

SHARED CARE FRAMEWORK APC BOARD DATE: 28 SEP 2017

CICLOSPORIN

Background Licensed Indications	Ciclosporin is a potent immunosuppressant Indications, dose adjustments and monitoring requirements for disease modifying drugs (DMDs) (licensed and unlicensed indications) included in this Framework are in line with national guidance published by the British Society for Rheumatology 2017. • Rheumatoid arthritis.
	 Psoriasis in patients in whom conventional therapy is inappropriate or ineffective. Atopic dermatitis when systemic therapy is required Inflammatory eye disease Transplant indications are not included
3. Locally agreed off-label use	 Interstitial lung disease Other rheumatology indications Myasthenia gravis Connective tissue disease Psoriatic arthritis Connective tissue disease Autoimmune and inflammatory kidney conditions Sarcoidosis Atypical neuro-inflammatory disease
4. Initiation and ongoing dose regime	Transfer of monitoring and prescribing to Primary care is normally after 3 months The duration of treatment will be determined by the specialist based on clinical response and tolerability Rheumatoid arthritis Initiation: For the first 6 weeks of treatment the recommended dose is 2.5 mg/kg/day orally given in 2 divided doses. If the effect is insufficient, the daily dose may then be increased gradually as tolerability permits, but should not exceed 5 mg/kg. Maintenance The dose will be titrated individually by the specialist to the lowest effective level according to tolerability, usually 2.5 -

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(or earlier if there is significant new evidence relating to this recommendation)

4mg/kg per day orally. **Psoriasis** Initiation: The recommended initial dose is 2.5 mg/kg/day orally given in 2 divided doses. If there is no improvement after 1 month, the daily dose may be gradually increased, but should not exceed 5 mg/kg. Initial doses of 5 mg/kg/day are justified in patients whose condition requires rapid improvement. Maintenance: Doses have to be titrated individually to the lowest effective level, and should not exceed 5 mg/kg/day. 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 Dose increases should be monitored by FBC creatinine/ eGFR, ALT and/or AST and albumin every 2 weeks for 6 weeks after the dose increase, then revert back to previous schedule. Termination of treatment will be the responsibility of the specialist. Baseline 5. Baseline investigations, initial Height, weight, BP, FBC, creatinine/ eGFR, ALT and /or monitoring and dose titration to AST. albumin. be undertaken by specialist • Vaccinations against pneumococcus and influenza are recommended. Shingles vaccine (Zostavax) is recommended as per the JCVI for eligible patients. Specialist to highlight in the first clinic letter notifying the GP of the decision to initiate DMDs that the GP will need to give the shingles vaccine if the patient is older than 69 years and the pneumococcal vaccine if this has not already been given. The GP should also be advised to add the patient to the influenza vaccine list. Patients should be assessed for comorbidities that may influence DMD choice, including evaluation of respiratory disease and screening for occult viral infection. Initiation • FBC, creatinine/ eGFR, ALT and /or AST and albumin every 2 weeks until on stable dose for 6 weeks; Once on stable dose, monthly FBC, creatinine/ eGFR, ALT and /or AST and albumin. Once patients have been stable for 12 months they can be considered for reduced frequency of monitoring, as advised by the specialist team. This monitoring consists of FBC, creatinine/ eGFR, ALT and/or AST and albumin at least every 12 weeks. BP and blood glucose should be measured at each monitoring visit. 6. Ongoing monitoring Monitoring Frequency Monthly until stable for 12 months,

FBC, creatinine/ eGFR,

then every 12 weeks or as advised

by the specialist. The exact frequency of the monitoring to be

ALT and/or AST,

Glucose

Albumin, BP, Blood

requirements to be undertaken

by primary care.

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	CRP and ESR (rheumatology patients only)	communicated by the specialist in all cases.	
N.B. For Rheumatology patients only - under the care of St Helens and Knowsley Hospitals: GP to choose whether monitored under Option 1 or Option 2	Option 1: GP to prescribe DMARD while monitoring undertaken via computerised Rheumatology Monitoring System (RMS). For patients with GPs who have access to Whiston pathology ICE system – results will be available via ICE For patients with GPs who do not have access to Whiston ICE, patients will be provided with blue record card of results which they will be advised to be made available to GP when writing prescription. N.B. Option 1 will be implemented by the Rheumatology Team if the patient's GP has not responded to the request for shared care after 21 days Option 2: GP to prescribe DMARD and monitoring to be undertaken via GP surgery.		
7. Pharmaceutical aspects	Route of administration	oral	
	Formulation Administration details	10mg, 25mg, 50mg, 100mg capsules, 100mg/1ml oral solution The daily doses of ciclosporin should	
	Administration details	be given in two divided doses The solution should be diluted, preferably with orange or apple juice, however, other drinks, such as soft drinks, can be used. (Rinse with more to ensure total dose is taken). Do not mix with grapefruit juice.	
	Other important information	Concomitant intake of grapefruit juice has been reported to increase bioavailability of ciclosporin.	
8. Contraindications Please note this does not replace the Summary of Product Characteristics (SPC) and should be read in conjunction with it.	 Hypersensitivity to the active substance or excipients. Concomitant use of tacrolimus Abnormal renal function, uncontrolled hypertension, uncontrolled infections or any malignancy SPC states live vaccines should be avoided; however JCVI and BSR recommend that oral DMD therapy at standard doses is not a contraindication in most patients, clinician discretion is advised. 		
9. Significant drug interactions	For a comprehensive list consult the BNF or Summary of Product Characteristics. SPC Ciclosporin interacts with all statins and is contraindicated with rosuvastatin MHRA Safety alert: Statins: interactions, and updated advice for atorvastatin Seek advice from the initiating Specialist if there are any concerns about interactions.		
10. Adverse Effects and	Result	Action	
managements	Abnormal bruising or severe sore throat Fall in WCC <3.5 x 109/l	Stop drug until FBC results available, contact Specialist Nurse (SN) Stop drug. Contact SN for advice	
	Fall in neutrophils <2.0 x 10 ⁹ /l Fall in platelets <140 x	and management	

