Efficacy and Safety of Tildrakizumab in Older Patients: Pooled Analyses of Two Randomized Phase III Clinical Trials (reSURFACE 1 and reSURFACE 2) Through 244 Weeks

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The evidence on treating older patients with psoriasis with modern biologics is scarce. This study compared the efficacy and safety of tildrakizumab among younger and older patients with psoriasis (<65/≥65 years) in a post hoc analysis of 2 phase III trials (reSURFACE1/2, n = 1,862). Tildrakizumab 100 mg/200 mg was administered at weeks 0/4/every 12 weeks thereafter. At week 28, patients with ≥75% improvement in baseline Psoriasis Area and Severity Index (PASI75) in reSURFACE1 were re-randomized to the same tildrakizumab dose or placebo; in reSURFACE2, PASI75 responders to 200 mg were re-randomized to tildrakizumab 100 mg or 200 mg; PASI75 responders to 100 mg maintained their dose. At weeks 64/52 (reSURFACE1/2), PASI50 responders entered an extension period (weeks 256/244). Outcomes were proportion of patients with PASI<3, Dermatology Life Quality Index (DLQI) 0/1, comorbidities, comedication, and side-effects. The proportion of patients with a PASI<3 was similar and maintained (tildrakizumab 100 mg and 200 mg, week 244: 83.3% and 84.1%/92.3% and 100.0%); DLQI 0/1 proportions at week 52 were 66.8% and 72.0%/68.3% and 81.3%. Comorbidity and comedication were more common in older patients. The safety profile of tildrakizumab appeared favourable in both groups. Tildrakizumab in patients ≥65 years appears effective and safe in long-term psoriasis management. These findings might assist treatment selection and overcome treatment reluctance.

Key words: adverse drug events; clinical efficacy; clinical trials, Phase III as topic; older adults, frail; tildrakizumab.

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Psoriasis is a chronic inflammatory disease with a worldwide prevalence rate of approximately 2–3% (1). Older patients (age≥65 years) represent an increasing proportion of patients with psoriasis and 15% of them have moderate to severe disease (2). With a steadily aging population (3), physicians are faced with an increasing number of older patients with psoriasis. However, optimal treatment selection might be difficult due to the presence of comorbidities (4), comedication (5, 6), and adverse events (AEs) (7), which also influence patient treatment preferences (8).

Generally, biologics have demonstrated even better efficacy than conventional systemics (9, 10), with lower rates of AE than conventional systemics (10). However, the elderly population is often excluded from clinical trials based on age or on age-related factors (e.g. comorbidities) (11), and representation of older patients in the available trial literature is low. However, a recent registry reported that discontinuation of biologics due to AEs did not occur more frequently in older compared with younger patients (12). Older patients tended to have more serious infections, non-melanoma skin cancer (NMSC) and malignancies than younger patients, possibly due to the ageing process and more extensive duration of disease (9, 13).

In the case of tildrakizumab (TIL), specifically, there is almost no evidence available regarding older patients, and the first report in clinical practice was provided by Ruggiero et al. (14). This study included only 6 older patients, but they reported similar results to those of randomized clinical trials (15). Although biologics seem to be relatively safe, this limited evidence-based management could trigger treatment reluctance to prescribe (newer) biologics in older patients for fear of lower efficacy or worse tolerability. Thus, more robust, comprehensive data regarding biologics in older patients are needed.

SIGNIFICANCE

This study compared the efficacy and safety of tildrakizumab among younger and older adults with psoriasis (<65/≥65 years). High and similar proportions of patients in both groups achieved improvement of skin lesions and disease-related quality of life during the first year, which was maintained up to 5 years. The most frequent side-effect was nasopharyngitis. Although older patients presented more comorbidities and comedication, they showed a similar and favourable safety profile, demonstrating that tildrakizumab appears to be a good and safe treatment for psoriasis in both older and younger patients with psoriasis.
The aim of the present study is to compare the pooled efficacy and safety of TIL 100 mg and 200 mg for 244 weeks among younger and older patients from the 2 pivotal reSURFACE trials (reSURFACE 1 and reSURFACE 2) (16), including long-term extension periods (17, 18).

MATERIALS AND METHODS

This is a post hoc pooled analysis of 2 3-part, randomized, double-blind, placebo-controlled, parallel-group, phase III trials (reSURFACE 1 and reSURFACE 2, ClinicalTrials.gov NCT01722331 and NCT01729754) that evaluated the efficacy and safety of TIL in patients with moderate to severe chronic plaque psoriasis for up to 5 years (17, 18). reSURFACE 2 included etanercept as an active comparator (16). reSURFACE 1 was conducted from 10 December 2012 to 28 October 2015. reSURFACE 2 was conducted from 12 February 2013 to 28 September 2015.

