**SHARED CARE GUIDELINE**

**Drug:** Sodium Aurothiomalate (Gold injection)

| Introduction | Indications:  
Licensed – Active, progressive rheumatoid arthritis, progressive juvenile chronic arthritis especially if polyarticular or seropositive.  
Unlicensed – skin diseases including pemphigus  

Background:  
The mechanism of action of Sodium aurothiomalate is not known.  
Benefit should not be expected until a cumulative dose of at least 300-500mg has been given. If there is no response after a cumulative dose of 1000mg has been given, alternative DMARD therapy will be considered.  

Definitions:  
Stable dose – the dose will be titrated to achieve efficacy at the lowest dose. Once efficacy achieved and provided the patient can tolerate the dose, this will be termed "stable dose"  
Stable bloods – results of blood tests remain below the “alert” thresholds as set by national guidelines and have stayed at similar levels for at least two consecutive tests.  
N.B. The patient can continue to have active disease despite being on a stable dose or having stable bloods, so the “patient” is not referred to as “stable”

| Form | Myocrisin 100mg/ml solution for injection, 0.5ml ampoules  

| Dose & Administration | • Sodium aurothiomalate should only be administered by deep intramuscular (IM) injection followed by gentle massage of the area. The patient should remain under medical observation for a period of 30 minutes after drug administration.  
• Typical dose: An initial 10mg test dose (administered in secondary care) in the first week, followed by 50mg doses weekly until signs of remission occur.  
• In patients showing signs of remission, 50mg doses should be given at two weekly intervals until full remission occurs.  
• With full remission, the interval between injections should be increased progressively to three and then four weeks.  
• After 18 months to 2 years, the interval between injections is to be increased to six weeks.  
• If after reaching a total dose of 1000mg (excluding the test dose), no major improvement has occurred other forms of treatment are to be considered.  
N.B. Do not use a darkened solution (more than pale yellow).  

| Secondary Care Responsibilities | • Confirm the diagnosis.  
• Check for absence of pregnancy in women of child-bearing age and ensure the patient understands the importance of contraception.  
• Discuss the benefits and side effects of treatment with the patient. Ensure that the patient understands which warning signs and symptoms to report.  
• Perform pre-treatment screening: height, weight, blood pressure, FBC, LFT, albumin and, creatinine/ calculated GFR, urinalysis for blood and protein and chest x-ray  
• Patients should be assessed for co-morbidities, including evaluation for respiratory disease and screening for occult viral infection  
• Administer a 10mg test dose and observe the patient for 30 minutes for signs of allergic reaction.  
• Ensure that the patient understands not to expect improvement for the first few injections.  
• Provide the patient with prescriptions for Sodium Aurothiomalate (Myocrisin®) injection until on stable dose and undergoing 3 monthly monitoring. 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.

- 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 frequency of monitoring, frequency of injections and when to stop treatment.
- Ensure that clear backup arrangements exist for GPs to obtain advice.

| **Primary Care Responsibilities** | • Provide the patient with prescriptions for Sodium aurothiomalate (Myocrisin®) once on stable dose and undergoing 3 monthly monitoring and make the necessary arrangements for administration of the injection.
• 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. |
| **Immunisations** | Annual flu vaccine is recommended
Pneumococcal vaccination recommended
In patients exposed to chicken pox or shingles, if required, passive immunisation should be considered for varicella. Refer to Green book: Varicella: the green book, chapter 34 - Publications - GOV.UK |
| **Common Drug Interactions** | This list is not exhaustive, please refer to SPCs and BNF
• ACE inhibitors
• Penicillamine |
| **Cautions** | • Elderly
• Moderate renal or hepatic impairment
• History of urticaria or eczema
• History of colitis
• If phenylbutazone or oxyphenbutazone are administered concurrently
• Irreversible skin pigmentation (chrysiasis) can occur in sun-exposed areas after prolonged treatment with sodium aurothiomalate. Patients should be advised to limit exposure to the sun by wearing protective clothing and using high factor sunscreens. |
| **Contraindications** | • Severe renal or hepatic impairment
• History of blood disorders or marrow aplasia
• Exfoliative dermatitis
• Systemic lupus erythematosus
• Necrotising enterocolitis
• Pulmonary fibrosis
• Acute Porphyria
• Pregnancy and breastfeeding
• Co-prescribing of penicillamine |

This guidance does not replace the SPC’s, which should be read in conjunction with this guidance.
**MONITORING AND ADVERSE EFFECTS**

<table>
<thead>
<tr>
<th>Treatment Status</th>
<th>FBC</th>
<th>LFT</th>
<th>Albumin</th>
<th>Creatinine / calculated GFR</th>
<th>Urinalysis (blood and protein)</th>
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</thead>
<tbody>
<tr>
<td>Initial monitoring until on stable dose for 6 weeks</td>
<td>Every 2 weeks</td>
<td>Every 2 weeks</td>
<td>Every 2 weeks</td>
<td>Every 2 weeks</td>
<td>Prior to each dose</td>
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<tr>
<td>For next three months</td>
<td>Monthly</td>
<td>Monthly</td>
<td>Monthly</td>
<td>Monthly</td>
<td>Prior to each dose</td>
</tr>
<tr>
<td>Thereafter</td>
<td>Every 3 months</td>
<td>Every 3 months</td>
<td>Every 3 months</td>
<td>Every 3 months</td>
<td>Prior to each dose</td>
</tr>
</tbody>
</table>

*Please note:* If the patient is also being treated with leflunomide, increased monthly monitoring is required, as specified in the leflunomide shared care guidance. (Where other biologic/DMARDs are used in combination with sodium aurothiomalate, the standard monitoring requirements, as outlined above, continue to apply).

