



Lithium for patients within adult services

Shared Care Protocol

Specialist responsibilities

- Diagnose the patient; ensure that this diagnosis is included in section 2 and communicated to primary care.
- Discuss the benefits and risks of the treatment with the patient and provide the appropriate counselling (see section 11) to enable the patient to reach an informed decision.
- Provide verbal and written patient information including a completed lithium booklet which includes the record book and alert card.
- Inform patients of the signs of toxicity
- Inform patients of child-bearing potential, of the possible lithium teratogenic effects
- Provide advice on the need for contraception to patients on initiation of lithium, and at each review.
- Give advice to primary care on continuing treatment if a woman becomes or wishes to become pregnant.
- Assess for contraindications and cautions (see section 4) and consider potential drug interactions jointly with Primary Care (see section 7).
- Conduct required baseline investigations and initial monitoring (see section 8).
- Initiate and optimise treatment as outlined in section 5.
- Prescribe the maintenance treatment for at least 4 weeks and until optimised.
- Determine the target plasma lithium range for each patient and advise the primary care prescriber accordingly
- Once treatment is optimised, specialist should contact the patient's GP in writing to request shared care, providing details of the diagnosis, current and ongoing dose, any relevant test results and when the next monitoring is required. Request that lithium is prescribed by brand and that blood test results are recorded in the lithium record book before issuing a repeat prescription.
- Ask GPs to carry out maintenance monitoring as in section 9.
- Prescribe sufficient medication to enable transfer to primary care.
- Conduct the required monitoring in section 8.
- Provide advice to primary care on any switches between brands or formulations of lithium
- Provide advice to primary care on the management of adverse effects if required.
- Review patient at least every 12 months to assess their mental health, effectiveness of treatment and the ongoing need for lithium
- Provide a clear plan for lithium treatment to primary care, including anticipated duration of treatment and advise if treatment should be discontinued.
- Document all communication with primary care in the patient's electronic health record.
- When a patient is considered to have been stable mentally and functionally for a suitable period of time, liaise
 with the GP about the possibility of fully discharging the patient back to the care of the GP. The GP must be
 able to access a fast-track referral back to a secondary care specialist if needed at any point. The specialist will
 be available for queries could be directed to an AMHT pharmacist if appropriate.
- Alternative to above When a patient is considered to have been stable mentally and functionally for a
 suitable period of time, liaise with the GP about the possibility of fully discharging the patient back to the care
 of the GP. The specialist will remain available for advice via email and telephone for the duration of the
 patient's treatment



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Primary care responsibilities

- Respond to the request from the specialist for shared care in writing within 14 days.
- If accepted, prescribe ongoing treatment as detailed in the specialist's request and as per section 5 taking into any account potential drug interactions in section 7.
- Always prescribe lithium by brand and form. Discuss any changes in the brand or formulation with the specialist.
- Adjust the dose of lithium prescribed as advised by the specialist.
- Conduct the required monitoring as outlined in section 9.
- Assess for possible interactions with lithium when starting new medicines (see section 7).
- Manage any adverse effects as detailed in section 10 and discuss with specialist team when required.
- Be familiar with the symptoms of lithium toxicity and the common causes
- Discuss other adverse effects with the specialist team as clinically appropriate (see section 10).
- Contact the specialist team immediately if the patient becomes or plans to become pregnant.
- Be aware that abrupt discontinuation of lithium increases the risk of relapse and discontinuation should be gradual over a period of several weeks under specialist advice
- Stop treatment as advised by the specialist.

Patient and/or carer responsibilities

- Agree to treatment and monitoring after making an informed decision
- Agree to being under the shared care of the GP and specialist
- Take lithium as prescribed and not stop taking it without speaking to their primary care prescriber or specialist.
- Be familiar with their brand of lithium and check that they have received the correct one.
- Take doses of lithium at the same time each day and if a dose is missed, take the next scheduled dose as usual
- Attend regularly for monitoring and review appointments with primary care and specialist. Request that results are recorded in lithium booklet and bring this to each appointment.
- Be familiar with symptoms of toxicity and the common causes.
- Maintain adequate fluid intake, particularly in hot weather or during periods of increased activity
- Report adverse effects to their primary care prescriber. Seek immediate medical attention if they develop any symptoms as detailed in section 11.
- Inform healthcare professionals that lithium is being taken when seeking medical or pharmacy advice
- Report the use of any over the counter medications to their primary care prescriber and be aware they should discuss the use of lithium with their pharmacist before purchasing any OTC medicines.
- Women of child-bearing potential should use reliable contraception and inform the specialist or primary care prescriber immediately if they become pregnant or wish to become pregnant.





