Urticaria Treatment Pathway

Guideline Summary

This clinical guideline outlines the treatment pathway for adult patients with urticaria

1. Scope

This treatment pathway applies to adult patients with a diagnosis of urticaria and is primarily to set out the different treatment options available in secondary care

2. Rationale

This treatment pathway provides an evidence-based approach for the treatment of urticaria whilst maximising cost effectiveness and clinical outcome for use by all healthcare professionals involved in patient care.

3. Background

Chronic urticaria (CU) is a disease characterised by pruritic weals, angio-oedema or both occurring for at least 6 weeks. Around half of patients present with weals alone, 40% with weals and angio-oedema and 10% with angio-oedema only. It encompasses chronic spontaneous urticaria (CSU) and chronic inducible urticarias.

For information and guidance on management of urticaria in primary care please refer to the <u>NICE Clinical Knowledge Summary (CKS)</u> on urticaria

Patients can have an inducible element to their urticaria which is triggered by heat, cold, pressure, vibration, water, ultraviolet light (UV), etc. These urticarias are induced reproducibly after a specific physical stimulus is applied, however there can be a certain degree of overlap between spontaneous and inducible urticarias.

Unlicensed use of medicines: a number of medications recommended within the pathway are not licensed for use in urticaria and so are being used 'off-label'. These have been highlighted with an asterisk * throughout the document for information. Patients should be informed and consent to receive such treatments.

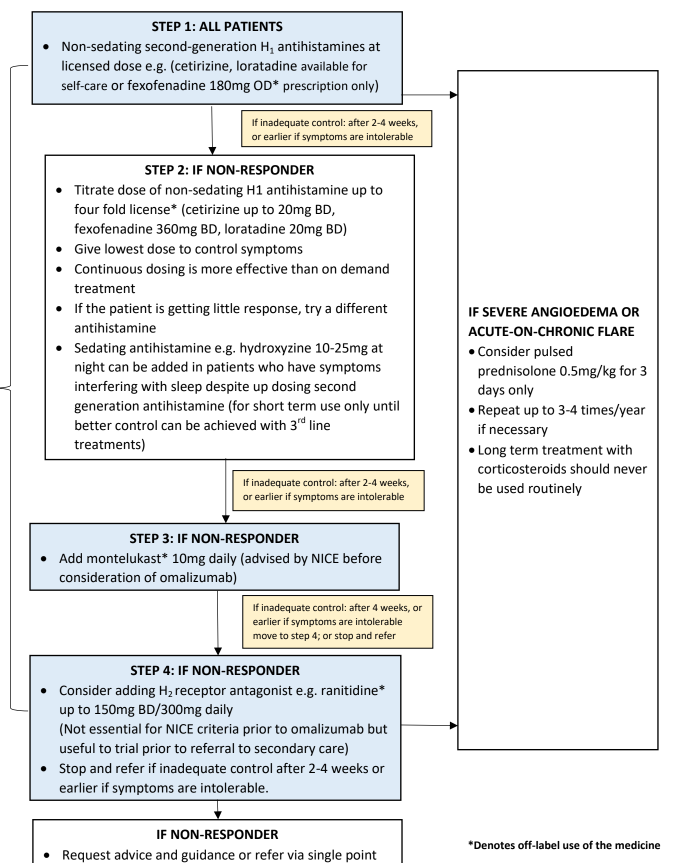
The use of Omalizumab within secondary care for the inducible urticarias listed in section 2 of the treatment pathway requires the submission of a category B* notification form to the relevant CCG.

See appendix 1 for prescribing responsibilities and RAG rating as per the South East London Joint Formulary.

4. Treatment Pathway

Primary care

For ALL types of urticaria

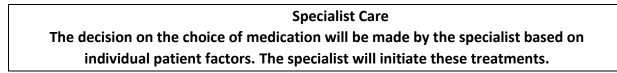


South East London Area Prescribing Committee. A partnership between NHS organisations in South East London: Bexley, Bromley, Greenwich, Lambeth, Lewisham and Southwark Clinical Commissioning Groups (CCGs) and GSTFT/KCH /SLAM/ & Oxleas NHS Foundation Trusts/Lewisham & Greenwich NHS Trust

of referral ERS to community dermatology service

Review Date: December 2020 (or sooner if evidence or practice changes)

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1) CHRONIC SPONTANEOUS URTICARIA



Trial of omalizumab if CSU severe despite optimal antimediator treatments in line with <u>NICE TA339</u> See dosing schedule below



Consider tranexamic acid* 500mg BD- 1.5gram TDS

IF AUTOIMMUNE URTICARIA suspected (e.g. positive basophil histamine release assay) Consider ciclosporin* 4mg/kg/day starting dose reducing to 2mg/kg if possible (treatment duration generally ≤ 1 year).

