

Approved by Hertfordshire and West Essex Area prescribing Committee

#### NHS Hertfordshire and West Essex

#### Kev to terms:

**DMARD:** disease-modifying anti-

rheumatic drug

ILi: interleukin inhibitors that target type 17,12 or 12/23 interleukins

JAKi: Janus kinase inhibitor MTX: methotrexate

PASI 75 response: reduction in psoriasis area severity index (PASI) score of at least 75% from baseline

**PsARC:** psoriatic arthritis response criteria

TA: NICE technology appraisal
TNFi: tumour necrosis factor

inhibitor

### Pathway for treatment of Psoriatic Arthritis in adults.

Trial of Has the patient had adequate trials of at least 2 conventional DMARDs (administered conventional either individually or in combination) **DMARDs** YES, but failing to respond or patient cannot tolerate or has contraindications Patient does not meet pathway No Does the patient have peripheral arthritis, with 3 or more tender joints and 3 or more criteria. Use standard systemic swollen joints? treatments and best supportive care. Yes

- 1. Adalimumab biosimilar is the best value biologic and preferred 1st line
- 2. For every line of therapy, the least expensive appropriate treatment should be chosen
- 3. If treatment is not tolerated or response is inadequate consider changing to an alternative treatment from a different class.

  NOTE if however, intolerance occurs within the first 3 months, another agent from that class can be trialled prior to moving

NOTE If, however, intolerance occurs within the first 3 months, another agent from that class can be trialled prior to moving to next line.

4 If the patient has both psoriatic arthritis and psoriasis, take into account both conditions before initiating or making changes to treatment with high cost drugs. The specialty for the more severe condition would generally be expected to take responsibility for prescribing high cost drug treatment *Note tofacitinib and golimumab are not commissioned for psoriasis*.

#### **BOX 1. First line biologic treatment options**

Usually start treatment with least expensive appropriate treatment
Options listed by mechanism of action and overall cost from left to right

TNFi: Adalimumab biosimilar TA199 – usual first line choice,

Etanercept biosimilar TA199, Infliximab biosimilar TA199, Certolizumab TA445, Golimumab TA220

JAKi: Upadacitinib TA768, Tofacitinib (with MTX) TA543

IL-17i: Secukinumab 150mg TA445, Bimekizumab TA916, Ixekizumab

TA537, Secukinumab 300mg TA445 see Note 1

<u>IL-23i</u>: Guselkumab NICE TA 815 see Note 2 <u>IL-12 /IL-23i</u>: Ustekinumab TA340 see Note 3

#### Box1.a First line apremilast

#### Apremilast (TA433) oral tablet

Apremilast is a clinically effective treatment compared with placebo but not as clinically effective as TNFis.

For use in patients who are needle phobic for whom JAKi (oral tablets) are not appropriate or are contraindicated.

In line with TA433 apremilast is NOT commissioned for use after TNFi.

Assess Response. Relevant NICE TAs state that the first response should be measured at 12 weeks - 24 weeks depending on treatment modality.

#### Adequate response to treatment is defined as:

- improvement in at least 2 of the 4 PsARC Criteria (see Note 4), 1 of which must be the joint tenderness or swelling score AND
- no worsening in any of the 4 criteria

For patients whose PsARC response does not justify continuation of treatment but who have psoriasis and a PASI 75 response, treatment should be assessed by a dermatologist to determine whether continuing is appropriate on the basis of skin response. (See Note 5)

# BOX 2. Second line treatment options Consider alternative treatment from a different class. The least expensive appropriate treatment should be

**TNFi** listed in BOX 1 may be used second line (local agreement) also **see Note 6** 

JAKi: Upadacitinib TA768, Tofacitinib (with MTX) TA543

<u>IL-17i</u>: Secukinumab 150mg TA445, Bimekizumab TA916, Ixekizumab TA537, Secukinumab 300mg TA445 **see Note 1** 

IL-23i: Guselkumab TA815, Risankizumab TA803 see Note 7

IL-12 /IL-23i: Ustekinumab TA340

Assess response as for first line. Stop treatment if adequate response not maintained / intolerance and consider next pathway step.