1. Background

Lithium is licensed for the treatment and prevention of mania, bipolar depression, recurrent depression (unipolar) and aggressive/self-mutilating behaviour. It is also recommended for augmentation of antidepressants in major depressive disorder.

Lithium can be very effective for acute episodes of mental illness, following which it is often continued. Likewise in prophylaxis, but longer periods of treatment may be required to establish its benefits. Not all patients respond to lithium, so the benefits and risks of continuation should be regularly and individually assessed. Lithium treatment should not be stopped suddenly, as this can cause relapse.

The benefits and many of the adverse effects of lithium relate to its plasma concentration. Lithium has a narrow therapeutic window of between 0.4 and 0.8 mmol/L for most indications, although a narrower range may be specified for individual patients. Higher target plasma levels (0.8–1 mmol/litre) are occasionally recommended for acute episodes of mania, for patients who have previously relapsed or when subthreshold symptoms of illness are associated with functional impairment. The specialist service will determine the target range for each patient and advise the primary care prescriber accordingly.

The plasma concentration of lithium is a function of absorption, distribution, and elimination. In salt form, lithium is readily absorbed from the gastrointestinal tract, but the rate and extent of absorption may differ between formulations. Levels fluctuate during distribution, so measurements are made 12 hours post-dose for monitoring purposes. Lithium is almost exclusively eliminated by the kidneys.

Lithium has numerous mild side effects but can be toxic if the dose is too high. Toxicity usually occurs with levels above 1.5 mmol/L but can emerge at lower levels in susceptible patients such as the elderly or those with renal impairment. Excluding excessive ingestion, toxicity most commonly arises due to a reduced elimination of lithium. Elimination is sensitive to sodium handling, so low-salt diets, dehydration, certain drug interactions and medical conditions such as Addison's disease are risk factors. Lithium toxicity can itself impair renal function, so rapid escalations in plasma levels may occur. Patients, carers, and clinicians should be familiar with the features of lithium toxicity, the common causes, and how to seek appropriate help.

With long-term use, lithium can have adverse effects on the kidneys, the thyroid, and the parathyroid glands. Routine monitoring of function is therefore required.

Lithium should always be prescribed by brand and form. Extra care must be taken when prescribing liquid forms, with clarity over the name and strength of the preparation. Patients should be involved in treatment decisions.

2. Indications

(Please state whether

licensed or unlicensed)

Licensed indications:

- Treatment and prophylaxis of mania
- Treatment and prophylaxis of bipolar disorder
- Treatment and prophylaxis of recurrent depression
- Treatment and prophylaxis of aggressive or self-harming behaviour





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	Augmentation of antidepressants in patients with treatment resistant
	depression [‡] See <u>NICE CG90: Depression in adults</u>
	[‡] Off-label indications. (Please note licensed indications vary by manufacturer).
3. Locally agreed off-label	To be agreed and completed locally (include supporting information)
use	
4. Contraindications and	Contraindications:
cautions	Hypersensitivity to lithium or any of the excipients. Excipients vary according
Please note this does not replace the Summary of Product Characteristics (SPC) and should be read in conjunction with it.	to the preparation being used. Tablets might include glycerol monostearate, glycerol distearate, mannitol, acacia, sodium lauryl sulfate, magnesium stearate, maize starch, sodium starch glycolate, gelatin and lactose. Liquid preparations contain ethanol. • Addison's disease
	Cardiac disease associated with rhythm disorder
	Cardiac insufficiency
	 Family or personal history of Brugada syndrome Patients with abnormal sodium levels, including dehydrated patients or those on low sodium diets Untreated hypothyroidism Severe renal impairment Breastfeeding
	Cautions:
	Mild to moderate renal impairment
	 Use in elderly patients Adequate and stable sodium and fluid intake should be maintained. This may be of special importance in hot weather, or during infectious diseases, including influenza, gastro-enteritis or urinary infections, when dose reduction may be required. Review lithium dose if diarrhoea and / or vomiting present and in cases where the patient has an infection and / or profuse sweating. Adjustments
	 may be required. Risk of seizures may be increased if co-administered with drugs that lower the seizure threshold, or in patients with epilepsy.
	 Cardiac disease May exacerbate psoriasis Surgery: discontinue 24 hours prior to major surgery and re-commence post-operatively once kidney function and fluid-electrolyte balance is normalised. Discontinuation is not required prior to minor surgery, providing fluids and electrolytes are carefully monitored.
	Please see <u>SPC</u> for comprehensive information.
5. Initiation and ongoing	Initial stabilisation:
dose regime	Usual starting dose for doses for all preparations are adjusted according to patient response and serum lithium concentration.
Note - •Transfer of monitoring and prescribing to primary care is normally after the	





