Tralokinumab for moderate-to-severe atopic dermatitis: results from two 52-week, randomized, double-blind, multicentre, placebo-controlled phase III trials (ECZTRA 1 and ECZTRA 2)*

A. Wollenberg ⁽¹⁾ A. Blauvelt ⁽¹⁾, ² E. Guttman-Yassky ⁽¹⁾, ³ M. Worm ⁽¹⁾, ⁴ C. Lynde, ^{5,6} J.-P. Lacour ⁽¹⁾, ⁷ L. Spelman ⁽¹⁾, ^{8,9} N. Katoh ⁽¹⁾, ¹⁰ H. Saeki ⁽¹⁾, ¹¹ Y. Poulin, ¹² A. Lesiak, ¹³ L. Kircik ⁽¹⁾, ^{14,15} S.H. Cho, ¹⁶ P. Herranz ⁽¹⁾, ¹⁷ M.J. Cork ⁽¹⁾, ¹⁸ K. Peris ⁽¹⁾, ¹⁹ L.A. Steffensen ⁽¹⁾, ²⁰ B. Bang ⁽¹⁾, ²⁰ A. Kuznetsova, ²⁰ T.N. Jensen, ²⁰ M.L. Østerdal, ²⁰ E.L. Simpson²¹ and on behalf of the ECZTRA 1 and ECZTRA 2 study investigators

¹Department of Dermatology and Allergy, Ludwig Maximilian University of Munich, Munich, Germany

²Oregon Medical Research Center, Portland, OR, USA

³Department of Dermatology and the Immunology Institute, Icahn School of Medicine at Mount Sinai, New York, NY, USA

⁴Division of Allergy and Immunology, Department of Dermatology, Venereology and Allergy, Charité – Universitätsmedizin Berlin, Berlin, Germany

⁵Lynde Dermatology, Probity Medical Research, Markham, ON, Canada

⁶Department of Medicine, University of Toronto, Toronto, ON, Canada

⁷Department of Dermatology, University Hospital of Nice, Nice, France

⁸Veracity Clinical Research, Brisbane, QLD, Australia

⁹Probity Medical Research, Woolloongabba, QLD, Australia

¹⁰Department of Dermatology, Graduate School of Medical Science, Kyoto Prefectural University of Medicine, Kyoto, Japan

¹¹Department of Dermatology, Nippon Medical School, Tokyo, Japan

¹²Laval University and Centre Dermatologique du Québec Métropolitain and Centre de Recherche Dermatologique du Québec Métropolitain, Québec, QC, Canada

¹³Department of Dermatology and Pediatric and Oncologic Dermatology, Medical University of Łódź, Łódź, Poland

¹⁴Icahn School of Medicine at Mount Sinai, New York, NY, USA

¹⁵Indiana University Medical Center, Indianapolis, IN, USA

¹⁶Department of Dermatology, The Catholic University of Korea, Seoul, South Korea

¹⁷Department of Dermatology, Hospital Universitario La Paz, Madrid, Spain

¹⁸Sheffield Dermatology Research, Department of Infection, Immunity, and Cardiovascular Disease, The University of Sheffield and Sheffield Teaching Hospitals NIHR Clinical Research Facility, Sheffield, UK

¹⁹Dermatology, Catholic University and Fondazione Policlinico Universitario A. Gemelli, IRCCS, Rome, Italy

²⁰LEO Pharma A/S, Ballerup, Denmark

²¹Department of Dermatology, Oregon Health & Science University, Portland, OR, USA

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Summary

Correspondence

Andreas Wollenberg. Email: wollenberg@lrz.uni-muenchen.de

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The tralokinumab ECZTRA 1 and 2 studies were sponsored by LEO Pharma A/S (Ballerup, Denmark). Medical writing and editorial assistance were provided by Kathryn Woods, PhD, and Jane Beck, MA, from Complete HealthVizion, funded by LEO Pharma A/S. Background Tralokinumab, a fully human monoclonal antibody, specifically neutralizes interleukin-13, a key cytokine driving peripheral inflammation in atopic dermatitis (AD). In phase II studies, tralokinumab combined with topical corticosteroids provided early and sustained improvements in AD signs and symptoms.

Objectives To evaluate the efficacy and safety of tralokinumab monotherapy in adults with moderate-to-severe AD who had an inadequate response to topical treatments. Methods In two 52-week, randomized, double-blind, placebo-controlled, phase III trials, ECZTRA 1 and ECZTRA 2, adults with moderate-to-severe AD were randomized (3 : 1) to subcutaneous tralokinumab 300 mg every 2 weeks (Q2W) or placebo. Primary endpoints were Investigator's Global Assessment (IGA) score of 0 or 1 at week 16 and \geq 75% improvement in Eczema Area and Severity Index (EASI 75) at week 16. Patients achieving an IGA score of 0 or 1 and/or EASI 75 with tralokinumab at week 16 were rerandomized to tralokinumab

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Conflicts of interest

Conflicts of interest statements are listed in Appendix 1.

A full list of study investigators is provided in Appendix S1 (see Supporting Information).

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Q2W or every 4 weeks or placebo, for 36 weeks. The trials were registered with ClinicalTrials.gov: NCT03131648 and NCT03160885.

Results At week 16, more patients who received tralokinumab vs. placebo achieved an IGA score of 0 or 1: $15\cdot8\%$ vs. $7\cdot1\%$ in ECZTRA 1 [difference $8\cdot6\%$, 95% confidence interval (CI) $4\cdot1-13\cdot1$; P = $0\cdot002$] and $22\cdot2\%$ vs. $10\cdot9\%$ in ECZ-TRA 2 ($11\cdot1\%$, 95% CI $5\cdot8-16\cdot4$; P < $0\cdot001$) and EASI 75: $25\cdot0\%$ vs. $12\cdot7\%$ ($12\cdot1\%$, 95% CI $6\cdot5-17\cdot7$; P < $0\cdot001$) and $33\cdot2\%$ vs. $11\cdot4\%$ ($21\cdot6\%$, 95% CI $15\cdot8-27\cdot3$; P < $0\cdot001$). Early improvements in pruritus, sleep interference, Dermatology Life Quality Index, SCORing Atopic Dermatitis and Patient-Oriented Eczema Measure were observed from the first postbaseline measurements. The majority of week 16 tralokinumab responders maintained response at week 52 with continued tralokinumab treatment without any rescue medication (including topical corticosteroids). Adverse events were reported in $76\cdot4\%$ and $61\cdot5\%$ of patients receiving tralokinumab in ECZTRA 1 and ECZTRA 2, respectively, and in $77\cdot0\%$ and $66\cdot0\%$ of patients receiving placebo in ECZTRA 1 and ECZTRA 2, respectively, in the 16-week initial period.

Conclusions Tralokinumab monotherapy was superior to placebo at 16 weeks of treatment and was well tolerated up to 52 weeks of treatment.

What is already known about this topic?