## **BOX 3. Third, fourth- and fifth-line treatment options** (TAs and local agreement)

Consider alternative treatment from a different class. The least expensive appropriate treatment should be chosen

JAKi: Upadacitinib TA768, Tofacitinib (with MTX) TA543

<u>I IL-17i</u>: Secukinumab 150mg TA445, Bimekizumab TA916, lxekizumab TA537, Secukinumab 300mg TA445 see Note 1 <u>IL-23i</u>: Guselkumab TA815, Risankizumab TA803 see Note 7

IL-12 /IL-23i: Ustekinumab TA340

Assess response as for first line. Stop treatment if adequate response not maintained / intolerance and consider next pathway step.

If treatment not tolerated, or no response or inadequate response, or loss of response move to next line in pathway

## If treatment is tolerated and there is adequate response

Maintain same treatment and monitor patient and PsARC. Clinical review to be carried out at 6-12 monthly intervals.

Stop treatment if adequate response not maintained / intolerance and consider next pathway step

**NOTE 1:** Cost order is dependent on dose :secukinumab 150mg maintenance dose is less expensive than bimekizumab and ixekizumab, secukinumab 300mg maintenance dose is more expensive than bimekizumab and ixekizumab.

NOTE 2: Guselkumab is only commissioned first line if TNFi are contra-indicated. NOTE 3: Ustekinumab is only commissioned first line if TNFi are contra-indicated. NOTE 4: The PsARC comprises:

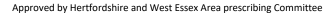
- A patient global self-assessment (on a 0–5 Likert scale); a physician global assessment (on a 0–5 Likert scale) with improvement defined as a decrease by at least 1 unit, and worsening defined as an increase by at least 1 unit)
- A tender joint score: a swollen joint score (with improvement defined as a decrease of at least 30%, and worsening defined as an increase of at least 30% from baseline.

NOTE 5: see local pathway and TAs103,134,146,180,350, 419,442,574,521 and 596 for guidance on the use of biologic drugs for the treatment of Psoriasis

NOTE 6: certolizumab is only commissioned second line if disease has stopped responding after the first 12 weeks of 1<sup>st</sup> line TNFi

**NOTE 7**: Risankizumab is only commissioned for patients who have had at least one biologic treatment and also have moderate to severe psoriasis (a body surface area of at least 3% affected by plaque psoriasis and a Psoriasis Area and Severity Index [PASI] score greater than 10. If both guselkumab and risankizumab are suitable the least costly treatment should be chosen.

Maximum of five sequential treatments per patient are routinely commissioned.







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Developed by	Pharmacy and Medicines Optimisation Team Hertfordshire and West Essex (HWE) ICB with relevant HWE ICS stakeholders.
Review Date	This HWE APC recommendation is based upon the evidence available at the time of publication. This recommendation will be reviewed upon request in the light of new evidence becoming available
Superseded version	V1.0 Risankizumab added up to 5 <sup>th</sup> line and guselkumab criteria amended, in line with NICE TA 803 and 815 and existing pathway principles.  V2.0 Bimekizumab added up to 5 <sup>th</sup> line in line with NICE TA 916 and existing pathway principles. Nov 2023  V3.0 Amended to remove note 7 'reduced dose interval of 4-weekly is not routinely commissioned for guselkumab'. Wording aligned with psoriasis pathway and text box advising updates to pathway.

NICE recommends if patients and their clinicians consider a medicine to be one of a range of suitable treatments, the least expensive treatment should be chosen, taking into account administration costs, dosage, price per dose and commercial arrangements. Therefore, in line with this recommendation and HWE APC agreed principles the order of preference of treatments within this pathway will be updated accordingly as prices change or biosimilar medicines become available.