Main interventions

The main inclusion and exclusion criteria at baseline were similar between trials. Baseline study inclusion and exclusion criteria, patient characteristics, treatment, and methodology of these 2 pivotal clinical trials have been reported previously (16, 18). A total of 1,862 patients ≥ 18 years with moderate to severe chronic plaque psoriasis diagnosed ≥ 6 months prior to enrolment, with a body surface area ≥ 10%, a Physician’s Global Assessment ≥ 3 and a Psoriasis Area and Severity Index (PASI) ≥ 12, were included (reSURFACE 1, n = 772; reSURFACE 2, n = 1,090) (16). In reSURFACE 1, patients were randomized to TIL 100 mg, 200 mg, placebo or etanercept 50 mg (2:2:1:2). Tildrakizumab was administered at weeks 0, 4 and every 12 weeks afterwards. Responders were defined as patients with ≥75% improvement in baseline PASI (PASI75). At week 28, PASI75 responders in reSURFACE 1 were re-randomized to continue with the same TIL dose or to receive placebo; in reSURFACE 2, PASI75 responders to TIL 200 mg were re-randomized to TIL 100 mg or 200 mg, while PASI75 responders to TIL 100 mg maintained the same dose. At week 64 (reSURFACE 1) or week 52 (reSURFACE 2), patients with ≥50% improvement from baseline PASI score entered an optional 192-week extension period, until week 256 (reSURFACE 1) or week 244 (reSURFACE 2) (17, 18).

Both reSURFACE trials were conducted in accordance with Good Clinical Practice guidelines and the principles of the Declaration of Helsinki 1964, and its successive amendments. The study protocols received local institutional review board or ethics committee approvals. All patients gave informed consent to participate in the trials.

Main outcome measures

Medical history, including comorbidities and comedications, for each age group were summarized with descriptive statistics. Main efficacy outcomes were defined as the proportion of patients achieving absolute PASI < 3 over 5 years of treatment; that is, at weeks 28, 52 and 244, and Dermatology Life Quality Index (DLQI)/DLQI-Relevant (DLQI-R) 0/1 responses at weeks 28 and 52. In the DLQI, non-relevant responses (NRR) are scored as having no impact on patient quality of life, artificially improving patients’ DLQI scores (19). The new DLQI-R scoring avoids the bias of the NRR option by adjusting the total score for relevant items (20). Proportions of patients achieving absolute PASI < 5 and < 1 were also evaluated. Analyses were stratified by age groups and TIL dose, attending to the following groups: < 65 years and ≥ 65 years, TIL 100 mg and 200 mg.

Safety assessments included a description of AEs. Pre-specified treatment-emergent AEs (TE-AEs) comprised severe infections, malignancies, NMSC, melanoma, confirmed extended major adverse cardiovascular events (MACE), injection site reaction and drug-related hypersensitivity reactions (16, 18). Adverse events were assessed at all study visits and classified according to age and dose split. Preferred terms from the Medical Dictionary for Regulatory Activities (MedDRA) were assigned to the treatment dose that the patient was actively receiving when the AE occurred.

Statistical analysis

Current post hoc analyses focus on differences between age groups in demographics (including comorbidities and comedications), absolute PASI response, DLQI and DLQI-R response (by adjusting the total questionnaire score by the number of NNRs indicated by a patient) (21), and safety.

No formal hypothesis testing was performed for these post hoc analyses. All subjects randomized to TIL 100 mg and 200 mg who received at least 1 dose of study medication were included for week 28 efficacy analyses (TIL 100 mg: n = 593, 541 patients aged < 65 years and 52 patients aged ≥ 65 years).
years; TIL 200 mg: \( n = 597 \), 547 patients aged < 65 years and 50 patients aged ≥ 65 years) (Fig. 1). All patients who were responders (i.e. PASI75) at week 28 and who continued treatment with the same TIL dose were included for the long-term efficacy analyses (weeks 52 and 244) (TIL 100 mg: \( n = 329 \), 303 patients aged < 65 years and 26 patients aged ≥ 65 years; TIL 200 mg: \( n = 227 \), 211 patients aged < 65 years and 16 patients aged ≥ 65 years) (Fig. 1).

