug rounds or theatre schedules for surgery.

Disruption to someone’s medication routine can have significant implications for the person with Parkinson’s and the level of support they require from hospital ward staff.

In extreme cases, missed doses may lead to the potentially fatal neuroleptic-like malignant syndrome.

Because specific Parkinson’s medication routines can be complex, it can be a challenge to maintain the right preparation, dosage and timing. Staff are also often required to give more assistance to a person with Parkinson’s.

However, there is a lot that you can do as a hospital pharmacist to make sure that routines are maintained to help people with Parkinson’s stay in control of their condition while they are in hospital.

What can I do to make sure people with Parkinson’s get the most out of a hospital stay?

Make sure people get their medication on time

The key issue with Parkinson’s medication is timing. Medication needs to be taken at specific times so that symptoms are controlled consistently without deterioration.

People with Parkinson’s tell us that when they are able to get their medication on time in hospital it helps them stay in control of their own condition and speeds up the time it takes them to recover.

Not getting medication on time can mean the difference between someone being able to function independently and someone becoming reliant on others for simple everyday activities such as walking and eating.

Get it on time

The Parkinson’s UK Get It On Time campaign raises awareness of the significance of drug timings in controlling the condition for people in hospitals and care homes. You can help us to get the message out to staff in hospitals that people with Parkinson’s need their medication on time – every time.

Find out more about the campaign and order our booklet and other materials at parkinsons.org.uk/getitontime
Parkinson’s UK has other information that might help you explain the importance of medication timing to other staff including:

- Caring for your patient with Parkinson’s – our booklet for ward staff
- Medicine optimisation for patients with Parkinson’s – our online video/DVD for ward staff

These are available to view and order from parkinsons.org.uk/professionals/resources

The Parkinson’s optimal dose calculator can also help you care for someone with Parkinson’s in an emergency setting. Find out more at parkinsons.org.uk/medicationcalculator

Help people with Parkinson’s to take control of their condition

- People with Parkinson’s are encouraged to bring a small supply of their medication in its original packaging to hospital with them so that it can be used during an admission, however this may not be available if they are admitted in an emergency.
- Assess patients to see if they are able to self-administer medication and support them to do so if they can. A person’s capability may change during an admission, such as immediately after an operation, so it is important that information about their self-administration status is accurately updated on all documentation and communicated to all relevant staff. However, the aim should always be to encourage people with Parkinson’s to be as independent with their medication as possible.

Prepare Parkinson’s medications

- Liaise with medical and nursing colleagues to make sure they have a comprehensive list of a person’s medications including information about preparations, dosages and timings. This can help with accurate prescribing and administration. People with Parkinson’s and their carers are usually aware of how important their medication is in managing their condition and may have this information with them. Make sure you ask them about this and check any formal documentation that they have.
- Medicine reconciliation should be a priority. This involves making a list of all medications a person with Parkinson’s is taking, to make sure they can continue to take them in hospital. It is important that procedures can support this at any time, including out-of-hours.
- Make sure a wide range of Parkinson's medication is stocked or that you are aware of how and where to get supplies so that individual routines can be maintained.
- Parkinson’s medications must be on the trust’s critical medicines list so that supplies can be accessed at any time.
- Use branded preparations where prescribed and don’t substitute with generic versions. This is because the level of active ingredients may vary. Also, non-active components of the drug may affect how well active ingredients are absorbed. The differences between formulations are small and can be regulated, but they might be clinically significant in terms of symptom control. Always explain to the person or their carer why a different preparation has been dispensed to help reduce any anxiety or confusion. For example, if medicines are not available locally.
- Print out timings on the pharmacy label or add them to any blister packs issued to help make sure that ward staff can support people with Parkinson’s to get their medication on time.
Adjust and plan hospital activities

- Make sure that other hospital staff understand the importance of keeping to the prescribed medication routine and adjust activities accordingly. Ward staff will need to ensure that a person with Parkinson’s gets their medication on time – even if this does not fit in with the usual timings of drug rounds, or if this coincides with other ward activities, such as mealtimes.

- Surgery or clinical tests will need to be planned around the timings of someone’s medication routine. Ideally, someone should be put at the start of operating lists to optimise their medication. People with Parkinson’s can still take prescribed oral medication, with small amounts of clear fluids, up to two hours before elective surgery¹.

- Anaesthetists should consider regional anaesthesia rather than general anaesthetic so that a person’s usual medication routine can continue.

- As patients may transfer between areas of the hospital during their admission, it may be necessary for you to advise a range of staff in different wards or departments.

Support people to take control post discharge

- If there have been any changes to a person’s medication routine while they have been in hospital, make sure you inform the person with Parkinson’s and their carers of these changes, including why they were needed and if they should look out for any specific side effects.