	10 ⁹ /l		
	Increased MCV >105fl	Check folate, B12 & TSH.Treat if abnormal, contact SN for advice if normal.	
	Unexplained reduction in albumin <30g/l Abnormal LFTs – AST or ALT > 100U/l	Contact SN for advice and management	
	'Significant' increase in fasting lipids		
	Rash:	Stop drug and contact SN.	
	Taste loss:	Reassure, continue drug.	
	Nausea, vomiting, diarrhoea	Discuss with SN	
	Increase in serum creatinine >30% over period of 12 months or less OR decline in eGFR > 25%	Contact specialist nurse if there is new or unexplained renal impairment	
	Hyperkalaemia:	Stop drug and contact SN	
	Hypertension:	Consider anti-hypertensive agent. If hypertension persists, stop drug and contact SN	
	Gingival hypertrophy	Send patient for dental advice	
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 drugs.		
12. Pregnancy and breast feeding	Ciclosporin can be used during pregnancy at the lowest effective dose after considering the potential risks and benefits. BP, renal function and drug level should be monitored.		
	Mothers receiving treatment with ciclosporin should not be discouraged from breastfeeding.		
	(BSR & BHPR guideline on prescribing in pregnancy and breastfeeding)		
13. Specialist contact information	See appendix 2		
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	BSR monitoring guideling	nes	
16. To be read in conjunction	Policy for shared contact the shared contact t		
with the following documents.	2. Shared care agree	ement form	
		are initiated, one shared care completed for all relevant drugs.	

Appendix 1

Policy for Shared Care

Shared care is only appropriate if it provides an optimum solution for the patient and it meets the criteria outlined in the Shared Care section of the Pan Mersey **Definitions and Criteria for Categorisation of Medicines in the Pan Mersey Formulary** document.

Before prescribing responsibilities are transferred to primary care:

- Prescribing responsibility will only be transferred when the consultant and the patient's GP
 agree that the patient's condition is stable.
- All information required by the shared care framework for the individual medicine has been provided to the patient's GP.
- Patients will only be referred to the GP once the GP has agreed to the Shared Care Agreement and returned signed copies.

Inherent in any shared care agreement is the understanding that participation is at the discretion of the GP, subject to the availability of sufficient information to support clinical confidence.

Specialist Responsibilities in Shared Care

- To initiate the medicine, prescribe, monitor for toxicity and efficacy as described by the shared care framework until the patient is stabilised.
- To ensure the patient or their carer:
 - o Is counselled with regard to the risks and benefits of the medicine.
 - Provide any necessary written information to the patient with regard to the individual medicine including patient information leaflets on individual drugs.
 - Obtain and document informed consent from the patient when any medicines is prescribed for an off-label indication for any condition
- To be familiar with the shared care framework.
- To provide all information to the patient's GP as required by the shared care framework when
 prescribing responsibility is initially transferred and at any subsequent times as necessary for
 safe and effective treatment of the patient.
- To assess the patient regularly as necessary for the duration of therapy.
- To review the patient promptly if required by the GP concerned.
- To meet any additional requirements as required by the individual medicine shared care framework.
- To communicate failure of a patient to attend a routine hospital review and advise the GP of appropriate action to be taken.
- Addition of a second DMD: Following the addition of a new drug to an existing regime covered by a Shared Care Agreement, the Specialist must initiate, prescribe and monitor the

new drug in accordance with the relevant shared care agreement including subsequent review and inform the GP of this. A new Shared Care Agreement must then be initiated for the new drug.