Efficacy analyses used an observed case approach. A multiple imputation approach (10 imputations) was used for missing data as sensitivity analyses for the PASI outcome, as described previously (18).

A mixed model was performed in the observed case population to evaluate possible changes in the absolute PASI at weeks 28 and 244 according to the following independent factors: age group, treatment, week, prior biological therapy for psoriasis, smoking habit, diabetes mellitus, history of psoriatic arthritis. We also included the age group x week interaction term into the model (looking at whether the behaviour of the variable under study in the 2 age groups is different over time, regardless of treatment). The model was covaried by baseline PASI and body mass index (BMI). This analysis was repeated with a multiple imputation approach. Analyses of comorbidities and comedications were performed in all randomized patients (\( n = 1,190 \)). Concomitant medications were collected over the 5-year study period. Safety analyses were performed in all patients who received at least 1 dose of study drug by treatment received (\( n = 1,800 \)). Safety data from week 0 to 5 years were pooled between reSURFACE 1 (up to week 256) and reSURFACE 2 (up to week 244) and presented for patients who received TIL during any part of the study with age of 65 years as the comparison threshold. Safety data are reported as number of events per 100 patient-years of exposure; exposure-adjusted incidence rates and 95% confidence intervals (95% CIs) were calculated as described previously (16, 18).

Analyses were performed with SAS software, version 9.4 (TSIM7) (© 2016 by SAS Institute INC, Cary, NC, USA), on the X64_10PRO platform for Windows, extension package SAS/STAT® software, version 15.2.

RESULT

Demographic and baseline characteristics

A summary of baseline characteristics is shown in Table I. The percentage of women was slightly higher in the older vs younger group and baseline DLQI score was significantly lower in older vs younger patients (\( p = 0.002 \)). The median (range) age for each age group was 44.0 (18.0–64.0) years in the younger, and 68.0 (65.0–82.0) years in the older group (Fig. S1). Both age groups showed no differences in previous experience with systemic biologic or non-biologic treatments.

The most common comorbidities in patients < 65 years vs ≥ 65 years were musculoskeletal and connective tissue disorders (26.4% vs 43.1%, \( p < 0.001 \)), metabolic and nutrition disorders (26.0% vs 56.9%, \( p < 0.001 \)), vascular disorders (24.4% vs 70.6%, \( p < 0.001 \)), and immune system disorders (22.3% vs 20.6%, \( p = 0.80 \)). The complete medical history, with comorbidities, by age group is shown in Table SI.

The proportions of patients < 65 years vs ≥ 65 years taking comedication at baseline were 56.3% vs 87.3%. Table SII shows comedication reported by patients over the 5-year study period.

Efficacy outcomes

PASI score. Fig. 2 shows the absolute mean PASI score over time by age group and TIL dose. Throughout the first 28 weeks, and especially between weeks 0 and 8, a decrease in the absolute mean PASI was observed in the 2 age groups and for each dose, from mean scores of ≥ 19 at baseline to means ≤ 6 from week 12. This trend was then maintained until week 244.

The proportion (95% CI) of TIL-treated patients aged < 65 vs ≥ 65 years achieving an absolute PASI < 3 for TIL 100 mg at week 28 was 66.4% (62.1–70.4%) vs 51.9% (37.6–66.0%). At week 244, it was 83.3% (77.8–88.0%) vs 92.3% (64.0–99.9%). The proportion (95% CI) of TIL-treated patients aged < 65 vs ≥ 65 years achieving an absolute PASI < 3 for TIL 200 mg at week 28 was 70.4% (66.3–74.2%) vs 58.3% (43.2–72.4%). At week 244, it was 84.1% (77.5–89.3%) vs 100.0% (75.3–100.0%) (comparison by age groups (combining TIL doses) week 244: \( p = 0.09 \)).

Table I. Demographic and baseline characteristics of the study population by age group and tildrakizumab dose