- Atopic dermatitis (AD) is a chronic interleukin (IL)-13-mediated disease.
- There is a need for safe and effective long-term treatment options for AD.
- Tralokinumab is a fully human monoclonal antibody that binds specifically to IL-13 with high affinity, thereby preventing receptor interaction and subsequent downstream signalling.
- Tralokinumab combined with topical corticosteroids showed early and sustained efficacy and safety in a 12-week, phase IIb trial in moderate-to-severe AD.

What does this study add?

- These are the first pivotal phase III trials demonstrating that by specifically targeting IL-13 alone, patients can achieve significant improvements in AD signs and symptoms and quality of life, and maintain these improvements over time without the requirement for topical corticosteroids.
- These trials provide evidence that tralokinumab offers a long-term, well-tolerated treatment option for patients with moderate-to-severe AD.

Atopic dermatitis (AD) is a chronic inflammatory skin disorder with an estimated prevalence in adults of between 2.1% and 4.9% across North America, Europe and Japan.¹ As we gain a better understanding of the inflammatory pathways driving AD, targeted treatment options are being developed to improve long-term therapeutic options for patients with moderate-to-severe AD.^{2–4} Recent evidence identified the type 2 cytokine interleukin (IL)-13 as a key driver of the underlying inflammation in AD.^{5–7} IL-13 is implicated in skin barrier disruption, skin inflammation, increased risk of skin infections, itch signalling and epidermal hyperplasia,⁶ with levels of IL-13 in lesional skin correlating with AD severity.^{8–13} Currently, dupilumab (an IL-4 receptor α antagonist) is the only biologic available for the treatment of moderateto-severe AD.

Tralokinumab, a fully human IgG4 monoclonal antibody, specifically binds with high affinity to IL-13 alone, preventing its interaction with the receptor and subsequent downstream signalling.¹⁴ In a phase II trial, tralokinumab provided early and sustained improvement in disease signs and symptoms in adult patients with moderate-to-severe AD.¹⁵ Here we report the results of the first phase III trials to investigate the long-term efficacy and safety of IL-13specific inhibition with tralokinumab monotherapy compared with placebo in adult patients with moderate-to-severe AD for up to 1 year, as assessed by the severity and extent of AD, itch, Staphylococcus aureus colonization and health-related quality of life.

Patients and methods

Study design and oversight

ECZTRA 1 (NCT03131648) and ECZTRA 2 (NCT03160885) were identically designed 52-week, multinational, randomized, double-blind, placebo-controlled trials. Patients were randomized 3 : 1 to subcutaneous tralokinumab 300 mg, after a 600-mg loading dose on day 0, or placebo every other week for 16 weeks (Figure S1; see Supporting Information). Randomization was performed using a computer-generated randomization schedule stratified by region and baseline disease severity. Treatment allocation was blinded to patients and investigators (Appendix S2; see Supporting Information). After a 16-week initial treatment period, tralokinumab-treated patients who achieved the prespecified criteria for clinical response of Investigator's Global Assessment (IGA) score of 0 (clear) or 1 (almost clear), or $\geq 75\%$ improvement in Eczema Area and Severity Index (EASI 75) were rerandomized 2 : 2 : 1 to tralokinumab 300 mg every 2 weeks (Q2W) or every 4 weeks (Q4W), or placebo for a 36-week maintenance treatment period.

Patients who achieved the clinical response criteria with placebo continued to receive placebo Q2W to maintain blinding of the study and were not included in analyses after week 16. Patients not achieving the clinical response criteria at week 16 were transferred to open-label tralokinumab 300 mg Q2W with optional topical corticosteroids (TCS). Additionally, patients who, during maintenance treatment, experienced decline in effect by meeting specific protocol-defined transfer criteria over a 4-week period (Appendix S2) were transferred to open-label tralokinumab. All patients had a final safety follow-up 16 weeks after the last dose of study medication, unless transferred to the long-term ECZTEND trial (NCT03587805).

Prior to randomization, AD treatments were washed out: 4 weeks for systemic treatments and 2 weeks for TCS and other topical treatments. Rescue treatment (Appendix S2) could be used at the discretion of the investigator to control intolerable symptoms and did not disqualify patients from continuing to randomized or open-label treatment. However, patients who received rescue treatment (including TCS) were considered nonresponders in the primary analyses (see Statistical analysis). Patients were instructed to use an emollient twice daily throughout the trials.

The trials were sponsored by LEO Pharma and conducted in accordance with the ethical principles derived from the Declaration of Helsinki and Good Clinical Practice guidelines and were approved by the local institutional review board or ethics committee of each institution. All patients provided written informed consent. Patients were enrolled from 30 May 2017 to 5 March 2018 in ECZTRA 1 and from 29 June 2017 to 26 April 2018 in ECZTRA 2.

Study population

Patients were ≥ 18 years of age, with a diagnosis of AD for ≥ 1 year, who were candidates for systemic therapy due to a recent (within 1 year) history of inadequate response to treatment with topical treatments or for whom topical treatments were medically inadvisable. Patients were required to have EASI ≥ 12 at screening and ≥ 16 at baseline, an IGA score of ≥ 3 and AD involvement of $\geq 10\%$ of body surface area at screening and baseline, and worst daily pruritus numerical rating scale (NRS) average score ≥ 4 during the week prior to baseline (Appendix S2; see Supporting Information).

Efficacy outcomes

Outcomes were analysed according to a prespecified hierarchy (Figure S2; see Supporting Information). Primary endpoints were IGA score of 0 (clear skin) or 1 (almost clear skin) at week 16 and EASI 75 at week 16.

Key secondary endpoints at week 16 included in the hierarchy were reduction of weekly average worst daily pruritus NRS of \geq 4 points, change in SCORing Atopic Dermatitis (SCORAD)¹⁶ and change in Dermatology Life Quality Index (DLQI),¹⁷ all from baseline to week 16.

Maintenance endpoints assessed at week 52 and included in the hierarchy were IGA score of 0 or 1 in patients initially randomized to tralokinumab, with IGA score of 0 or 1 at week 16 achieved without rescue medication (including TCS), and EASI 75 in patients initially randomized to tralokinumab, with EASI 75 at week 16 achieved without rescue medication (including TCS).

Additional secondary endpoints included the proportions of patients achieving \geq 50% and \geq 90% improvement in EASI (EASI 50 and EASI 90), change in EASI, change in worst daily pruritus NRS, reduction in worst daily pruritus NRS by \geq 3 points, change in Patient-Oriented Eczema Measure (POEM) and reduction in DLQI by \geq 4 points at week 16.

S. aureus colonization on lesional skin was assessed in ECZ-TRA 1 and antidrug antibodies were assessed in both studies (Appendix S2; see Supporting Information).

Safety assessments

All adverse events (AEs) were recorded and classified by severity, causality and outcome. Clinical laboratory tests, vital signs and other safety assessments were performed at scheduled visits. A data-monitoring committee independent of the trial and LEO Pharma reviewed safety data during the study.

