

# Shared care guidelines

Drug	FIRST GENERATION ANTIPSYCHOTIC LONG-ACTING INJECTIONS (Depots) <b>Flupentixol decanoate, Haloperidol decanoate &amp; Zuclopenthixol decanoate</b>
Specialty	ALL SPECIALTIES ( <u>excluding</u> Children & Young People's Services)
Indication	SCHIZOPHRENIA and other psychoses
Overview	<p>There are currently three first generation antipsychotic (FGA) long-acting injections (LAIs) recommended within the <a href="#">TEWV Long-Acting Injectable (LAIs) Antipsychotics procedure</a> - Flupentixol decanoate, zuclopenthixol decanoate and haloperidol decanoate.</p> <p>FGA LAIs should be initiated by a specialist with expertise in psychotic disorders as part of a comprehensive treatment plan but prescribing, administration &amp; monitoring responsibility can transfer to GPs under these shared care guidelines.</p>
Specialist responsibilities	<p><b>Pre-treatment:</b> (see each individual product <a href="#">SPC</a>) for full details of contra-indications &amp; cautions)</p> <p>Assess suitability for treatment with a FGA LAI by reviewing the patient's medical history, completing a physical examination and completing the baseline monitoring as detailed in <a href="#">TEWV Psychotropic Monitoring Guide</a> - Antipsychotics. It should be noted that dose adjustment is required in patients 65 years or older.</p> <p><b>Initial prescription - dosage and administration:</b> (see <a href="#">BNF</a>, <a href="#">SPC</a>) and <a href="#">TEWV Long-Acting Injectable (LAIs) Antipsychotics procedure</a> for full details)</p> <p>A small test dose should be administered to assess tolerance, then dose and dose interval should be adjusted according to the patient's symptoms and response to treatment.</p> <p><b>Monitoring</b></p> <p>The efficacy and tolerability of antipsychotic medication should be established by the specialist team by use of objective and validated measures, prior to transfer and at each review. Physical health monitoring should be completed by the specialist team for the first 12 months, then at each review (at least annually)</p> <ul style="list-style-type: none"> <li>• <b>Side effects</b> – use LUNSERS or GASS to assess tolerability at each review</li> <li>• <b>Physical Health monitoring</b> – for the first 12 months of treatment, then at each review (at least annually); see physical parameters in <a href="#">TEWV Psychotropic Monitoring Guide</a> - Antipsychotics.</li> <li>• <b>Clinical response</b> – use an appropriate measure, e.g. PANSS (positive and negative syndrome scale), CGI (clinical global impressions) and GAF (global assessment of functioning), to assess response prior to transfer and at each review.</li> </ul> <p>Where tolerability or clinical response is not demonstrated, the LAI should be stopped. On-going clinical need and patient preference for a LAI should be reviewed at least annually.</p> <p><b>Transfer of prescribing / communication – see checklist</b></p> <p>Prescribing, administration and monitoring responsibility may be transferred to the patient's GP after 3 months or once the treatment has been stabilised, whichever is the longer. The request must be made using the attached form with a covering clinic letter and a copy of this guideline (with contact details added) – the following details should be clearly communicated at transfer, and after each subsequent review:</p> <ul style="list-style-type: none"> <li>• <b>Diagnosis</b></li> <li>• <b>Dose</b> of FGA LAI (must not exceed BNF max. – if HDAT then not suitable for transfer)</li> <li>• <b>Date</b> and <b>site</b> of last administration, and <b>date when next dose is due</b></li> <li>• <b>Action</b> to be taken by GP if the patient does not attend for their scheduled dose</li> <li>• Completed and required <b>monitoring</b></li> <li>• <b>Discontinued medication</b> for same diagnosis</li> <li>• <b>Date</b> of next specialist review</li> </ul> <p>The transfer request should be sent one month in advance of the patient needing their next dose. Acceptance should not be assumed until the GP responds positively using the attached form (scanned &amp; e-mailed to the specialist team).</p>

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Approved by	Drug & Therapeutics Committee	Date of Approval	28 <sup>th</sup> November 2024
Protocol Number	PHARM-0142 –v2	Date of Review	1 <sup>st</sup> December 2027

## Specialist responsibilities (continued)

## GP responsibilities

### Discharge

Patients prescribed a FGA LAI should not be discharged from secondary mental health services. In exceptional circumstances an individual agreement for discharge may be considered for a patient who expressly indicates that they do not want to be seen by secondary mental health services. However, discharge should only be considered if FGA LAI treatment is stable, and the patient is adherent to treatment and compliant with monitoring requirements. Discharge arrangements should involve a proper discussion with the GP and the rationale for discharge must be clearly documented.

### Transfer of prescribing / communication:

Notify specialist immediately (within 2 weeks) if transfer of prescribing and monitoring responsibility is accepted or not so that alternative arrangements can be put in place if necessary. Contact specialist if communication of prescribing, administration & monitoring requirements is not clear.