As per secondary care responsibilities, for clarity the frequency of monitoring should be specified in the initial shared care request.

Dose increases should be monitored by FBC, creatinine / calculated GFR, albumin and LFTs every 2 weeks until on stable dose for 6 weeks and then revert to previous schedule.

The team responsible for prescribing the medication should also hold responsibility or monitoring i.e. prescribing to be carried out in Primary care only once patient on stable dose and undergoing 3 monthly monitoring

- The patient should be asked about the presence of rash, unusual bruising or mouth ulcers, unexplained breathlessness or cough. If present, withhold until discussed with specialist team.
- Results of FBC, including numerical platelet count, at the time of each injection need not be available before the injection is given, but must be available before the next injection. However urinalysis must precede monthly administration.

If 2+ proteinuria or more check MSSU. If infection present treat appropriately. If sterile and 2+ proteinuria or more persists on two consecutive occasions, STOP sodium aurothiomalate and discuss with the specialist team.

In the event of the following adverse laboratory results or patient reported symptoms, withhold sodium aurothiomalate injections until discussed with specialist team and repeat the test after two weeks:

- WCC < 3.5 x 10^9/L or less than the lower limit of reference
- Neutrophils < 1.6 x 10^9/L or less than the lower limit of reference
- Platelets < 140 x 10^9/L or less than the lower limit of reference
- AST/ALT > 100U/I
- MCV > 105Fl
- Creatinine increase >30% over 12 months and / or calculated GFR <60ml/min
- Unexplained eosinophilia >0.5 x 10^9/l
- Unexplained reduction in albumin <30g/l
- Rash or oral ulceration
- Abnormal bruising or severe sore throat: Check FBC immediately
- New or increasing dyspnoea or dry cough STOP sodium aurothiomalate as a precaution and discuss urgently with specialist team.
As well as responding to absolute values in laboratory tests, it is also relevant to observe trends in results (e.g. gradual decreases in white blood cells or albumin, or increasing liver enzymes). If urgent clinical abnormalities arise emergency access to specialist rheumatology advice should be sought.

**Other adverse effects:**

- Haematuria - requires investigation
- Anaphylactoid reactions are rare but may occur a few minutes after the injection. Advise the specialist team and do not give any further doses.
- Blood dyscrasias, hepatotoxicity, peripheral neuropathy and Guillain-Barre syndrome.
- Colitis

This list is not exhaustive; please refer to SPCs and BNF.

**References**

1. [https://www.medicines.org.uk/emc/medicine/18613](https://www.medicines.org.uk/emc/medicine/18613) SPC Myocrisin 100mg/ml Solution for Injection
2. BSR/BHPR Non-Biologic DMARD Guidelines 2017

**RELEVANT CONTACT LIST**

<table>
<thead>
<tr>
<th>Speciality</th>
<th>Name and Title</th>
<th>Tel. No.</th>
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Shared Care Agreement - Disease Modifying Drugs (DMARDs)

Request by specialist Clinician for the patient’s GP to enter into a shared care agreement

Reference:                     Date:

Patient name:                RXR/NHS number:

Patient address:

Diagnosis: ________________________________

In accordance with the shared care guidelines I kindly request that you prescribe:

1. ____________________ Dose ___________ Frequency ___________

2. ____________________ Dose ___________ Frequency ___________

3. ____________________ Dose ___________ Frequency ___________

for the above named patient:

Shared care guidelines available @ http://www.elmmb.nhs.uk/policies-and-guidelines/shared-care-guidelines/

Last Prescription issued: ________________ Next prescription due: ________________

Date of last blood test: ________________ Date of next blood test: ________________

Frequency of Blood test: ________________________________

I can confirm that the patient has been stabilised and reviewed on the above regime in accordance with the Shared Care guideline.

If this is a Shared Care Agreement for a drug indication which is unlicensed or off label, I confirm that informed consent has been received.

I will accept referral for reassessment at your request. The clinical team in the rheumatology department are available to give you advice.

Details of Specialist Clinician

Name: ________________________________ Date: ________________

Consultant/ Associate Specialist/ Specialist Registrar /Specialist Nurse (circle or underline as appropriate)

When the request for Shared Care is made by a specialist nurse, it is the supervising consultant who takes medicolegal responsibility for the agreement.

Consultant: ________________________________

Contact details for rheumatology specialist nurses ELHT: elht.rheumatologynurses@nhs.net

Telephone number: 01254 734491 or 01254 734569

Unless we hear from you within 14 days, we will assume that the Shared Care agreement has been accepted.

Yours sincerely,

The Rheumatology Directorate, ELHT