Statistical analysis

For the primary endpoint of IGA score of 0 or 1 at week 16, a sample size of 780 patients randomized 3 : 1 would provide > 99% power to detect a target difference between the two arms with a response rate of 30% in the tralokinumab arm and 10% in the placebo arm, with a 5% two-sided level of

significance. For EASI 75 at week 16, a sample size of 780 would provide > 99% power to detect a target difference between tralokinumab and placebo at week 16, assuming EASI 75 response rates of 40% and 15% for tralokinumab and placebo, respectively (Appendix S2; see Supporting Information). To control the overall type 1 error rate at a 5% significance level, a prespecified testing hierarchy was used for assessment of the primary, key secondary and maintenance endpoints (Figure S2; see Supporting Information).

A number of prespecified statistical analyses were conducted for binary and continuous endpoints applying different analytical approaches to handle missing data, rescue medication use and permanent discontinuation (Appendix S2). The primary analytical approach for the binary endpoints considered patients who received rescue medication (including TCS) and patients with missing data to be nonresponders. The difference in response rates between treatment groups was analysed using the Cochran–Mantel–Haenszel test stratified by region and baseline disease severity (IGA score of 3 or 4). An alternative analysis was also applied, where all observed data were used irrespectively of rescue medication use, with missing data imputed as nonresponse.

For the primary analyses of the continuous endpoints, a repeated-measurements model was used, where data collected after permanent discontinuation or initiation of rescue medication were excluded from the analysis, and the model included all data up to the timepoint of discontinuation or rescue medication use. The model included baseline IGA, region and treatment-by-week interaction as factors, and interaction between week and baseline value as a covariate. An unstructured covariance matrix was used to model the within-patient errors.

For the maintenance endpoints of IGA score of 0 or 1 and EASI 75 at week 52, patients who, prior to week 52, received rescue medication and/or were transferred to open-label treatment were considered nonresponders. The differences in response rates were analysed using the Cochran–Mantel–Haenszel test stratified by region.

Results

Patients

In total, 802 patients were enrolled in ECZTRA 1 and 794 patients were enrolled in ECZTRA 2 (Figure S3; see Supporting Information). Baseline demographics and disease characteristics were well balanced across treatment groups (Table 1). About half of all patients had severe AD (IGA score of 4). The median EASI at baseline ranged from 28.2 to 30.3, the median disease duration was 25-28 years and the median affected body surface area was approximately 50% in all randomized groups. Almost all patients (98-100%) had a history of TCS use prior to enrolment; systemic corticosteroids had been used by 59.2-69.1% and ciclosporin by 32.3-37.6% (Table S1; see Supporting Information). Discontinuation rates prior to week 16 were similar between treatment groups in ECZTRA 1 but greater in the placebo group in ECZTRA 2 (Figure S4; see Supporting Information). Rescue medication was used in the first 16 weeks by 35.8% and 22.8% of patients

 Table 1 Demographics and clinical characteristics of randomized patients at baseline

	ECZTRA 1 ^a		ECZTRA 2 ^b		
	Placebo, n = 199	Tralokinumab Q2W, n = 603	Placebo, n = 201	Tralokinumab Q2W, n = 593	
Age (years), median (IQR)	37.0 (26.0-49.0)	37.0 (27.0-48.0)	30.0 (23.0-46.0)	34.0 (25.0-48.0)	
Male sex, n (%)	123 (61.8)	351 (58·2)	114 (56.7)	359 (60.5)	
Race, n (%)					
White	138 (69.3)	426 (70.6)	123 (61.2)	374 (63.1)	
Black	18 (9.0)	41 (6.8)	17 (8.5)	43 (7.3)	
Asian	40 (20.1)	120 (19.9)	52 (25.9)	154 (26.0)	
Other or missing data	3 (1.5)	16 (2.6)	9 (4.5)	22 (3.7)	
Disease duration (years), median (IQR)	28.0 (18.0-41.0)	27.0 (19.0-38.0)	25.0 (18.0-36.0)	25.5 (17.0-39.0)	
Affected BSA (%), median (IQR)	52.5 (31.0-77.0)	50.0 (33.0-70.0)	50.0 (31.0-74.0)	50.0 (31.0-74.0)	
EASI, median (IQR)	30.3 (22.0-41.5)	28.2 (21.3-40.0)	29.6 (20.6-41.4)	28.2 (19.8-40.8)	
IGA score of 4, n (%)	102 (51.3)	305 (50.6)	101 (50.2)	286 (48.2)	
SCORAD score, median (IQR)	70.8 (63.8-81.0)	69.2 (61.5–79.1)	69.9 (61.9–79.1)	69.5 (60.5-79.1)	
Weekly average worst daily pruritus NRS, median (IQR)	7.9 (6.9–8.7)	7.9 (6.7–8.9)	8.1 (7.1–9.0)	8.0 (7.0–9.0)	
DLQI score, median (IQR)	16.0 (13.0-22.0)	17.0 (12.0-22.0)	18.0 (12.5-24.0)	18.0 (13.0-23.0)	
Weekly average eczema-related sleep interference NRS, median (IQR)	7.0 (5.7–8.0)	7.1 (5.7–8.4)	7.9 (6.4–8.6)	7.4 (6.2–8.7)	
POEM score, median (IQR)	24.0 (20.0-27.0)	24.0 (20.0-27.0)	24.0 (20.0-27.5)	24.0 (20.0-27.0)	

BSA, body surface area; DLQI, Dermatology Life Quality Index; EASI, Eczema Area and Severity Index; IGA, Investigator's Global Assessment; IQR, interquartile range; NRS, numerical rating scale; POEM, Patient-Oriented Eczema Measure; Q2W, every 2 weeks; SCORAD, SCORing Atopic Dermatitis. ^aECZTRA 1 enrolled patients from France, Germany, Japan, Spain and the USA. ^bECZTRA 2 enrolled patients from Australia, Canada, Denmark, Italy, Korea, Poland, Russia, the UK and the USA.

receiving tralokinumab in ECZTRA 1 and ECZTRA 2, respectively, and was used at a greater rate in the placebo groups (46.2% and 44.3%, respectively). The majority of patients using rescue medication used TCS, and use of systemic rescue treatment was greater in the placebo groups (Table S2; see Supporting Information). The average time to initiation of rescue medication was shorter in the placebo groups (Figure S4).

Primary outcomes

In both studies, achievement of an IGA score of 0 or 1 and EASI 75 at week 16 was significantly higher with tralokinumab vs. placebo (Table 2). IGA 0 or 1 was achieved by 15.8% with tralokinumab vs. 7.1% with placebo in ECZTRA 1 [difference 8.6%, 95% confidence interval (CI) 4.1–13.1, P = 0.002] and by 22.2% with tralokinumab vs. 10.9% with placebo in ECZTRA 2 (difference 11.1%, 95% CI 5.8–16.4; P < 0.001). These results are depicted in Figure 1(a), Table 2 and Table S3 (see Supporting Information).

EASI 75 was achieved by 25.0% with tralokinumab vs. 12.7% with placebo in ECZTRA 1 (difference 12.1%, 95% CI 6.5–17.7; P < 0.001) and by 33.2% with tralokinumab vs. 11.4% with placebo in ECZTRA 2 (difference 21.6%, 95% CI 15.8–27.3; P < 0.001), as shown in Figure 1(b), Table 2 and

Table S4 (see Supporting Information). Following an alternative analysis, irrespectively of rescue medication use, response rates were improved for all treatment arms in both studies (Table S5; see Supporting Information).

