SHARED CARE GUIDELINE

Drug: Methotrexate

Contact Details Name: Tel 🕿: Location:							
					Date:		NHS Number:
					Introduction	 Indication: Licensed: Rheumatoid arthritis, severe psoriasis, severe active juvenile idiopathic arthritis, severe psoriatic arthritis and Crohn's disease (pen device is unlicensed in Crohn's disease). Unlicensed: Severe Eczema, Lichen Planus, Felty's syndrome N.B. Not all brands/formulations are licensed for all indications – please refer to individu SPCs Background: Methotrexate is a folic acid antagonist and its major site of action is the enzyme dihydrofolate reductase. Its main effect is inhibition of DNA synthesis but it also impairs RNA and protein synthis may not account however for its action in rheumatoid arthritis or psoriasis which is not full understood. Response to treatment cannot be expected before two or three months and may not occur un six months of treatment. In patients with psoriasis response to treatment is also variable and take up to a month or more before any significant effect. Patients commenced on methotrexate are usually commenced on oral methotrexate. They n switched to methotrexate. Definitions: Stable dose – the dose will be titrated to achieve efficacy at the lowest dose. Once efficacy at and provided the patient can tolerate the dose, this will be termed "stable dose" 	
						and have stayed at similar levels for a	e active disease despite being on a stable dose or having stable
Form	_	e used to avoid confusion; do not use 10mg) ² on ranging from 7.5mg to 30mg in pre-filled pen) ³ .					
Dose & Administration	characteristics such as age, renal fur may be increased incrementally by 2.	ing on the indication and severity of the condition and patient action and other comorbid conditions. The dose of methotrexate 5-5mg every 1-6 weeks until disease is stabilised. The maximum exceptionally the dose may be increased to 30mg weekly. Folic					

 Fersponsibilities Exclude active infections. Check for absence of pregnancy in women of child-basing age and ensure the patient understands the importance of contraception. Reliable contraception should be used by both men and women whills on methortoxate and for at least 6 months after stopping methorexate. Discuss the boeffis and side effects of treatment with the patient. Ensure that the patient understands that dosing is ONCE WEEKLY and which warning symptoms to report. Perform pre-treatment screening: FBC, LFTs, U&Es, creatinine/ GFR and chest vary (unless done within 6 months). Pulmonary function tests should be consisted in patients with absential should be used by both men and women while on the patient. Ensure that the patient the the attent is now when and where to attend for monitoring. Encourage the patient to take responsibility for ensuring that results of tests are entered in the monitoring booklet. If initiating medication specifying the DAY OF THE WEEK on the prescription; don't use the dose term 'as directed'. Make arrangements for shared care with the patient's GP. Review the patient regularly to monitor the patient's response to therapy. Advise the GP on management of any dose adjustments and when to stop treatment. Ensure that clear backup arrangements exist for GPs to obtain advice. Methotrexate Injection pen: If the decision is made to switch to methotrexate injection pen provide one month's supply and a purple lided cytotoxic sharps bin. The Sharp Sate and Sharps Guard cyto com bins are examples of bins which will hold the pen davice. Provide training on self-administration of methotrexate injection with the dose. Provide training on self-administration of methotrexate injection with the patient. Provide the patient with prescriptions for methotrexate 2.5mg tablets and folic acid 5mg tablets. Do not prescribe		
Primary Care Responsibilities Methotrexate tablets: • Provide the patient with prescriptions for methotrexate 2.5mg tablets and folic acid 5mg tablets. Do not prescribe the 10mg tablets of methotrexate. Methotrexate injection pen: • Provide the patient with prescriptions for methotrexate pen as advised by the specialist and a 1L purple lidded cytotoxic sharps bin as required. The Sharp Safe and Sharps Guard cyto com bins are examples of bins which will hold the pen device. • Ensure systems are in place for the patient to receive their weekly injection if they are not self-administering. For both: • Ensure that the patient understands that dosing is ONCE WEEKLY and which warning symptoms to report. • Specify the DAY OF THE WEEK on the prescription; don't use the dose term 'as directed'. • Reinforce advice about using reliable contraception for both men and women whilst on methotrexate and for at least 6 months after stopping methoterxate • Monitor at the recommended frequencies (see MONITORING below) and ensure that test results are recorded in the monitoring booklet. • Report any adverse events to the consultant or specialist nurse and stop treatment on their advice or immediately if an urgent need arises (see MONITORING below). • Report any worsening of control of the condition to the consultant or the specialist nurse. • Follow recommended immunisation programme.	Secondary Care Responsibilities	 Exclude active infections. Check for absence of pregnancy in women of child-bearing age and ensure the patient understands the importance of contraception. Reliable contraception should be used by both men and women whilst on methotrexate and for at least 6 months after stopping methotrexate. Discuss the benefits and side effects of treatment with the patient. Ensure that the patient understands that dosing is ONCE WEEKLY and which warring symptoms to report. Perform pre-treatment screening: FBC, LFTs, U&Es, creatinine/ eGFR and chest x-ray (unless done within 6 months). Pulmonary function tests should be considered in patients with abnormal shadowing on x-ray. Dermatologists should include P3NP screening for patients with psoriasis. Provide the patient with a monitoring and dosage record booklet and ensure that the patient knows when and where to attend for monitoring. Encourage the patient to take responsibility for ensuring that results of tests are entered in the monitoring booklet. If initiating medication specifying the DAY OF THE WEEK on the prescription; don't use the dose term 'as directed'. Make arrangements for shared care with the patient's GP. Review the patient regularly to monitor the patient's response to therapy. Advise the GP on management of any dose adjustments and when to stop treatment. Ensure that clear backup arrangements exist for GPs to obtain advice. Methotrexate Injection pen: If the decision is made to switch to methotrexate injection pen provide one month's supply and a purple lidded cytotoxic sharps bin. The Sharp Safe and Sharps Guard cyto com bins are examples of bins which will hold the pen device. The first injection of Metoject[®] PEN should be performed under direct medical or nursing supervision in secondary care. Provide training on self-administration of methotrexate injection with the pen.
 Responsibilities Provide the patient with prescriptions for methotrexate 2.5mg tablets and folic acid 5mg tablets. Do not prescribe the 10mg tablets of methotrexate. Methotrexate injection pen: Provide the patient with prescriptions for methotrexate pen as advised by the specialist and a 1L purple lidded cytotoxic sharps bin as required. The Sharp Safe and Sharps Guard cyto com bins are examples of bins which will hold the pen device. Ensure systems are in place for the patient to receive their weekly injection if they are not self-administering. For both: Ensure that the patient understands that dosing is ONCE WEEKLY and which warning symptoms to report. Specify the DAY OF THE WEEK on the prescription; don't use the dose term 'as directed'. Reinforce advice about using reliable contraception for both men and women whilst on methotrexate and for at least 6 months after stopping methotrexate Monitor at the recommended frequencies (see MONITORING below) and ensure that test results are recorded in the monitoring booklet. Report any adverse events to the consultant or specialist nurse and stop treatment on their advice or immediately if an urgent need arises (see MONITORING below). Report any worsening of control of the condition to the consultant or the specialist nurse. Follow recommended immunisation programme. 		
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	 Pneumococcal vaccination recommended In patients exposed to chicken pox or shingles, if required, passive immunisation should be considered for varicella. Refer to Green book: <u>Varicella: the green book, chapter 34 - Publications - GOV.UK</u> Live vaccinations to be avoided. Shingles vaccine can be given as a precaution if dose of methotrexate is <0.4mg/Kg/week
Adverse Effects	 N.B. Please see MONITORING below for ADVERSE EFFECTS which require an intervention. This list is not exhaustive, please refer to SPCs and BNF. Headache, tiredness, drowsiness, erythema, pruritus, exanthema, dyspepsia, anorexia, leucopenia, anaemia, thrombopenia, pneumonia, elevated transaminases, nausea and vomiting, diarrhoea. Decreased resistance to infections.
Common Drug Interactions	 Trimethoprim and co-trimoxazole must be avoided Antifolate effect of methotrexate also increased by phenytoin. Caution with drugs with potential hepatotoxic or nephrotoxic effects. Tolbutamide – increases serum concentration of methotrexate NSAIDs, aspirin and penicillins are known to reduce the excretion of methotrexate causing an increase in serum level (increased risk of toxicity) but are not contraindicated.
Cautions	 Alcohol – cautions required, advise to stay well within national recommendations Ulcers of the oral cavity and known gastrointestinal ulcer disease Current illness that may cause renal impairment
Contra-indications	 Pregnancy – both men and women are advised to take contraceptive precautions while on methotrexate and for 6 months after stopping methotrexate. Breastfeeding. Serious active infection (suspected local or systemic). Severe renal or hepatic impairment. High alcohol intake/ alcohol abuse. Pre-existing blood dyscrasias, such as bone marrow failure or significant anaemia. Hypersensitivity to methotrexate. Some live vaccines – see under immunisation.