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patient's dose has been optimised and with satisfactory investigation results for at least 4 weeks

- •The duration of treatment & frequency of review will be determined by the specialist, based on clinical response and tolerability.
- •All dose or formulation adjustments will be the responsibility of the initiating specialist unless directions have been discussed and agreed with the primary care clinician
- •Termination of treatment will be the responsibility of the specialist.

Most patients are prescribed lithium in tablet form (lithium carbonate). Doses may initially be divided throughout the day but once-daily administration is preferred when serum-lithium concentration is stabilised to target range (specified by specialist team).

In practice, the typical starting dose is 400 mg once daily, adjusted according to patient response and 12-hour plasma levels. Lower starting doses (such as 200 mg once daily) are preferable in the elderly and/or cases in which caution is required.

In some scenarios, such as acute mania, a higher starting dose (loading) may be preferable. The BNF outlines the typical starting doses by indication and brand.

Lithium citrate is absorbed at a different rate and to a different extent (bioavailability) compared to tablet forms. Extra care must be taken when prescribing lithium in liquid form, as some offer different strengths (mg/ml) under the same brand name (Li-liquid®) and some brand names (Priadel®) are used for the liquid and tablet forms. Switches between tablet and liquid formulations should be overseen by specialist services as dose conversions require the calculation of milligram equivalence between lithium carbonate and lithium citrate.

The loading period must be prescribed by the initiating specialist.

Maintenance dose (following initial stabilisation):

Individualised, to achieve plasma levels in the range specified for the patient. The initial maintenance dose must be prescribed by the initiating specialist.

Conditions requiring dose adjustment:

Lower doses may be required in older or physically frail/ low body weight patients, in mild to moderate renal impairment and electrolyte imbalance. Dose adjustments may also be required in patients prescribed interacting medicines.

Stopping lithium treatment

The decision to stop treatment will be the responsibility of the specialist. Clinicians, patients, and carers should be aware that abrupt discontinuation of lithium increases the risk of relapse. If lithium is to be stopped, the dose should gradually be reduced over a period of at least four weeks but preferably over a period of up to three months.

	period or op to the order	
6. Pharmaceutical aspects	Route of administration:	Oral
	Formulation:	Lithium is available as lithium carbonate (tablet formulations) and lithium citrate (liquid formulations). The patient should be maintained on the same brand and formulation of lithium. If a switch in brand or formulation is considered, refer to the specialist team.
		 Lithium Carbonate: Priadel® 200 mg and 400 mg prolonged-release tablets Camcolit® 400 mg controlled release tablets Liskonum® 450 mg controlled release tablets