### Maintenance (repeat) prescription:

Prescribe the FGA LAI in accordance with specialist advice received on transfer and following reviews. The maximum dose of each FGA LAI that the GP would be expected to prescribe is as follows:

**Zuclopenthixol decanoate:** 600 mg weekly (max single dose: 600mg)

**Flupentixol decanoate:** 400 mg weekly (max single dose: 400mg)

**Haloperidol decanoate:** 300 mg every 4 weeks (max single dose: 300mg)

Any dose above these is classed as "high dose antipsychotic treatment" and is not suitable for shared care (**RED** classification)

### Administration:

See [SPC](#) and see Appendix C&D of [TEWV Long-Acting Injectable \(LAIs\) Antipsychotics procedure](#) for detailed information regarding sites of administration, volumes of administration and action to take in response to missed or delayed doses.

**Monitoring** – annually (additionally as clinically indicated) – see [TEWV Psychotropic Monitoring Guide](#) – Antipsychotics for details:

- Weight (+BMI and waist circumference where possible)
- Urea & electrolytes, including creatinine
- eGFR
- Lipid profile (total cholesterol, HDL-cholesterol, total/HDL-cholesterol ratio, triglycerides – fasting sample)
- Full blood count
- HbA1c
- Blood pressure (sitting/lying and standing) and pulse
- ECG – only if clinically indicated or patient is "high risk"

Any physical health concerns and/or recent monitoring tests should be communicated to the specialist prior to the next annual review.

### Referral:

Refer back to the specialist should any of the following occur:

- Significant adverse reaction or intolerable side effects
- Lack of efficacy / patient's condition deteriorates
- Development of co-morbidities / necessity to prescribe interacting drugs
- Pregnancy
- Failure to attend for administration of dose within permitted timeframe

## Adverse events

See [SPC](#) and [BNF](#) for full details of known adverse effects

Common side effects include agitation, constipation, dizziness, drowsiness, dry mouth, arrhythmias and erectile dysfunction. Please note that side effects may persist after stopping a LAI FGA until the drug has cleared from its depot site. Report any suspected adverse events to MHRA via the [Yellow Card scheme](#)

## Specialist contact details

(to be added by specialist prescriber when transferring prescribing)

Name:

Base:

Telephone no:

E-mail address:

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<b>AMBER ▲</b>	<b>TRANSFERRING PRESCRIBING OF: ANTIPSYCHOTIC LONG ACTING / DEPOT INJECTIONS</b>		
<b>GP details:</b>			
<b>Patient details</b> (name/address/DOB/NHS number):			
<b>Diagnosis:</b> Specify clinical rationale if first line option or standard formulation not prescribed			
<b>Checklist for transfer:</b> <ul style="list-style-type: none"> <li><input type="checkbox"/> The patient has completed at least 3 months of treatment and is suitable for repeat prescription of the antipsychotic depot/LAI</li> <li><input type="checkbox"/> The medication and patient's mental health are stable (i.e. the patient has completed their response to medication and there are no recognised problems with attendance or significant acute risks of harm to self or to others).</li> <li><input type="checkbox"/> A minimum of one month's notice is being provided to the GP to ensure adequate time to add the prescription to the GP system</li> <li><input type="checkbox"/> The patient &amp; medication meets all of the criteria defined within the shared care protocol</li> <li><input type="checkbox"/> A clear and a copy of the shared care protocol has been sent to the GP</li> <li><input type="checkbox"/> Arrangements have been made to continue prescribing until the GP agrees to shared care being established for this patient</li> <li><input type="checkbox"/> Arrangements have been made for the necessary secondary care responsibilities to be carried out (as defined in the protocol)</li> <li><input type="checkbox"/> There has been consideration of STOMP (if applicable)</li> </ul>			
<b>Medication details:</b> (generic & brand name, dose and dose interval):			
<b>Discontinued medication</b> (list any medicines discontinued when this AMBER treatment initiated):			
<b>Last Administration</b> (details of date and site of administration and date next dose due):			
<b>Monitoring results:</b>			
<b>Secondary care review frequency:</b>			

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**Actions requested of GP:**

**Please continue to issue prescriptions, administer and monitor the FGA LAI detailed above until advised**

The treatment has been explained to the patient and they understand they should contact you for future prescriptions.

You will be informed of any changes to treatment, if you are not required to issue prescriptions or if treatment is to be discontinued.

Please contact the prescriber on the number below if there is any change in the patient's condition, if the patient is late or fails to attend for administration of doses, or you require advice.

Secondary care contacts:	Contact details (address/telephone no):
Care coordinator (name):	
Consultant (name):	
Prescriber (name):	
<b>Signature &amp; date:</b>	

**Acceptance of prescribing responsibility by GP:**

<b>Patient's name:</b>	<b>NHS Number:</b>
<b>Address:</b>	
<b>Medication:</b>	
I confirm receipt of prescribing transfer information for the above patient and accept prescribing responsibility	
<b>GP's name:</b> <i>(Please print name in BLOCK CAPITALS)</i>	
<b>Signature/ Practice Stamp:</b>	
<b>Date:</b>	
<b>Please scan &amp; e-mail back to (e-mail address):</b>	
<b>or return as soon as possible to (postal address):</b>	

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