Key secondary outcomes at week 16

Tralokinumab showed a significant improvement vs. placebo in all key secondary endpoints. Reduction of weekly average worst daily pruritus NRS by \geq 4 points from baseline to week 16 was achieved by 20.0% with tralokinumab vs. 10.3% with placebo in ECZTRA 1 (difference 9.7%, 95% CI 4.4–15.0; P = 0.002; Table 2) and by 25.0% vs. 9.5% in ECZTRA 2 (difference 15.6%, 95% CI 10.3–20.9; P < 0.001; Table 2). The adjusted mean changes in SCORAD from baseline to week 16 were -25.2 vs. -14.7 in ECZTRA 1 (difference -10.4%, 95% CI -14.4 to -6.5; P < 0.001; Table 2; and Tables S6 and S7; see Supporting Information) and -28.1 vs. -14.0 in ECZTRA 2 (difference -14.0%, 95% CI -18.0 to -10.1; P < 0.001; Table 2; and Tables S6 and S7; see Supporting Information)

The adjusted mean changes in DLQI from baseline to week 16 were -7.1 for tralokinumab vs. -5.0 for placebo in ECZ-TRA 1 (difference -2.1%, 95% CI -3.4 to -0.8; P = 0.002; Table 2; and Tables S8 and S9; see Supporting Information)

Table 2 Efficacy outcomes for the initial treatment period: primary and key secondary endpoints, full analysis set

	ECZTRA 1			ECZTRA 2			
	Placebo, n = 197	Tralokinumab Q2W, n = 601	Difference vs. placebo (95% CI)	Placebo, n = 201	Tralokinumab Q2W, n = 591	Difference vs. placebo (95% CI)	
Primary endpoints							
IGA score of 0 or 1 at week 16, $n (\%)^{a}$	14/197 (7.1)	95/601 (15.8)	$8.6 (4.1 \text{ to } 13.1);^{b}$ P = 0.002^{c}	22/201 (10·9)	131/591 (22·2)	11.1 (5.8 to 16.4); ^b $P < 0.001^{c}$	
EASI 75 at week 16, n (%) ^a	25/197 (12·7)	150/601 (25·0)	$12.1 (6.5 \text{ to } 17.7);^{\text{b}}$ $P < 0.001^{\text{c}}$	23/201 (11·4)	196/591 (33·2)	21.6 (15.8 to 27.3); ^b $P < 0.001^{c}$	
Key secondary endpoint	ts						
Improvement in weekly average of worst daily pruritus NRS \geq 4 points from baseline to week 16, n/N (%) ^{a,d}	20/194 (10·3)	119/594 (20·0)	$9.7 (4.4 \text{ to } 15.0);^{\text{b}}$ P = 0.002^{c}	19/200 (9.5)	144/575 (25·0)	15.6 (10.3 to 20.9); ^b $P < 0.001^{c}$	
Adjusted mean change (SE) from baseline in SCORAD score at week 16 ^e	-14.7 (1.80)	-25.2 (0.94)	-10.4 (-14.4) to -6.5 ; P < 0.001	-14.0 (1.79)	-28.1 (0.92)	-14.0 (-18.0) to -10.1 ; P < 0.001	
Adjusted mean change (SE) from baseline in DLQI score at week 16 ^e	-5.0 (0.59)	-7.1 (0.31)	-2.1 (-3.4) to -0.8 ; P = 0.002	-4.9 (0.60)	-8.8 (0.30)	-3.9 (-5.2) to -2.6 ; P < 0.001	

CI, confidence interval; DLQI, Dermatology Life Quality Index; EASI, Eczema Area and Severity Index; IGA, Investigator's Global Assessment; NRS, numerical rating scale; Q2W, every 2 weeks; SCORAD, SCORing Atopic Dermatitis; SE, standard error. ^aPatients who received rescue medication were considered nonresponders. Patients with missing data at week 16 were imputed as nonresponders. ^bMantel–Haenszel risk difference, stratified by region and baseline IGA. ^cCochran–Mantel–Haenszel test, stratified by region and baseline IGA. ^dBased on patients in the full analysis set with a baseline pruritus NRS weekly average ≥ 4 . ^eData collected after permanent discontinuation of the investigational medicinal product or initiation of rescue medication not included. Repeated-measurements model on postbaseline data: change = (Treatment × Week) + (Baseline measure × Week) + Region + Baseline IGA.



Figure 1 Achievement of (a) Investigator's Global Assessment (IGA) score of 0 or 1 and (b) \geq 75% improvement in Eczema Area and Severity Index (EASI 75) in the 16-week initial treatment period in ECZTRA 1 and ECZTRA 2. Patients who received rescue medication were considered nonresponders. Patients with missing data at week 16 were imputed as nonresponders. The Cochran–Mantel–Haenszel test was used, stratified by region and baseline IGA. *P < 0.05 vs. placebo, **P < 0.01 vs. placebo, ***P < 0.001 vs. placebo, $^{+P} = 0.002$ vs. placebo. Q2W, every 2 weeks.

and $-8\cdot8$ vs. $-4\cdot9$ in ECZTRA 2 (difference $-3\cdot9\%$, 95% CI $-5\cdot2$ to $-2\cdot6$; P < 0.001; Table 2; and Tables S8 and S9; see Supporting Information).

Significant improvements in favour of tralokinumab were observed using the primary and alternative analytical approaches (Tables S10 and S11; see Supporting Information).

Maintenance outcomes at week 52

In ECZTRA 1 and ECZTRA 2, 185 and 227 patients were rerandomized 2 : 2 : 1 to continue tralokinumab Q2W, to reduce the dosing frequency of tralokinumab to Q4W or to switch to placebo Q2W (Figure S3; see Supporting Information). In patients who achieved IGA 0 or 1 with tralokinumab at week 16, IGA 0 or 1 was maintained at week 52 without rescue medication (including TCS) in 51% with continued tralokinumab Q2W vs. 47% with tralokinumab Q2W to placebo (difference 6.0%, 95% CI -21.8 to 33.7; P = 0.68) in ECZTRA 1 and in 59% with continued tralokinumab Q2W vs. 25% with tralokinumab Q2W to placebo (difference 34.1%, 95% CI 13.4-54.9; P = 0.004) in ECZTRA 2 (Figure 2a and Table 3).

For the other maintenance-phase endpoints in the testing hierarchy, EASI 75 at week 52 was maintained by 60% with continued tralokinumab Q2W vs. 33% with tralokinumab Q2W to placebo (difference 21·2%, 95% CI -0.2 to 42·6; P = 0.056) in ECZTRA 1 and 56% with continued tralokinumab Q2W vs. 21% with tralokinumab Q2W to placebo (difference 33·7%, 95% CI 17·3–50·0; P < 0·001) in ECZTRA 2 (Figure 2b and Table 3).