<table>
<thead>
<tr>
<th>Age group</th>
<th>&lt; 65 years</th>
<th>≥ 65 years</th>
</tr>
</thead>
<tbody>
<tr>
<td>100 mg</td>
<td>200 mg</td>
<td>Total</td>
</tr>
<tr>
<td>n (541)</td>
<td>n (547)</td>
<td>n (1,088)</td>
</tr>
<tr>
<td>Age, years, ( n ), mean (SD)</td>
<td>541, 43.2 (11.3)</td>
<td>547, 43.8 (11.9)</td>
</tr>
<tr>
<td>Female, %</td>
<td>161/541 (29.8)</td>
<td>142/547 (26.0)</td>
</tr>
<tr>
<td>BMI, kg/m², ( n ), mean (SD)</td>
<td>540, 30.0 (7.1)</td>
<td>547, 29.8 (7.5)</td>
</tr>
<tr>
<td>Weight, kg, ( n ), mean (SD)</td>
<td>541, 89.1 (23.1)</td>
<td>547, 89.1 (23.2)</td>
</tr>
<tr>
<td>PASI score, ( n ), mean (SD)</td>
<td>541, 20.2 (7.9)</td>
<td>547, 20.3 (8.0)</td>
</tr>
<tr>
<td>BSA, %, ( n ), mean (SD)</td>
<td>541, 31.7 (18.4)</td>
<td>542, 31.3 (17.4)</td>
</tr>
<tr>
<td>DLQI score, ( n ), mean (SD)</td>
<td>538, 14.0 (7.4)</td>
<td>541, 13.5 (7.0)</td>
</tr>
<tr>
<td>PsA, %</td>
<td>89/541 (16.5)</td>
<td>87/547 (15.9)</td>
</tr>
<tr>
<td>PGA ≥ 4, %</td>
<td>177/541 (32.7)</td>
<td>185/547 (34.0)</td>
</tr>
<tr>
<td>Previous experience with systemic 94/541 (17.4)</td>
<td>95/547 (17.4)</td>
<td>189/1,088 (17.4)</td>
</tr>
<tr>
<td>Previous experience with systemic 174/541 (32.1)</td>
<td>194/547 (35.5)</td>
<td>386/1,088 (33.8)</td>
</tr>
</tbody>
</table>

\( ^a n = 26 \) after W28 re-randomization; \( ^b n = 16 \) after W28 re-randomization; \( ^c \) Students’ t-tests are used for numerical variables and \( ^x ^2 \) tests for categorical data. Comparisons involve the 2 total age groups; \( ^{x 3} \)including phototherapy. When presenting percentages, numerator and denominator are reported. BMI: body mass index; BSA: body surface area; DLQI: Dermatology Life Quality Index; NA: not applicable; PASI: Psoriasis Area and Severity Index; PGA: Physician Global Assessment; PsA: psoriatic arthritis; SD: standard deviation; \( w \): week.
The proportions of patients with PASI < 5 coincided with those found for PASI < 3, with no differences between age groups at week 244. The proportions of patients with PASI < 1 was lower compared with PASI < 3 and < 5, with a slight tendency to show a benefit in patients aged ≥ 65 vs < 65 years at week 244 (see Appendix S1).

Absolute mean PASI and absolute PASI < 3, < 5 and < 1 results evaluated by the sensitivity analysis (multiple imputation) is shown in Fig. S2 and Appendix S2, respectively.

There was no effect of age group on absolute PASI at weeks 28 and 244. There was a significant effect of baseline PASI, treatment, week and smoking status \((p < 0.001)\) on absolute PASI at week 28. At week 244, there were significant effects on absolute PASI for baseline PASI \((p < 0.001)\), treatment \((p = 0.02)\), BMI \((p = 0.001)\), prior biological therapy for psoriasis \((p = 0.01)\), smoking status \((p = 0.007)\), and the interaction term age group \(\times\) week \((p = 0.003)\) (Table SIII). The mixed model on PASI course for sensitivity analysis with multiple imputation is shown in Table SIV.

**DLQI and DLQI-R.** The proportion (95% CI) of TIL-treated patients aged < 65 vs ≥ 65 years achieving a DLQI 0/1 for TIL 100 mg at week 28 was 53.8% (49.5–58.2%) vs 53.9% (49.5–58.2%), and at week 52, it was 66.8% (61.1–72.1%) vs 68.2% (56.8–76.5%). The proportion (95% CI) of TIL-treated patients aged < 65 vs ≥ 65 years achieving a DLQI 0/1 for TIL 200 mg at week 28 was 61.1% (56.8–65.3%) vs 60.4% (54.3–74.2%), and at week 52 it was 72.0% (65.2–78.1%) vs 81.3% (54.4–96.0%) (see Fig. 3) (comparison by age groups (combining TIL doses) at week 52: \(p = 0.54\)).

The absolute mean DLQI scores and DLQI-R results are shown in Appendix S3. The mean absolute DLQI
and DLQI-R scores were similar, except for patients aged ≥65 years for TIL 100 mg at week 28, where the mean DLQI-R was higher (4.5 (5.2) vs 3.4 (3.9)). The proportions of patients for each age group maintained the same trend for DLQI-R as those observed for DLQI, although the proportions are lower for the former.