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		 Lithium Citrate: Priadel® Liquid: 520 mg/5 mL strength sugar-free, pineapple flavoured syrup Li-Liquid®: 509 mg/5 mL and 1,018 mg/5 mL strength cherry flavoured syrup Always prescribe lithium by brand name. Switching preparation (either between brands of the same form or changing between tablets and liquid) additional monitoring to ensure that the 12-hour plasma lithium level remains in the desired range Particular care should be taken if prescribing liquid preparations; lack of clarity may lead to the patient receiving
		a sub-therapeutic or toxic dose. Consistency is paramount in lithium treatment and monitoring. Doses should be taken regularly, at the same time
	Administration details:	every day. Lithium carbonate tablets should not be crushed or chewed. Priadel® 200mg and 400mg tablets have score lines and can
		be divided accurately to provide dosage requirements as small as 100mg within product license.
		Liskonum® 450mg tablets are licensed to be halved for the purposes of dose adjustment.
		Other brands may be scored to facilitate breaking for ease of swallowing, and not to divide into equal doses. Breaking these tablets is not expected to alter their release properties but the accuracy of the division is not established
	Other important information:	If a dose is missed, then the next scheduled dose should be taken as usual; <u>a double dose should not be taken to make up for a missed dose.</u>
7. Significant medicine interactions	information and I	is not exhaustive; please see SPC for comprehensive recommended management. Igs must not be prescribed without consultation with
For a comprehensive list consult the BNF or Summary of Product Characteristics. SPC	 specialists: Drugs that may increase plasma lithium concentrations (by reducing renal elimination) and so risk toxicity: NSAIDs (including cyclo-oxygenase 2 inhibitors). If NSAID use is unavoidable, a dose reduction of lithium may be required and levels should be monitored more frequently. 'As required' use of NSAIDs should be avoided where possible since it may cause fluctuations in 	

lithium levels and makes monitoring levels challenging.

o Diuretics, particularly thiazide diuretics





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- Angiotensin converting enzyme (ACE) inhibitors and angiotensin II receptor antagonists
- Other drugs which alter electrolyte balance with the potential to alter lithium clearance e.g. steroids.
- Certain antibiotics including metronidazole and tetracyclines
- Drugs that may decrease plasma lithium concentrations (by increasing renal elimination) and so risk loss of efficacy:
 - Theophylline
 - o Products which contain sodium bicarbonate e.g. antacids
- Drugs that may increase risk of neurotoxicity when co-administered with lithium:
 - Calcium channel blockers (e.g. verapamil, diltiazem)
 - Antipsychotics (e.g. haloperidol, olanzapine, clozapine, flupentixol, chlorpromazine)
 - o Antidepressants with a serotonergic action (e.g. SSRIs, tricyclic antidepressants, venlafaxine, duloxetine)
 - Carbamazepine
- Drugs associated with QT prolongation (e.g. amiodarone, macrolides, tricyclic antidepressants) – potential for additive effects when co-administered with lithium.
- Drugs that lower seizure threshold (e.g. SSRIs, tricyclic antidepressants, antipsychotics) – increased risk of seizures

Care should be taken on initiation, dose adjustment or discontinuation of any interacting medicines. The onset and degree of the interaction can vary and additional lithium monitoring is likely to be indicated, with doses adjusted accordingly.

8. Baseline investigations, initial monitoring and ongoing monitoring to be undertaken by specialist

Monitoring at baseline and during initiation is the responsibility of the **specialist.** Only once lithium therapy is optimised on the chosen formulation with no anticipated further changes expected in immediate future will prescribing and monitoring be transferred.

Recent and relevant investigation results must be documented in the corresponding letter from specialist. In the event that baseline monitoring cannot be completed, for example due to patient refusal, this will also be documented in the letter from the specialist.

Baseline (all indications):

- Urea and electrolytes (U+Es), including calcium and eGFR
- Thyroid function tests (TFTs)
- Electrocardiogram (ECG) recommended for patients with existing cardiovascular disease (CVD) or risk factors
- Full blood count (FBC)
- Height, weight and body mass index (BMI)

Additional baseline investigations (bipolar disorder):

Cardiovascular status including pulse and blood pressure (BP)





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	profile. Liver function tests (LFTs). Initial monitoring: 12-hour plasma lithium levels on any change in dose or formulatio monitored weekly until the desir Ongoing monitoring:	e week after initiation and one week after on. Typically this means levels will be ed level and clinical effect is achieved. months to assess their mental health, he ongoing need for lithium.
9. Ongoing monitoring	Monitoring – all indications	Frequency
requirements to be	The GP must be able to access a	ı fast-track referral back to a secondary
undertaken by primary		point. The specialist will be available for
care.	1	an AMHT pharmacist if appropriate.
	Plasma lithium level taken 12 hours post-dose. In the event of twice daily dosing, withhold the morning dose before blood test for level. Record results in patient's NPSA purple lithium pack, NHS Health Monitor for Lithium app, or other suitable recording mechanism. It is advisable to document the actual time interval between the last dose and the blood sample	Measure the person's plasma lithium level every 3 months for the first year. After the first year, measure plasma lithium levels every 6 months, or every 3 months for people in any of the following groups: Older people People taking drugs that interact with lithium People who are at risk of impaired renal or thyroid function, raised calcium levels or other complications People who have poor symptom control People with poor adherence People whose last plasma lithium level was 0.8mmol per litre or higher.
	U+Es (including calcium and eGFR), TFTs Height, weight, and BMI.	Every 6 months. More frequent monitoring (particularly renal function) may be advised by the specialist team in some circumstances (e.g. elderly, renal impairment, altered TFTs, concurrent interacting medicines).
	Signs of toxicity	At every consultation