The percentage of patients who maintained IGA 0 or 1 at week 52 when rerandomized to tralokinumab Q4W was 39% in ECZTRA 1 (difference vs. tralokinumab Q2W to placebo -9.5%, 95% CI -37.1 to 18.0; P = 0.50) and 45% in ECZTRA 2 (difference 19.9%, 95% CI -1.2 to 40.9; P = 0.084) (Figure 2a and Table 3). For EASI 75 the proportion with tralokinumab Q2W to Q4W was 49% (difference vs.



Figure 2 Maintenance of (a) Investigator's Global Assessment (IGA) score of 0 or 1 and (b) \geq 75% improvement in Eczema Area and Severity Index (EASI 75) clinical response at week 52 in ECZTRA 1 and ECZTRA 2. Patients who, after week 16, received rescue medication or were transferred to open-label treatment were considered nonresponders at week 52. Missing values were imputed as nonresponse. The Cochran– Mantel–Haenszel test was used, stratified by region. Maintenance of IGA 0 or 1 was assessed in patients achieving the week 16 primary outcome of IGA score of 0 or 1 without use of rescue medication after initial randomization to tralokinumab. EASI 75 was assessed in patients achieving the week 16 primary outcome of EASI 75 without use of rescue medication after initial randomization to tralokinumab. **P < 0.01, vs. placebo ***P < 0.001 vs. placebo. Q2W, every 2 weeks; Q4W, every 4 weeks.

tralokinumab Q2W to placebo 11.7%, 95% CI -8.7 to 32.0; P = 0.27) in ECZTRA 1 and 51% (difference 30.0%, 95% CI 13.7-46.4; P = 0.001) in ECZTRA 2 (Figure 2b and Table 3).

Additional efficacy analyses

Changes from baseline in SCORAD, worst daily pruritus NRS, POEM and DLQI were greater with tralokinumab vs. placebo throughout the initial 16 weeks, and separation between tralokinumab and placebo was observed from week 1 onwards for weekly average worst daily pruritus NRS (P < 0.001) and from week 2 onwards for SCORAD (P < 0.001) and DLQI (P < 0.01) (Table 4; and Figure S5; see Supporting Information).

A greater reduction in weekly average eczema-related sleep interference was observed with tralokinumab vs. placebo from week 1 onwards (P < 0.01) in both studies (Figure S5; see Supporting Information). The difference between tralokinumab and placebo in the adjusted mean change from baseline in eczema-related sleep interference at week 16 was -0.7 in ECZTRA 1 and -1.4 in ECZTRA 2 (Table 4).

In both studies, more patients achieved EASI 50 and EASI 90 with tralokinumab vs. placebo at week 16 (Table 5; and Figure S6; see Supporting Information) and EASI 50 at each scheduled assessment (P < 0.01) from week 2. EASI 90 response was greater than with placebo from week 4 to week 16, with a separation between arms (P < 0.01) from week 6. A greater percentage change in EASI was observed with tralokinumab vs. placebo at week 16, with a separation between arms (P < 0.001) from week 2 onwards (Table 5, Figure 3; and Figure S7; see Supporting Information).

Safety

The incidence of AEs was comparable between tralokinumab and placebo in the initial treatment period of both studies (Table 6). The majority of AEs were nonserious and mild or

	ECZTRA 1			ECZTRA 2			
Rerandomization at week 16	Tralokinumab Q2W to placebo, n = 35	Tralokinumab Q2W to Q2W, n = 68	Tralokinumab Q2W to Q4W, n = 76	Tralokinumab Q2W to placebo, n = 46	Tralokinumab Q2W to Q2W, n = 91	Tralokinumab Q2W to Q4W, n = 89	
IGA score of 0 or 1 at week 52, n/N (%) ^{a,b}	9/19 (47)	20/39 (51)	14/36 (39)	7/28 (25)	32/54 (59)	22/49 (45)	
Difference in percentage vs. placebo (95% CI) ^c	10 (20 (22)	6.0 (-21.8 to 33.7); $P = 0.68^{d}$	-9.5 (-37.1 to 18.0); P = 0.50 ^d		34.1 (13.4 to 54.9); $P = 0.004^{d}$	19.9 (-1.2 to 40.9); P = 0.084 ^d	
EASI /5 at week 52, n/N (%) ^{b,e}	10/30 (33)	28/4/ (60)	28/5/ (49)	9742 (21)	43/77 (56)	38/74 (51)	
Difference in percentage vs. placebo (95% CI) ^c		21.2 (-0.2 to 42.6); P = 0.056 ^d	11.7 (-8.7 to); $P = 0.27^{d}$		33.7 (17.3 to 50.0); $P < 0.001^{d}$	30.0 (13.7 to 46.4); $P = 0.001^{d}$	

Table 3	Efficac	y outcomes a	t week 5	2 following	the maintenance	treatment	period:	maintenance	analys	is set
		/								

CI, confidence interval; EASI, Eczema Area and Severity Index; IGA, Investigator's Global Assessment; Q2W, every 2 weeks; Q4W, every 4 weeks. ^aAmong patients with an IGA score of 0 or 1 at week 16 achieved without rescue medication after initial randomization to tralok-inumab. ^bPatients who received rescue medication or were transferred to open-label treatment were considered nonresponders. Missing data at week 52 were imputed as nonresponse. ^cMantel–Haenszel risk difference compared with placebo, stratified by region. ^dCochran–Mantel–Haenszel test, stratified by region. ^eAmong patients with \geq 75% improvement in EASI at week 16 achieved without rescue medication after initial randomization to tralokinumab.

Table 4 Efficacy outcomes for the initial treatment period: other endpoints, full analysis set

	ECZTRA 1	ECZTRA 1			ECZTRA 2			
	Placebo, n = 197	Tralokinumab Q2W, n = 601	Difference vs. placebo (95% CI)	Placebo, n = 201	Tralokinumab Q2W, n = 591	Difference vs. placebo (95% CI)		
Adjusted mean change (SE) from baseline in SCORAD score at week 2 ^a	-5.0 (0.92)	-10.6 (0.53)	-5.6 (-7.7) to -3.5 ; P < 0.001	-3.9 (0.84)	-10.8 (0.49)	-6.9 (-8.8) to -5.0 ; P < 0.001		
Adjusted mean change (SE) from baseline in weekly average of worst daily pruritus NRS at week 1 ^a	-0.2 (0.07)	-0.7 (0.04)	-0.4 (-0.6) to -0.3 ; P < 0.001	-0.3 (0.08)	-0.7 (0.05)	-0.4 (-0.6) to -0.2 ; P < 0.001		
Adjusted mean change (SE) from baseline in DLQI at week 2 ^a	-2.5 (0.39)	-4.4 (0.22)	-2.0 (-2.8) to -1.1 ; P < 0.001	-2.2 (0.39)	-4.7 (0.23)	-2.5 (-3.4) to -1.7 ; P < 0.001		
Adjusted mean change (SE) from baseline in weekly average of eczema-related sleep interference NRS at week 16 ^a	-1.9 (0.23)	-2.6 (0.12)	-0.7 (-1.2) to -0.2 ; P = 0.007	-1.5 (0.22)	-2.9 (0.12)	-1.4 (-1.9) to -0.9 ; P < 0.001		
Adjusted mean change (SE) from baseline in POEM score at week 16^a	-3.0 (0.66)	-7.6 (0.35)	-4.6 (-6.0) to -3.1 ; P < 0.001	-3.7 (0.66)	-8.8 (0.33)	-5.1 (-6.5) to -3.6 ; P < 0.001		