**Safety outcomes**

Summary of exposure adjusted rates of AEs attending TIL dose and age are shown in **Table II**. The cumulative incidence of TEAEs for TIL 100 mg/TIL 200 mg in patients <65 vs ≥65 years was 4,717/5,032 vs 515/545 per 100 patient-years of exposure, within which the highest cumulative incidence of infections were 621/592 vs 23/51 per 100 patient-years of exposure for nasopharyngitis, followed by 168/203 vs 11/10 for other upper respiratory tract infection, and 76/94 vs 3/10 for influenza.

With regards to TEAEs of special interest, the cumulative incidence for TIL 100 mg/TIL 200 mg in patients <65 vs patients ≥65 years was 17/11 vs 4/6 per 100 patient-years of exposure for malignancy excluding NMSC, 6/10 vs 8/6 for NMSC, and 14/21 vs 1/3 for confirmed extended MACEs.

A total of 6 (0.2%) drug-related serious AEs (SAEs) per 100 patient-years of exposure (100 mg TIL)/4 (0.2%) (200 mg TIL) led to discontinuation in the younger group, and 3 (12.5%)/1 (0.5%) in the older group. The specific drug-related SAEs are shown in **Table III**.

**DISCUSSION**

The increasing number of elderly patients (≥65 years) with moderate to severe psoriasis in daily practice represents a challenge for dermatologists. However, evidence in this patient population is limited to a few biological agents and small-molecule inhibitors (22). This is one of the first studies to depict the efficacy and safety data of TIL for older vs younger patients from randomized clinical trials. This comparison is important because of possible differences in patient profile and the increasing number of older patients with psoriasis needing a safe and effective treatment. TIL in patients ≥65 years appears to be effective and safe in long-term psoriasis management, which was comparable to younger patients.

Differences in comorbidities were evident between both age groups. The current study found a higher proportion of musculoskeletal, metabolic, and vascular disorders, proportionally, in older patients. These disorders are more common in old age (23–25), and have been (partially) related to the existence of psoriasis (26–28). Biologics appear to have a good safety profile and are usually well tolerated. In addition, in terms of comedication, older patients had a higher intake of drugs related to cardiac or gastric problems. Since TIL is cleared from the body by general protein catabolism processes, and is not eliminated by renal or hepatic pathways, no interaction between TIL and the medications taken in this population has been described (29).

Although the current study found some differences among the presence of comorbidities and comedication, comparable long-term PASI and DLQI responses were found in younger and older patients, independently of the administration dose, without safety concerns. In the long-term (week 244), the current study found that more than 80% and 90% of younger and older patients, respectively, showed a PASI<3. These results are consistent with other studies on the long-term effects of different biologics on PASI responses (30).

**Table II. Exposure-adjusted rates of adverse events (AEs) and treatment-emergent AEs (TEAEs) by age group and tildrakizumab dose**