Enquire about and document signs

indicate toxicity, e.g. paraesthesia,

and symptoms which might





NHS Foundation Trust ataxia, tremor, cognitive impairment. Additional monitoring - bipolar Frequency disorder Annually as part of physical health check Diet, nutritional status and level of recommended by NICE (CG185 Bipolar physical activity. disorder: assessment and management). **Cardiovascular status including** pulse and BP. Metabolic status including fasting blood glucose, HbA_{1c} and blood lipid profile. LFTs. 10. Adverse effects and Result Action for GP 12-hour plasma lithium level. managements NB: range for each patient to be determined by the specialist team. Any serious adverse reactions should be reported to the MHRA **Below range** Assess adherence, including discussion via the Yellow Card scheme with patient and check of GP clinical www.mhra.gov.uk/yellowcard systems. Offer advice on adherence if appropriate (e.g. daily routines, reminders). Ensure level was taken 12 hours after lithium dose. Consider other factors e.g. drug interactions, excess fluid intake. Recheck level / contact specialist team for advice if suspected that the dose is too low. Within range Contact specialist team for advice. Referral to secondary care may be (signs of toxicity) required depending on the severity of symptoms and the certainty of toxicity. Use clinical judgement to determine the urgency of referral. Within range Repeat level. Assess adherence, including (but marked change since last discussion with patient and check of GP level and no dose change) clinical systems. Offer advice on adherence if appropriate (e.g. daily routines, reminders). Ensure level was taken 12 hours after lithium dose. More frequent monitoring may be required. Check if there is an explanation for the **Above range** (no signs of toxicity) high level e.g. dehydration, timing of level ie not 12 hrs post dose, interacting medicines, incorrect dose taken, patient's physical and mental status. Check for signs





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		of toxicity, Mithhold lithium and cont

of toxicity. Withhold lithium and contact specialist team for advice

If there is a possible explanation for high lithium level, correct where possible and recheck level.

If there is no explanation for the high level, recheck level, investigate renal function and if repeat level is higher than original target, refer back to specialist for advice.

If the trend is for the high end of the range but is not consistent with the range specified by the specialist team, decrease the dose, encourage fluids and recheck in one week.

If ≥2mmol/I – send patient to A&E and inform specialist team

Signs of toxicity

Typical signs and symptoms include diarrhoea, vomiting, loss of appetite, muscle weakness, lethargy, dizziness, ataxia, lack of coordination, tinnitus, blurred vision, coarse tremor of the extremities and lower jaw, muscle hyper-irritability, choreoathetoid movements, dysarthria, and drowsiness

If lithium toxicity is suspected, do an urgent lithium level immediately and seek specialist advice.

Referral to secondary care may be required depending on the severity of symptoms and the certainty of toxicity. Use clinical judgement to determine the urgency of referral.

Thyroid function

Altered TFTs without symptoms

Contact specialist team for advice. During lithium treatment, TFTs are commonly abnormal; the TSH can rise early in treatment but settle with time. Note that the symptoms of hypothyroidism can be difficult to discriminate from depression and the common side effects of lithium.

Subclinical hypothyroidism

- Raised TSH
- Normal T4

Contact specialist team for advice, which may include input from endocrinology services.