• For Rheumatology patients only - under the care of St Helens and Knowsley Hospitals:
where GP chooses Option 1 – Blood test monitoring will remain the responsibility of
Rheumatology department via Rheumatology Monitoring System. Rheumatology department
takes responsibility for actioning abnormal blood test results. Blood test results will be
available to GP via Whiston Pathology ICE (or for GP practices that do not have access to
this, via patient hand held blue results card)

Primary Care Responsibilities in Shared Care

 To reply to a written request for Shared Care within 21 days ensuring both copies of the Shared Care Agreement are signed if appropriate.

If agreeing to shared care, the GP is asked to:

- To provide prescribe or manage and monitor the medicine as advised by the Specialist and in line with the individual Shared Care Framework.
- For Rheumatology patients only under the care of St Helens and Knowsley

 Hospitals: where GP chooses Option 1 GP to prescribe medication and ensure patient
 has been attending for blood tests via rheumatology monitoring system and that blood test
 results are available (via Whiston Pathology ICE system or patient held blue result card
 blood test monitoring).
- To review the patient as required by the Shared Care Framework
- To make appropriate and contemporaneous records of prescribing and/or monitoring and to note the existence of the Shared Care Agreement on the patient's clinical record. A READ code of "6652 Shared Care- Specialist/GP" can be used.
- To be familiar with the individual Shared Care Framework.
- To report any adverse effects of treatment to the specialist team.
- To inform the Specialist of any relevant change in the patient's circumstances.
- To seek Specialist advice as appropriate.
- To meet any additional requirements as required by the individual Shared Care Framework.
- To respond to Specialist communication relating to any change or addition to the patients treatment covered by the Shared Care Agreement.

<u>Disease modifying drugs (DMDs)</u>
Request by Specialist Clinician for the patient's GP to enter into a shared care agreement

<u> Part 1</u>

To be signed by Consultant / Associate Specialist / Specialist registrar or Specialist Nurse (who must be a prescriber)

Date	
Name of patient	
Address	
Patient NHS No	If using addressograph label please attach one to each copy
Patient hospital unit No	
Diagnosed condition	
Dear Dr	
I request that you prescribe	
(1)	
(2)	
(3)	
(4)	
for the above patient in accordance with the enclose	d shared care framework.
Last Prescription Issued: / / Next Sup	ply Due: /
Date of last blood test: / / Date of nex	t blood test: /
Frequency of blood test:	
I confirm that the patient has been stabilised and	reviewed on the above regime in
accordance with the Shared Care Framework and	d Policy.
I confirm that if this is a Shared Care Agreement	for a drug indication which is unlicensed o
off label, informed consent has been received.	N/A

Details of Specialist Clinicians

Name	Date
Consultant / Associate Specialis	st / Specialist Registrar / Specialist Nurse *circle or <u>underline</u> as appropriate
Signature	
In <u>all</u> cases, please also provide	e the name and contact details of the Consultant.
When the request for shared ca takes medico-legal responsibilit	are is made by a Specialist Nurse, it is the supervising consultant who y for the agreement.
Consultant:	
Contact details:	
Telephone number:	Ext:
Address for return of documentation	
Part 2 To be completed by Prim	ary Care Clinician
I agree to prescribe the enclosed shared care frame	for the above patient in accordance with ework.
GP signature	Date
GP name	Please print
For <u>Rheumatology patients or</u> I would like monitoring to be und	nly under the care of St Helens and Knowsley Hospitals lertaken
	onitoring System Yes / No the Rheumatology Team if the patient's GP has not responded to the request for
shared care after 21 days. Option 2 - at GP surgery	Yes / No
GP: Please sign and return a c	copy within 21 calendar days to the address above
OR	

GP- If you do not agree to prescribe, please delete the section above and provide any supporting information as appropriate below:

St Helens Rheumatology Monitoring System (RMS)

St Helens Rheumatology Department has developed an in-house computerised blood monitoring system for patients on DMARD therapies which has now been running for over 15 years. It was upgraded to a web-based programme in 2009.

Overleaf is a flow chart of this system.

It has a number of advantages over tradition shared care monitoring (where blood tests are taken, checked and transcribed in to patient held monitoring booklet by hand).

- These include:
 - 1) It minimises the number of health professionals involved in the process, reducing the risk of miscommunication
 - 2) It ensures prompt action on any abnormality being taken by an experienced rheumatology nurse specialist
 - 3) It is an efficient use of human resources using the computer to do the detection of the abnormality
 - 4) It reduces risk of human error an abnormal result being overlooked, or inaccurate transcription of blood test result to patient held monitoring booklet.
 - 5) It has a robust mechanism for detecting DNAs and enabling the appropriate action to be taken.

However its major disadvantage is that the results of the tests are sent to the patient on a blue card but the prescribing GP is then reliant on either the patient remembering to bring the blue card record of all their blood tests to the surgery when requesting a repeat prescription or the GP checking the results on the Whiston pathology system assuming they have access to this or the GP trusting in our monitoring system (and I appreciate that they may not feel able to do so).



RHEUMATOLOGY MONITORING SYSTEM (RMS) PATHWAY (2018)