Monitoring and Adverse Effects	Treatment status	FBC	LFT	U+E	Creatinine/ eGFR	ESR or CRP	P3NP
	Initial monitoring	Weekly	Weekly	Weekly	Weekly	Every 3 months (for RA only)	N/A
	Once dose is stable (see definition)	Every 2 months	Every 2 months	Every 2 months	Every 2 months	Every 3 months (for RA only)	Annual for dermatology only (if elevated monitor every 3 months)
	 Once dose is stable monitoring frequency may be reduced to 3 monthly on consultant advice At dose increase changes advised by the specialist team, the monitoring will need to be weekly until dose and bloods stable. Thereafter revert back to standard monitoring as above. The patient should be asked about the presence of rash, oral ulceration, severe sore throat, abnormal bruising, diarrhoea, nausea and vomiting and whether they have new or increasing dyspnoea or cough, at each visit. If MCV > 105fL check thyroid function, B12 and folate. Treat any underlying abnormality but if these results are normal, discuss with specialist team for further advice. 						

	ving adverse laboratory results or patient reported symptoms, withhold ussed with specialist team:
 Significant deter Abnormal bruisin Rash, nausea an frequent toxic efference Cough or dyspnoe or dyspnoea met urgently. 	 < 3.5 x 10⁹/L or less than the lower limit of reference range as per lab < 2.0 x 10⁹/L or less than the lower limit of reference range as per lab < 150 x 10⁹/L or less than the lower limit of reference range as per lab > 2 times the upper limit of reference range ined fall (in absence of active disease) rioration in renal function. and vomiting, diarrhoea or oral ulceration. Diarrhoea and severe ulcerative stomatitis are fects and require interruption of therapy, otherwise haemorrhagic enteritis and death erforation may occur. and the interval of the and discussion with specialist team should take place

References

- 1. http://www.rheumatology.org.uk/includes/documents/cm_docs/2009/d/diseasemodifying_antirh eumatic_drug_dmard_therapy.pdf
- <u>http://www.medicines.org.uk/emc/medicine/24573/SPC/Methotrexate+2.5mg+Tablets+BP/</u>
 <u>https://www.medicines.org.uk/emc/medicine/28982</u>
- 4. BNF 66 September 2013-March2014
- 5. http://cks.nice.org.uk/dmards#!scenariorecommendation:9