- After a person is discharged it is very important that clear information about their medication is sent to the person’s GP to avoid medication errors.

- It is also recommend that discharge medicines are communicated (with patient consent) to the community pharmacist. People with Parkinson’s often have close links with their community pharmacy and this will ensure they receive the best support with their medication.

- Because of the complexity of Parkinson’s medication, a Medicines Use Review (MUR) could help make sure a person’s medication routine continues to work for them. A national target area for MURs in England is patients who have recently been discharged from hospital and had their medication routine changed¹¹. You can schedule a MUR with the person’s community pharmacist following their admission. This is an excellent way of strengthening your connections with community care.
Appendix 1

Optimal Parkinson’s medication guideline and conversion calculator

You can use the guideline and calculator below to help your patients maintain control of their Parkinson’s symptoms. This is important when they cannot take oral medications until support is available from local specialists.

There are two calculators. The first is for patients who can have an NG Tube and the second is for patients who cannot. You can access them online at parkinsons.org.uk/medicationcalculator

Appendix 2

Non-oral treatments for Parkinson’s

Apomorphine (APO–go)

Apomorphine is a dopamine agonist administrated via an intermittent sub-cutaneous injection or a continuous subcutaneous infusion via a pump – it is not morphine-based, is not an analgesic and is not a controlled drug.

Patients who are established on apomorphine need to be continued at the prescribed dose and frequency (injection) or rate (pump) – do not change the pump settings unless requested to do so. For further support, call the APO–go helpline on 0844 880 1327 or contact the relevant specialist, for example a person’s Parkinson’s nurse or specialist pharmacist.

Duodopa infusion

Duodopa is co-careldopa (levodopa and carbidopa) in gel form. It is delivered into the jejunum via a PEJ tube.

Patients who are established on a Duodopa routine need to be continued at the prescribed rate – providing gastric emptying is not delayed and the PEJ tube is unobstructed. If not, discontinue and commence on rotigotine patches.

Deep brain stimulation (DBS)

This involves stimulation of target sites within the brain through electrodes connected to a neurostimulator placed under the skin around the chest or stomach area. Target sites are in the thalamus, the globus pallidus or the subthalamic nucleus.

Patients who are established on DBS need to be maintained on the same routine. For further support, contact the neurological department that implanted the system. The patient should carry a patient ID card listing contact details and the model number of their DBS system.
Appendix 3

Complications with Parkinson's

• Delirium (acute confusion due to drugs or infection)
• Chest infection, especially aspiration pneumonia
• Urinary tract infections
• Postural hypotension and falls – check meds and BP lying/sitting then standing
• Neuroleptic-like malignant syndrome

Red flags that means a patient needs to be referred to a Parkinson's specialist:

• Fibrotic reactions with ergot-derived dopamine agonists. These include bromocriptine, pergolide and cabergoline where these have been prescribed. For example dyspnoea, persistent cough, chest pain, cardiac failure, abdominal pain or tenderness may occur and these patients should be having regular echocardiography and chest X-rays
• Signs of liver disorder with tolcapone, such as nausea, vomiting, fatigue, abdominal pain, dark urine or pruritus. These patients should be having regular liver function tests.
• Increased falling especially early in the condition
• Hallucinations/dementia/depression/cognitive decline, especially early in the condition

References
1. Parkinson's UK (2018) 'The incidence and prevalence of Parkinson's in the UK – Results from the Clinical Practice Research Datalink summary report'
More information and support for your patients

Our free confidential helpline can provide general support and information. Call 0808 800 0303 (calls are free from UK landlines and most mobile networks) or email hello@parkinsons.org.uk.

We run a peer support service if your patient would like to talk on the phone with someone affected by Parkinson’s who has faced similar issues to them. The service is free and confidential – our helpline can talk to your patient about being matched with a volunteer.

Our helpline can put your patients in touch with one of our Parkinson’s local advisers, who give one-to-one information and support to anyone affected by Parkinson’s. They can also provide links to local groups and services.

Our website parkinsons.org.uk has a lot of information about Parkinson’s and everyday life with the condition. There are details of local support teams and local group meetings at parkinsons.org.uk/localtoyou

Your patients can visit parkinsons.org.uk/forum to chat to other people with similar experiences on our online discussion forum.

Driving
Advise people with Parkinson’s of the need to inform relevant authorities of their condition at the time of diagnosis, such as the DVLA (DVA in Northern Ireland) and their car insurer.

For information on driving and Parkinson’s visit parkinsons.org.uk/driving

Thank you
Thank you very much to everyone who contributed to or reviewed this information sheet:

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The UK Parkinson’s Excellence Network is the driving force for improving Parkinson’s care, connecting and equipping professionals to provide the services people affected by the condition want to see.

This is your Network. Get involved at parkinsons.org.uk/excellencenetwork