CI, confidence interval; DLQI, Dermatology Life Quality Index; IGA, Investigator's Global Assessment; NRS, numerical rating scale; POEM, Patient-Oriented Eczema Measure; Q2W, every 2 weeks; SCORAD, SCORing Atopic Dermatitis; SE, standard error. ^aData collected after permanent discontinuation of the investigational medicinal product or initiation of rescue medication not included. Repeated-measurements model on postbaseline data: change = (Treatment \times Week) + (Baseline measure \times Week) + Region + Baseline IGA.

moderate in severity, with most resolved or resolving by the end of the treatment period; few patients had AEs leading to permanent discontinuation. Of the most frequently reported AEs (\geq 5% in any treatment group), upper respiratory tract infection (mainly reported as common cold) and conjunctivitis occurred more frequently with tralokinumab than with placebo, and dermatitis atopic and skin infection occurred more frequently with placebo (Table 6). The frequency of serious AEs (SAEs) was low and comparable between treatment groups in the initial treatment period of both studies; the majority of patients recovered from the events.

Table 5	Efficacy	outcomes	for the	initial	treatment	period:	additional	secondary	endpoints,	full a	analysis se	t
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	ECZTRA 1	l		ECZTRA 2			
	Placebo, n = 197	Tralokinumab Q2W, n = 601	Difference vs. placebo (95% CI)	Placebo, n = 201	Tralokinumab Q2W, n = 591	Difference vs. placebo (95% CI)	
Reduction of weekly average of worst daily pruritus NRS ≥ 3 at week 16, n/N (%) ^{a,b}	28/195 (14·4)	177/597 (29.6)	15.2 (9.2 to 21.3); ^c $P < 0.001^{d}$	28/200 (14·0)	199/583 (34.1)	20.1 (13.9 to 26.2); ^c $P < 0.001^{d}$	
Adjusted mean change (SE) from baseline in weekly average of worst daily pruritus NRS at week 16 ^e	-1·7 (0·21)	-2.6 (0.11)	-0.9 (-1.4) to -0.4 ; P < 0.001	-1.6 (0.21)	-2.9 (0.11)	-1.3 (-1.7) to -0.8 ; P < 0.001	
SCORAD score of 75 at week 16, $n/N \left(\%\right)^a$	6/197 (3·0)	53/601 (8.8)	5.7 (2.5 to 8.9); ^c $P = 0.007^{d}$	7/201 (3·5)	68/591 (11.5)	8.0 (4.4 to 11.6); ^c $P < 0.001^{d}$	
SCORAD score of 50 at week 16, $n/N \ (\%)^a$	23/197 (11·7)	156/601 (26.0)	14.1 (8.6 to 19.6); ^c $P < 0.001^{d}$	29/201 (14·4)	198/591 (33.5)	18.9 (12.8 to 25.1); ^c $P < 0.001^{d}$	
DLQI score reduction ≥ 4 at week 16, n/N (%) ^{a,f}	60/190 (31·6)	258/578 (44.6)	13.0 (5.4 to 20.5); ^c $P = 0.001^{d}$	54/198 (27·3)	325/577 (56·3)	28.9 (21.4 to 36.3); ^c $P < 0.001^{d}$	
Adjusted mean change (SE) from baseline in EASI at week 16^{e}	-9·0 (1·05)	-15.5 (0.55)	-6.4 (-8.8) to -4.1 ; P < 0.001	-7·0 (1·06)	-16.9 (0.55)	-9.9 (-12.2) to -7.5 ; P < 0.001	
EASI 50 at week 16, n/N (%) ^a	42/197 (21·3)	250/601 (41.6)	20.1 (13.3 to 26.8); ^c $P < 0.001^{d}$	41/201 (20·4)	295/591 (49·9)	29.3 (22.5 to 36.1); ^c $P < 0.001^{d}$	
EASI 90 at week 16, n/N (%) ^a	8/197 (4·1)	87/601 (14.5)	10.3 (6.4 to 14.1); ^c $P < 0.001^{d}$	11/201 (5·5)	108/591 (18·3)	12.7 (8.3 to 17.0); ^c $P < 0.001^{d}$	

CI, confidence interval; DLQI, Dermatology Life Quality Index; EASI, Eczema Area and Severity Index; IGA, Investigator's Global Assessment; NRS, numerical rating scale; Q2W, every 2 weeks; SCORAD, SCORing Atopic Dermatitis; SE, standard error. ^aPatients who received rescue medication were considered nonresponders. Patients with missing data at week 16 were imputed as nonresponders. ^bBased on patients in the full analysis set with a baseline pruritus NRS weekly average \geq 3. ^cMantel–Haenszel risk difference, stratified by region and baseline IGA. ^dCochran–Mantel–Haenszel test, stratified by region and baseline IGA. ^eData collected after permanent discontinuation of the investigational medicinal product or initiation of rescue medication not included. Repeated-measurements model on postbaseline data: change = (Treatment × Week) + (Baseline measure × Week) + Region + Baseline IGA. ^fAnalysis includes only patients with baseline DLQI \geq 4.

Conjunctivitis as an AE of special interest (AESI) occurred with greater frequency with tralokinumab vs. placebo (Table 6). Most cases of conjunctivitis were mild and resolved by the end of the treatment period; one case led to treatment discontinuation. Tralokinumab-treated patients had lower rates of eczema herpeticum vs. placebo in both studies, and lower rates of skin infections requiring systemic treatment were reported as an AESI for tralokinumab vs. placebo in ECZTRA 2 (Table 6).

Overall, in the maintenance treatment period, AEs were detected at a lower rate than with tralokinumab Q2W in the initial treatment period, and the pattern of events was comparable with that in the initial treatment period. AEs were more frequently reported in the continued tralokinumab Q2W group than in the tralokinumab Q2W to Q4W group, and a low number of SAEs and AEs leading to permanent discontinuation of tralokinumab was observed (Table S12; see Supporting Information).

No marked differences in SAEs were observed between the treatment groups within each treatment period and between

the treatment periods, and there was no clustering with respect to specific system organ class or event types.

In ECZTRA 1, the reduction in S. aureus colonization on lesional skin, from baseline to week 16, as assessed by quantitative polymerase chain reaction, was more than 10 times greater with tralokinumab vs. placebo: the median reduced from 969 to 22 gene copies per cm² with tralokinumab vs. 649 to 238 gene copies per cm² with placebo.

Neutralizing antidrug antibodies were detected in three and eight tralokinumab-treated patients in ECZTRA 1 and ECZTRA 2, respectively. Based on examination of tralokinumab concentrations, antidrug antibody titre levels, AEs, and IGA and EASI scores across the trials, it was considered that the presence of neutralizing antibodies did not have an impact on the efficacy and safety of tralokinumab for any of the patients.