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	Overt hypothyroidism High TSH	The optimal management of subclinical hypothyroidism during lithium treatment remains controversial, with different thresholds for treatment advocated. Anticipate the need for additional monitoring, investigations and potentially thyroid hormone replacement based on specialist recommendations. Contact specialist team for advice, which may include input from endocrinology services.
	Low T4Symptomatic	Thyroid hormone replacement is usually indicated and often continued throughout the course of lithium treatment.
	Renal function	
	Polyuria and polydipsia	Polyuria is common with lithium and often well tolerated. Advise the patient to maintain adequate fluid intake and advocate excellent oral hygiene.
		Contact specialist team for advice, which may include input from nephrology services. In some instances, dose adjustment or specific treatments may be advocated.
	U&Es (including calcium) out of range	Check that the most recent 12-hour plasma lithium level is in the desired range and act accordingly if not.
		Determine whether there are symptoms and signs related to the electrolyte disturbance or lithium toxicity.
		Consider arranging an ECG in those at risk for QT prolongation.
		Contact specialist team for advice. Changes in calcium levels may reflect parathyroid dysfunction and input from endocrinology services may be indicated.
	eGFR <45ml/min rapidly falling eGFR gradual decline in eGFR	The response to impaired or deteriorating renal function should be individualised. Contact specialist team for advice, which may include input from nephrology services. A cardiovascular risk profile may guide specialist advice and should be provided if available. Use clinical judgement to determine the urgency of consultation.





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		Anticipate the need for increased
		monitoring as trends in renal function are
		more useful than absolute values. In the
		elderly or those at the extremes of muscle
		mass, creatinine clearance provides a
		better estimate of renal function that
		eGFR.
		Adjustments to dose may be advised. If renal function is significantly compromised, lithium may no longer be an appropriate treatment and specialists will advise accordingly.
	Weight and BMI	Provide appropriate support on
	Outside healthy range	multicomponent interventions to increase
	, , , , ,	physical activity levels, improve eating
		behaviour and quality of diet.
		Consider measuring waist circumference for individualised monitoring.
	Physical health check (bi-polar	Any physical health problems should be
	disorder)	treated by the appropriate primary care
		health professional and communicated to
		the specialist team within 14 days.

11. Advice to patients and carers

The specialist will counsel the patient with regard to the benefits and risks of treatment and will provide the patient with any relevant information and advice, including patient information leaflets on individual medicines.

The patient should be advised to report any of the following signs or symptoms to their GP without delay:

- Lithium toxicity (diarrhoea, vomiting, loss of appetite, muscle weakness, lethargy, dizziness, ataxia, lack of coordination, tinnitus, blurred vision, coarse tremor of the extremities and lower jaw, muscle hyper-irritability, choreoathetoid movements, dysarthria, and drowsiness)
- Signs of hypothyroidism (e.g. fatigue, cold intolerance, weight gain, constipation and depression), renal dysfunction (including polyuria and polydipsia), and benign intracranial hypertension (persistent headache and visual disturbance).

Additional advice for patients/ carers:

- Lithium should be taken regularly, as prescribed. If doses are missed, patients should not attempt to catch up or double dose
- Patients should not stop taking lithium suddenly doing so increases the chance of relapse. If lithium is to be stopped, it should be reduced over at least four weeks and preferably three months.
- The same brand of lithium should always be taken unless otherwise instructed. Patients should become familiar with their brand and check they have received the correct one before taking.
- Changes in hydration and sodium balance can affect plasma lithium levels. Patients should maintain adequate fluid intake, particularly in hot weather or when activity levels change (such as increases in exercise or immobility).





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Large changes in dietary sodium should be avoided – changing dietary regime may inadvertently alter sodium intake.

- Substantial changes in plasma lithium levels can occur if patients develop diarrhoea or vomiting, or if they become acutely ill for any reason. Patients should seek medical advice in such instances.
- Excessive alcohol consumption should be avoided as it can lead to dehydration, increasing plasma lithium levels and so risk of toxicity.
- Patients should be warned about common drug interactions and advised to
 present their 'Lithium alert card' whenever they redeem a new prescription.
 They should specifically be advised not to take OTC NSAIDs as these can
 increase plasma lithium levels and so risk toxicity.
- Lithium may impair performance of skilled tasks (e.g. driving, operating machinery). Patients with a diagnosis of bipolar disorder must notify the Driver and Vehicle Licensing Agency (DVLA).
- Women of childbearing potential should be advised that lithium carries
 additional risks in pregnancy and is a potential teratogen. They should be
 aware of the need to use reliable contraception and that they should tell
 their doctor straight away if they become pregnant while taking lithium.
 Lithium should not be taken if breastfeeding.
- For acute indications such as mania or augmentation, patients may respond
 within days to weeks of starting lithium. Depending on episode frequency, it
 may take months or even years to determine whether lithium has proven
 effective for release prevention.

