Tralokinumab

We recommend tralokinumab in AE patients who are candidates for systemic treatment.

Tralokinumab: in licence for ≥ 12 years;
age 12-17: initially 600 mg s.c. day 1 followed by 300 mg Q2W
adults: initially 600 mg s.c. day 1 followed by 300 mg Q2W
At prescriber’s discretion, every fourth week dosing may be considered for patients who achieve clear or almost clear skin after 16 weeks of treatment.

Certainty of evidence(1, 2):
Short term (8–16 weeks) vs placebo (NMA medications used in clinical practice or likely to be approved soon)
★★★★ HIGH for mean difference/ standardized mean difference EASI, change in signs, POEM, peak pruritus
NRS, DLQI, QoL, itch
★★ ◯◯ LOW for undesirable effects

Mechanisms of action and efficacy

Tralokinumab is a fully human, high affinity IgG4 mAb, which neutralizes IL-13, and has been approved by the EMA in summer 2021.(3) In two 52-week, double-blind, placebo-controlled, phase III trials, adults with moderate-to-severe AE were randomized to subcutaneous tralokinumab 300 mg every 2 weeks or placebo.(4) Tralokinumab monotherapy was superior to placebo at 16 weeks of treatment. Co-primary endpoints were IGA score of 0 or 1 and EASI 75 at week 16. Patient achieving an IGA score of 0/1 and/or EASI 75 with tralokinumab at week 16 were re-randomized to tralokinumab Q2W or every 4 weeks or placebo for 36 weeks. The majority of week 16 tralokinumab-responders maintained response at week 52 with continued tralokinumab treatment without any rescue medication. In a randomised, double-blind, phase III trial 301 adolescent patients received either 300 mg or 150 mg of tralokinumab or placebo. After 16 weeks, significantly more patients in the tralokinumab arms showed an EASI 75 response (27.8%, 28.6%, 6.4%) or an IGA of 0 or 1 (17.5%, 21.4%, 4.3%). Subjects achieving a clinical response (IGA = 0, 1; or EASI75) at week 16 without use of rescue medication were re-randomized to maintenance dosing regimens. At 52 weeks, EASI-75 response ranged from 44.4% to 63.3% in the different maintenance dosing regimens.(5)

Dosage: acute flare, short term, long term

The recommended dosage is 300 mg every 2 weeks after a loading dose of 600 mg at treatment onset. At prescriber’s discretion, every fourth week dosing may be considered for patients who achieve clear or almost clear skin after 16 weeks of treatment.

Phase III trials have also investigated what happens when patients who do well for 16 weeks on tralokinumab continue treatment as labeled, reduce treatment frequency, or discontinue treatment. After 16 weeks, patients who reached EASI 75 or IGA success were re-randomized to continue treatment every two weeks, titrate down to every four weeks, or use placebo. At 52 weeks, without
TCS, more than 55% of patients who continued twice-monthly treatment maintained EASI 75, as did approximately 50% of patients treated monthly. More than 51% of patients who stayed on twice-monthly dosing maintained IGA 0 or 1, versus 39% and 45% of patients who switched to monthly dosing.

**Safety**

In the two studies, adverse events were reported in 76.4% and 61.5% of patients receiving tralokinumab and in 77.0% and 66.0% of patients receiving placebo in the 16-week initial period.

Notably, tralokinumab appears to have lower rates of ocular complications than dupilumab.(4) The combination therapy with TCS, TCI and UV light treatment is possible.

**Monitoring**

No biochemical or instrumental exams are reported to be required for the monitoring of the therapy.

**Combination with other treatments**

In an additional phase III double-blind, placebo study the efficacy and safety of tralokinumab in combination with TCS as needed in patients with moderate-to-severe AE were evaluated. At week 16, significantly more tralokinumab-treated patients than placebo achieved IGA 0/1 and EASI 75. Nine out of ten EASI 75 responders at week 16 maintained response at week 32 with continued tralokinumab and TCS as needed.(6)
References