There were no noteworthy differences between treatment groups in laboratory values, vital signs or electrocardiographic assessments. More patients treated with tralokinumab experienced eosinophilia during the initial treatment period, but the



Figure 3 Example of improvement in Eczema Area and Severity Index (EASI) from baseline to week 16 in a patient receiving tralokinumab. IGA, Investigator's Global Assessment; NRS, numerical rating scale. [Colour figure can be viewed at wileyonlinelibrary.com]

mean eosinophil levels returned to baseline values during the maintenance period, and the safety profile of patients with eosinophilia (> 1.5×10^9 L) was comparable with that in the total trial population.

Discussion

The efficacy and safety of tralokinumab monotherapy for the treatment of moderate-to-severe AD were examined over 1 year in two large, multinational phase III studies of identical design. Tralokinumab 300 mg Q2W demonstrated superiority over placebo during 16 weeks of treatment across multiple outcome measures reflecting the signs and symptoms of AD. Improvements in itch and sleep scores were apparent as early as week 1 for patients treated with tralokinumab vs. placebo, and the majority of patients who achieved response criteria for improvement in the extent and severity of AD at week 16 maintained response at week 52 without need for rescue medication, including TCS.

The studies were consistent in demonstrating the effectiveness of tralokinumab monotherapy, but there were some differences in the level of response vs. placebo for some endpoints. In particular, there was a greater difference between tralokinumab and placebo for the primary endpoints in ECZTRA 2 compared with ECZTRA 1. One potential explanation is the greater use of rescue medication in tralokinumab-treated patients in ECZTRA 1 (35.8%) compared with ECZTRA 2 (22.8%). The criteria surrounding use of rescue medication were strict; a single use of TCS was registered as treatment failure following the primary analysis approach. This does not reflect real-world use of biologic agents, which are initiated as add-on therapy to TCS for active lesions in 80– 90% of patients,^{18,19} and therefore the utility of these findings in daily practice is difficult to assess. The ECZTRA 3 study (NCT03363854), of tralokinumab in combination with TCS, is more reflective of the likely clinical use in daily practice.²⁰

More than 50% of patients who achieved clinical responses at week 16 with tralokinumab Q2W maintained that response through to week 52 without any rescue medication, including TCS, and 39-51% of patients maintained response when receiving tralokinumab Q4W. Unexpectedly, a proportion (21-47%) of tralokinumab responders at week 16 who were rerandomized to placebo maintained responses at week 52. Retained response over 36 weeks without active maintenance treatment or TCS suggests that tralokinumab could induce remission of AD and skin normalization for some patients. It has previously been shown that IL-13 expression is much lower in nonlesional than lesional AD skin.9,10,21 It is therefore possible that following a period of clear or almost clear skin achieved with tralokinumab Q2W, IL-13-mediated inflammation in the skin may have been extinguished, altering the natural disease course. Further studies are needed to identify whether some patients can experience a true remission or altered natural course upon stopping therapy. There were also patients who could maintain good disease control with Q4W dosing of tralokinumab. These observations suggest the possibility of less frequent maintenance dosing in some patients initially treated with tralokinumab Q2W.

Support for reversal of IL-13-associated skin abnormalities with tralokinumab was provided by the observed greater reduction in skin colonization with S. aureus, which was consistent with observations in the phase II study.¹⁵ In addition to decreasing S. aureus colonization, fewer skin infections requiring systemic treatment and a lower frequency of eczema

Table 6 Summary of adverse events (AEs) and AEs of special interest (AESIs) in the 16-week initial treatment period for the safety analysis set

	ECZTRA 1		ECZTRA 2		
	Placebo, n = 196, PYE = 57·13	Tralokinumab Q2W, $n = 602$, PYE = 177.6	Placebo, n = 200, PYE = 57.35	Tralokinumab Q2W, n = 592, PYE = 176·9	
AEs					
Total number of AEs	491	1482	408	997	
Total number of SAEs	11	24	6	10	
Patients with AEs					
$\geq 1 \text{ AE}$	151 (77.0)	460 (76.4)	132 (66.0)	364 (61.5)	
≥ 1 SAE	8 (4.1)	23 (3.8)	5 (2.5)	10 (1.7)	
Severity					
Mild	111 (56.6)	385 (64.0)	93 (46.5)	288 (48.6)	
Moderate	98 (50·0)	241 (40.0)	84 (42.0)	168 (28.4)	
Severe	16 (8.2)	41 (6.8)	16 (8.0)	24 (4.1)	
Leading to permanent discontinuation of IMP	8 (4.1)	20 (3.3)	3 (1.5)	9 (1.5)	
Outcome					
Not recovered/not resolved	35 (17.9)	106 (17.6)	25 (12.5)	61 (10.3)	
Recovering/resolving	7 (3.6)	36 (6.0)	15 (7.5)	20 (3.4)	
Recovered/resolved	139 (70.9)	429 (71.3)	125 (62.5)	340 (57.4)	
Recovered/resolved with sequelae	0	6 (1.0)	2 (1.0)	9 (1.5)	
Frequent AEs ^a					
Dermatitis atopic	75 (38.3)	156 (25.9)	67 (33.5)	98 (16.6)	
Viral upper respiratory tract infection	41 (20.9)	139 (23.1)	17 (8.5)	49 (8.3)	
Upper respiratory tract infection	2 (1.0)	9 (1.5)	17 (8.5)	59 (10.0)	
Conjunctivitis	4 (2.0)	43 (7.1)	3 (1.5)	18 (3.0)	
Skin infection	3 (1.5)	6 (1.0)	11 (5.5)	12 (2.0)	
Pruritus	10 (5.1)	32 (5.3)	5 (2.5)	12 (2.0)	
Headache	10 (5.1)	28 (4.7)	6 (3.0)	16 (2.7)	
AESI – eye disorders	7 (3.6)	62 (10.3)	6 (3.0)	33 (5.6)	
AESI Conjunctivitis ^b	7 (3.6)	60 (10.0)	5 (2.5)	31 (5.2)	
AESI Keratoconjunctivitis	0	1 (0.2)	0	2 (0.3)	
AESI Keratitis	0	3 (0.5)	1 (0.5)	1 (0.2)	
AESI – skin infections requiring systemic treatment	4 (2.0)	13 (2.2)	22 (11.0)	21 (3.5)	
AESI – eczema herpeticum	2 (1.0)	3 (0.5)	5 (2.5)	2 (0.3)	
AESI – malignancies diagnosed after randomization	0	0	0	1 (0.2)	

The data are presented as n or n (%). IMP, investigational medicinal product; PYE, patient-years of exposure; Q2W, every 2 weeks; SAE, serious AE. ^aPreferred term according to the Medical Dictionary for Regulatory Activities (MedDRA) 20-0, occurring in \geq 5% of patients in any randomized group. ^bIncludes the preferred terms conjunctivitis, conjunctivitis bacterial, conjunctivitis viral and conjunctivitis allergic.

herpeticum were seen with tralokinumab. These findings may be due to tralokinumab's effects on improving skin barrier integrity. Previous studies have also linked upregulation of IL-13 to decreased expression of antimicrobial genes;^{22,23} thus, tralokinumab could be enhancing skin antimicrobial responses in this manner. Analysis of biomarkers from skin biopsies and serum biomarkers collected in ECZTRA 1 may provide further insights into the effects of tralokinumab on the skin microbiome and epidermal changes at a molecular level.