<table>
<thead>
<tr>
<th>Condition</th>
<th>&lt;65 years</th>
<th>≥65 years</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total follow-up, patient-year</td>
<td>2,487.7</td>
<td>2,531.6</td>
</tr>
<tr>
<td>Any SAE</td>
<td>205 (8.2) [7.1–9.4]</td>
<td>206 (8.1) [7.0–9.3]</td>
</tr>
<tr>
<td>Drug-related SAEs</td>
<td>18 (0.7) [0.4–1.1]</td>
<td>11 (0.4) [0.2–0.7]</td>
</tr>
<tr>
<td>SAEs leading to discontinuation</td>
<td>22 (0.9) [0.5–1.3]</td>
<td>16 (0.6) [0.3–1.0]</td>
</tr>
<tr>
<td>Drug-related SAEs leading to discontinuation</td>
<td>6 (0.2) [0.0–0.4]</td>
<td>4 (0.2) [0.0–0.3]</td>
</tr>
<tr>
<td>Any TEAE</td>
<td>4,717 (189.6) [184.1–195.1]</td>
<td>5,032 (198.8) [193.2–204.4]</td>
</tr>
<tr>
<td>Drug-related TEAEs</td>
<td>752 (30.2) [28.1–32.4]</td>
<td>989 (39.1) [36.6–41.6]</td>
</tr>
<tr>
<td>TEAEs leading to discontinuation</td>
<td>42 (1.7) [1.2–2.2]</td>
<td>30 (1.2) [0.8–1.6]</td>
</tr>
<tr>
<td>Drug-related AEs leading to discontinuation</td>
<td>16 (0.6) [0.3–1.0]</td>
<td>8 (0.3) [0.1–0.5]</td>
</tr>
<tr>
<td>Deaths</td>
<td>9 (0.446) [0.1–0.6]</td>
<td>4 (0.2) [0.0–0.3]</td>
</tr>
<tr>
<td>TEAEs of special interest:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Severe infection*</td>
<td>31 (1.3) [0.8–1.7]</td>
<td>41 (1.6) [1.1–2.1]</td>
</tr>
<tr>
<td>Malignancy excluding NMSC</td>
<td>17 (0.7) [0.4–1.0]</td>
<td>13 (0.4) [0.2–0.7]</td>
</tr>
<tr>
<td>NMSC</td>
<td>6 (0.2) [0.0–0.4]</td>
<td>10 (0.4) [0.2–0.6]</td>
</tr>
<tr>
<td>Confirmed extended MACE</td>
<td>14 (0.6) [0.3–0.9]</td>
<td>21 (0.8) [0.5–1.2]</td>
</tr>
<tr>
<td>Injection-site reaction</td>
<td>66 (2.7) [2.0–3.3]</td>
<td>81 (3.2) [2.5–3.9]</td>
</tr>
<tr>
<td>Drug-related hypersensitivity reaction</td>
<td>14 (0.6) [0.3–0.9]</td>
<td>5 (0.2) [0.0–0.4]</td>
</tr>
</tbody>
</table>

Data shown as n (number of events per 100 patient-years of exposure) [95% CI]. *Severe infection was defined as any infection meeting the regulatory definition of a serious AE (i.e. resulted in death, was life-threatening, required inpatient hospitalization or prolongation of existing hospitalization, resulted in persistent or significant disability or incapacity, or required intervention to prevent 1 of the other outcomes listed), or any infection requiring intravenous antibiotic.

95% CI: 95% confidence interval; MACE: major adverse cardiovascular event; NMSC: non-melanoma skin cancer; SAEs: serious AEs.
Safety analysis showed a favourable tolerability profile in both age groups. Adverse events were consistent with the rates observed in other clinical trials with biologics (37). In terms of infection, both age groups showed a similar profile, sharing the highest incidence of nasopharyngitis and other respiratory tract infection, in line with those of phase II–III trials with biologics (38, 39) and available real-world evidence (RWE) registries (40), with no new safety evidence. However, in terms of TEAEs of special interest, older patients showed proportionally more cases of cardiovascular events, NMSC and other malignancies compared with younger patients, as previously described in the literature (13), most likely related to the ageing process and the longer psoriatic disease duration, and not due to the psoriatic treatment administered. In general, these results demonstrate the potential benefit of TIL in older patients without affecting their safety profile.

**Limitations**

This study has some limitations. The main limitation is the relatively small number of older patients. In addition, older patients with extensive multimorbidity and/or polypharmacy are less likely to be included in clinical trials. The investigated older patients may still be a relatively healthy older group, as they passed the clinical trial inclusion and exclusion criteria. In this vein, exclusion criteria for reSURFACE trials are not based on (or represent) RWE. For example, some of the common comorbidities associated with psoriasis or its variants (psoriatic arthritis, erythrodermic psoriasis) were exclusion criteria in the reSURFACE trials, as well as recurrent infections (41–43). In addition, patients with extensive pre-treatment were excluded, as they had to wait until their psoriasis showed a PASI ≥ 12. In clinical practice, this is not feasible, under-representing to the trial populations what clinicians see in daily clinical practice. Despite the fact that some data indicate that patients treated in routine practice with TIL differed substantially from those included in phase III studies (44), a RWE study has recently confirmed that there is no efficacy-effectiveness gap for TIL (45). Further research with a larger number of older patients in a real-world setting is needed to confirm these preliminary results. Another limitation is the lack of age-randomized groups and control settings.

**Conclusion**

In these current post hoc analyses, TIL demonstrated long-term control with a favourable safety in patients below and above 65 years of age. This confirms the limited RWE on the clinical effectiveness of TIL in older patients with moderate to severe psoriasis. Despite the differences between the age groups in terms of comorbidities and comedations, these results indicate a similar
percentage improvement in disease severity in the 2 age groups, with comparable improvements in quality of life and without major safety issues. Further confirmatory studies are desirable, with dedicated and real-world trials to better understand the profile of biological management of this group of patients.

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REFERENCES