- Other immunosuppressive agents for CSU and omalizumab failures are:
- Methotrexate* up to 30mg weekly $\ge 3/12$
- Mycophenolate mofetil* up to 1500mg BD or hydroxychloroquine* up to 200mg BD would be last line options
- Azathioprine* 2mg/kg/day up to 2.5mg/kg/day in steroid dependent patients

Consider doxepin* 25-50mg at night in the appropriate context (CSU often coexists with anxiety and depression)

If non-responder, reassess in a specialist urticaria clinic

Omalizumab in CSU

- 1. Eligibility
 - ≥ 12 years of age
 - Severity of condition is assessed objectively e.g. using a weekly urticaria activity score (UAS) of ≥ 28 or angioedema activity score (AAS) ≥ 28/week
 - Not responded to standard treatment with H₁-antihistamines and leukotriene receptor antagonists

2. Dosing

- 300mg every 4 weeks for 4 doses and review
- Omalizumab is stopped at or before the fourth dose if the condition has not responded
- Omalizumab is stopped at the end of a course of treatment (6 doses) if the condition has responded, to establish whether the condition has gone into spontaneous remission, and is restarted only if the condition relapses

3. Administration

• Omalizumab is administered as an outpatient in a secondary care specialist centre in dermatology, immunology or allergy clinic due to a very small risk of anaphylaxis

*Denotes off-label use of the medicine

2) INDUCIBLE URTICARIAS

a) SYMPTOMATIC DERMOGRAPHISM

1. Consider narrow band UVB phototherapy for at least 6 weeks

2. If DLQI ≥15 consider omalizumab* (150mg monthly increasing to 300mg monthly if necessary)

Omalizumab stopping criteria

Stop at or before the fourth dose if condition has not responded/DLQI does not reduce below 10 \mp No increase in objective wealing threshold from 36g/mm² using a calibrated dermographometer

b) CHOLINERGIC URTICARIA

- Consider danazol* 200-600mg daily (in divided doses) in men, propranolol* up to 40mg BD, or oxybutynin* 5mg 2-3 times a day increased to 5mg QDS if necessary. If failure of these consider propantheline* up to 30mg QDS or hyoscine butylbromide* 10mg TDS increased up to 20mg QDS. All of the above should be stopped if no response after 6 weeks at the maximum dose.
- 2. If DLQI ≥15 consider omalizumab* (150mg monthly increasing to 300mg monthly if necessary) Omalizumab stopping criteria

Stop at or before the fourth dose if condition has not responded/DLQI does not reduce below $10 \mp$ No subjective increase in exercise or heat tolerance

c) DELAYED PRESSURE URTICARIA

- 1. Consider dapsone* 50mg/day up to max. 150mg/day or sulfasalazine* 0.5-4g/day (if not aspirin sensitive). Stop after 6 weeks if no response at the maximum dose.
- 2. If DLQI ≥15 consider omalizumab* (150mg monthly increasing to 300mg monthly if necessary) <u>Omalizumab stopping criteria</u>

Stop at or before the fourth dose if condition has not responded/DLQI does not reduce below 10 Ŧ

d) COLD URTICARIA

- 1. If DLQI ≥15 consider omalizumab* (150mg monthly increasing to 300mg monthly if necessary)
- 2. Consider ciclosporin* (4mg/kg for 4 weeks reducing by 1mg/kg every 6 weeks to zero for omalizumab non-responder. Longer treatment is an option)

Omalizumab stopping criteria

Stop at or before the fourth dose if condition has not responded/DLQI does not reduce below 10 Ŧ No fall in temperature threshold using Temp Test

e) SOLAR URTICARIA

- 1. Consider ciclosporin* (4mg/kg for 4 weeks reducing by 1mg/kg every 6 weeks to zero. Longer treatment is an option)
- 2. If DLQI ≥15 consider omalizumab* (150mg monthly increasing to 300mg monthly if necessary)

Omalizumab stopping criteria

Stop at or before the fourth dose if condition has not responded/DLQI does not reduce below 10 F No objective improvement in monochromator phototest thresholds

Ŧ In line with licensed use, omalizumab should be stopped at 6 months if the condition has responded, and restarted only if the condition relapses.

• As part of the local commissioning arrangements for omalizumab in inducible urticarias, a category B* notification form will need to be submitted to commissioners for patients initiated on this treatment.

*Denotes off-label use of the medicine

3) OTHER URTICARIAS

- a) IDIOPATHIC PRURITIS
- Consider amitriptyline* (up to 75mg ON), pregabalin* (up to 75mg BD), or gabapentin* (up to 600mg TDS) can be used for symptoms of dysesthesia/pruritus.
- Naltrexone* (initially 25mg daily increased to 50mg per day. Total weekly dose may be divided and given on 3 days of the week – max. 350mg per week) may rarely be used for idiopathic pruritus that has not responded to amitriptyline* (up to 75mg ON), pregabalin* (up to 75mg BD) or gabapentin* (up to 600mg TDS), reviewed every 3 months. Stop if no clinical response
- b) URTICARIAL VASCULITIS OR AUTOINFLAMMATORY SYNDROMES
- Consider colchicine* 0.5mg BD up to 2.5mg daily (in divided doses)
- Hydroxychloroquine*, azathioprine*, methotrexate* or corticosteroids may also be considered

*Denotes off-label use of the medicine

Appendix 1

Traffic Light Status Information

- **Red** Specialist / hospital prescribing only.
- Amber 1 treatment can be initiated in primary care after a recommendation from an appropriate specialist
- Amber 2 Specialist initiation and supply followed by maintenance prescribing in primary care
- Amber 3 specialist initiation with ongoing monitoring required. Transfer of prescribing to the GP using either the approved APC GP Information sheets where applicable or full shared care (drugs indicated with **).
- Green specialist and non-specialist initiation

Green	Amber 1	Amber 2	Amber 3	Red
 Non-sedating second generation H1 antihistamines at licensed dose Non-sedating antihistamines up to four fold license Ranitidine Montelukast Prednisolone 		 Tranexamic acid Colchicine Amitriptyline Oxybutynin Propantheline Hyoscine butylbromide Pregabalin Gabapentin 	GP Information sheets: • Doxepin • Danazol • Naltrexone Full shared care: • Ciclosporin** • Mycophenolate** • Methotrexate ** • Hydroxychloroquine** • Sulfasalazine** Azathioprine**	Omalizumab Dapsone

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