Tralokinumab was well tolerated, with an overall frequency and severity of AEs comparable with placebo over 52 weeks, consistently with those observed in the phase II trial.¹⁵ Incidences of eye disorders were collected as AESIs. Conjunctivitis occurred in $\leq 10\%$ of patients receiving tralokinumab in the initial treatment period and in < 9% in any tralokinumab treatment arm in the maintenance periods. Almost all cases of conjunctivitis were mild or moderate, the majority recovered, and one case led to withdrawal. This side-effect has been observed in other studies of IL-4 or IL-13 inhibitors in AD and the aetiology remains unknown.

Consistently with other trials in AD, a limitation of these studies was the requirement to ensure adequate washout of existing treatments, particularly in a patient population with significant disease and high levels of prior medication use. The protocol-required 2-week washout period, during which no TCS use was allowed, may have been long enough to exacerbate AD in some patients, leading to use of rescue TCS early in the studies. In fact, approximately half of the patients who used rescue medications did so within the first 4 weeks of the trials.

Most currently available treatment options for AD were not designed to selectively target disease-specific pathways that have been established in the pathogenesis of the disease, and there is a need for additional long-term treatment options for patients with moderate-to-severe AD. These are the first phase III studies to demonstrate that specific IL-13 neutralization with tralokinumab monotherapy is efficacious for the treatment of AD, providing further evidence that IL-13 is a key cytokine in the pathogenesis of this disease.

In conclusion, tralokinumab monotherapy was superior to placebo at 16 weeks of treatment and was well tolerated up to 52 weeks of treatment in a large cohort of patients with moderate-to-severe AD.

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Appendix

Conflicts of interest

A.W. reports honoraria for conduct of the ECZTRA 1 and ECZ-TRA 2 trials to Ludwig Maximilian University of Munich from LEO Pharma A/S during the conduct of the study; personal fees from AbbVie, Chugai, Galderma, LEO Pharma, Lilly, Med-Immune, Novartis, Pfizer, Regeneron and Sanofi-Aventis; and grants from LEO Pharma outside the submitted work. A.B. reports personal fees for consulting and clinical study fees to Oregon Medical Research Center from LEO Pharma during the conduct of the study; personal fees for consulting and clinical study fees from AbbVie, Dermira, Incyte, Lilly, Pfizer and Regeneron; and personal fees for consulting from Sanofi outside the submitted work. E.G.-Y. declares they have no conflicts of interest. M.W. reports honoraria for advisory boards and lecture activities from ALK-AbbVie Deutschland GmbH & Co. KG, Abelló Arzneimittel GmbH, Actelion Pharmaceuticals Deutschland GmbH, Aimmune Therapeutics UK Limited, Allergopharma GmbH & Co. KG, Bencard Allergie GmbH, Biotest AG, DBV Technologies S.A., HAL Allergie GmbH, LEO Pharma GmbH, Lilly Deutschland GmbH, Mylan Germany GmbH, Novartis AG, Regeneron, Sanofi-Aventis Deutschland GmbH and Stallergenes GmbH outside the submitted work. C.L. reports honoraria for speaking and consulting, and investigator fees from AbbVie, Amgen, Bausch, Galderma, LEO Pharma, Pfizer and Sanofi Genzyme outside the submitted work. J.-P.L. reports grants from LEO Pharma during the conduct of the study; grants from AbbVie, Amgen, Boehringer Ingelheim, Dermira, Janssen and Pfizer; and grants and personal fees from Celgene, Galderma, Lilly, Novartis and Sanofi outside the submitted work. L.S. reports investigator fees from Amgen, Blaze, BMS, Boehringer Ingelheim, LEO Pharma, Pfizer, Sanofi, Sun and UCB; and investigator fees and fees for advisory board participation from AbbVie, Galderma, Janssen, Lilly and Novartis outside the submitted work. N.K. reports personal fees from AbbVie, Janssen, Maruho, Mitsubishi Tanabe and Taiho; and grants and personal fees from Lilly Japan, LEO Pharma and Sanofi outside the submitted work. H.S. reports personal fees from Kyorin, Kyowa Kirin, LEO Pharma, Sanofi, Taiho and Tokiwa; grants and personal fees from Maruho and Mitsubishi Tanabe; and grants from Eisai and Torii outside the submitted work. Y.P. reports grants from LEO Pharma during the conduct of the study; grant funding as an investigator from AnaptysBio, Arcutis, Asana, AstraZeneca, Baxalta, Baxter, Boehringer Ingelheim, Bond Avillion, Bristol Myers Squibb, Celgene, Dermira, Devonian, Galderma, Genentech, GlaxoSmithKline, Incyte, LEO Pharma, MedImmune, Merck, Novartis, Pfizer, Regeneron, Roche, Serono and Takeda; and grants and honoraria for services as an investigator, speaker and member of advisory boards from AbbVie, Amgen, Bausch, Janssen-Ortho and UCB. A.L. reports investigator fees paid to the Medical University of Łódź from Regeneron and personal fees and investigator fees paid to the Medical University of Łódź from AbbVie, LEO Pharma, Novartis and Pfizer outside the submitted work. L.K. reports personal fees as a speaker from Sanofi, grants from Arena and grants and personal fees as a consultant or speaker from Abb-Vie, Arcutis, Dermavant, Dermira, Incyte, LEO Pharma, Lilly, Novartis, Ortho Dermatologics, Pfizer and Regeneron outside the submitted work. S.H.C. reports personal fees and clinical study fees to the Catholic University of Korea from LEO Pharma during the conduct of the study and personal fees and clinical study fees to the Catholic University of Korea from AbbVie, Pfizer, Regeneron and Sanofi outside the submitted work. P.H. declares they have no conflicts of interest. M.J.C. has served as an investigator or consultant for AbbVie, Astellas, Boots, Dermavant, Galapagos, Galderma, Hyphens, Johnson & Johnson, LEO Pharma, L'Oréal, Menlo, Novartis, Oxagen, Pfizer, Procter & Gamble, Reckitt Benckiser, Regeneron and Sanofi Genzyme. K.P. reports grants and personal fees for participation in advisory boards from AbbVie and Galderma and personal fees for participation in advisory boards from Almirall, Janssen, LEO Pharma, Lilly, Novartis, Pierre Fabre, Sanofi and Sun Pharma outside the submitted work. L.A.S., B.B., A.K., T.N.J. and M.L.Ø. are employees of LEO Pharma A/S. E.L.S. reports grants from Celgene, Galderma, Merck, Novartis and Tioga; personal fees from Boehringer Ingelheim, Dermavant, Dermira, Forté Bio, Incyte, Menlo, Ortho Dermatologics, Pierre Fabre, Sanofi and Valeant; and grants and personal fees from AbbVie, Kyowa Hakko Kirin, LEO Pharma, Lilly, MedImmune, Pfizer and Regeneron outside the submitted work.

Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher's website:

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Appendix S2 Additional study methods.

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Figure S2 Testing hierarchy.

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