At the start of treatment patients should be given suitable information on lithium and means to keep a record of their serum lithium levels, for example the NHS Health Monitor for Lithium app, or a purple lithium pack.

Patient information on this medicine can be found at the following links:

- NHS: https://www.nhs.uk/medicines/lithium/
- MIND: https://www.mind.org.uk/information-support/drugs-and-treatments/lithium-and-other-mood-stabilisers/lithium/
- National Patient Safety Agency purple lithium pack: https://www.sps.nhs.uk/wp-content/uploads/2018/02/2009-NRLS-0921-Lithium-patientet-2009.12.01-v1.pdf

12. Pregnancy, paternal exposure and breast feeding

It is the responsibility of the specialist to provide advice on the need for contraception to male and female patients on initiation and at each review but the ongoing responsibility for providing this advice rests with both the GP and the specialist.

All patients should be informed of the risks and benefits of taking this medicine during pregnancy and breastfeeding.

<u>Pregnancy</u>: Lithium should not be used during pregnancy, especially in the first trimester (risk of teratogenicity, including cardiac abnormalities). In certain cases where a severe risk to the patient could exist if treatment were stopped, lithium has been continued during pregnancy; under these circumstances prescribing is the responsibility of the specialist team.

If a patient becomes pregnant whilst on lithium, the specialist team should be informed immediately (but do not stop the lithium).

Women of child-bearing potential should be advised to use a reliable form of contraception. It is the responsibility of the specialist to provide advice on the need for contraception to patients on initiation of lithium, and at each review.





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	Under shared care agreements, the ongoing responsibility for providing this advice rests with both the GP and the specialist.		
	Breastfeeding:		
	Lithium is secreted in breast milk and there have been case reports of neonates		
	showing signs of lithium toxicity. Lithium should be avoided during		
	breastfeeding.		
	Paternal exposure:		
	Animal studies have reported spermatogenesis abnormalities that may lead to		
	impairment of fertility- it is unknown if this risk applies to humans.		
13. Specialist contact	Oxford Health Main Switchboard		
information	Tel: 01865 901000		
14. Additional information	Where patient care is transferred from one specialist service or GP practice to		
	another, a new shared care agreement must be completed.		
15. References	eBNF accessed via <u>www.medicinescomplete.com</u> on 17/02/2021.		
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	Pharma. Date of revision of the text: 24/08/2020. Accessed via		
	https://products.mhra.gov.uk/ on 17/02/2021.		
	Patient Information Leaflet. Priadel® 520mg/5mL liquid. Essential Pharma. Date		
	of revision of the text: June 2020. Accessed via https://products.mhra.gov.uk/ on 23/02/2021.		
	 Summary of Product Characteristics. Camcolit 400 mg, controlled release Lithium 		
	Carbonate. Essential Pharma. Date of revision of the text: 28/09/2020. Accessed		
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	Date of revision of the text: 14/05/2020. Accessed via		
	https://products.mhra.gov.uk/ on 23/02/2021.		
	• Summary of Product Characteristics. Li-Liquid 509 mg/5mL oral syrup. Rosemont.		
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	https://www.sps.nhs.uk/articles/suggestions-for-therapeutic-drug-monitoring-		
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	13th ed. London: Wiley-Blackwell; 2018, pp. 205-213.		





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	disorder/prescribing-information/lithium/ on 17/02/2021.
	• NHS UK leaflet: Lithium. Accessed via https://www.nhs.uk/medicines/lithium/on
	17/02/2021.
	National Patient Safety Agency. Safer Lithium Therapy. 2009. Archived resources
	available via: https://www.sps.nhs.uk/articles/npsa-alert-safer-lithium-therapy-
	<u>2009/</u> .
16. To be read in	RMOC Shared Care Guidance
conjunction with the	 NHSE/NHSCC guidance – items which should not be routinely prescribed in
following documents	primary care: guidance for CCGs
	 NHSE policy- Responsibility for prescribing between Primary &
	Secondary/Tertiary Care
17. Local arrangements for	Each mental health specialist will contact the patient's GP in writing and within
referral	this communication, will advise the GP about how to refer / escalate patients
Define the referral procedure from	back into AMHT.
hospital to primary care prescriber &	
route of return should the patient's	
condition change.	