Long-term, durable, absolute Psoriasis Area and Severity Index and health-related quality of life improvements with risankizumab treatment: a post hoc integrated analysis of patients with moderate-to-severe plaque psoriasis

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Abstract

Background Risankizumab has demonstrated durable, high rates of efficacy in patients with moderate-to-severe plaque psoriasis as assessed by the achievement of relative Psoriasis Area and Severity Index (PASI) improvement and Dermatology Life Quality Index (DLQI) 0/1.

Objectives The aim of this post hoc analysis is to assess the achievement of absolute PASI thresholds and related improvements in health-related quality of life (HRQoL) in patients with moderate-to-severe plaque psoriasis treated with (i) risankizumab compared with ustekinumab, and (ii) long-term (>52 weeks to 172 weeks) risankizumab.

Methods Data from patients randomised to 150 mg risankizumab or 45 or 90 mg ustekinumab in replicate randomised controlled trials UltiMMa-1 and UltiMMa-2 were analysed for the achievement of absolute PASI thresholds PASI ≤ 3, PASI ≤ 1, and PASI = 0, time to achieve these thresholds, and combined PASI and DLQI endpoints. Data from patients initially randomised to risankizumab who continued on risankizumab in the open-label extension study LIMMitless were analysed for the achievement of absolute PASI levels, mean DLQI scores, and DLQI 0/1.

Results Significantly greater proportions of patients treated with risankizumab compared with ustekinumab achieved PASI ≤ 3, PASI ≤ 1, and PASI = 0, as well as combined endpoints for absolute PASI and DLQI [PASI ≤ 3 and DLQI ≤ 5] or (PASI ≤ 1 and DLQI 0/1)]. The median time to first achieve PASI ≤ 3, PASI ≤ 1, and PASI = 0 was significantly lower for risankizumab-treated patients compared with ustekinumab-treated patients. Among patients treated with long-term risankizumab, more than 90% achieved PASI ≤ 3 though week 172 and more than 80% achieved DLQI 0/1. Low absolute PASI scores corresponded with low mean absolute DLQI scores through week 172 of continuous risankizumab treatment.

Conclusions Risankizumab treatment demonstrated high rates of rapid and durable efficacy as measured by absolute PASI thresholds and improvements in patient HRQoL.

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Conflicts of interest

M Gooderham has been an investigator, speaker, adviser, or consultant for AbbVie, Amgen, Akros, Arcutis, Boehringer-Ingelheim, BMS, Celgene, Coherus, Dermira, Dermavant, Eli Lilly, Galderma, GSK, Incyte, Janssen, Kyowa Kirin Pharma, Leo Pharma, MedImmune, Merck, Novartis, Pfizer, Roche, Regeneron, Sanofi-Genzyme, Sun Pharma, UCB, and Valeant/Bausch. A Pinter has been an investigator, speaker, adviser, or consultant for AbbVie, Allmirall-Hermal, Amgen, Biogen Idec, Boehringer-Ingelheim, Celgene, GSK, Eli Lilly, Galderma, Hexal, Janssen, LEO Pharma, MC2, Medac, Merck Serono, MSD, Novartis, Pfizer, Tigercat Pharma, Regeneron, Roche, Sandoz Biopharmaceuticals, Schering-Plough, and UCB Pharma. LK Ferris has been an investigator for Amgen, AbbVie, Eli Lilly, Janssen, BMS, Arcutis, Dermavant, Regeneron, InflaRx, Novartis, and UCB; and a consultant for AbbVie, Eli Lilly, Janssen, BMS, Arcutis, Dermavant, and Pfizer. RB Warren received grants from AbbVie, Almirall, Amgen, Celgene, Janssen, Lilly, LEO Pharma, Medac, Novartis, Pfizer, and UCB and received personal fees from AbbVie, Almirall, Amgen, Arena Pharmaceuticals,
Introduction

Psoriasis is a chronic, immune-mediated, systemic inflammatory disease with prominent skin manifestations estimated to affect between 1% and 4% of the population in developed countries, approximately 100 million people worldwide.1–3 Approximately 20% of patients are estimated to have a more severe form of psoriasis, affecting ≥10% of body surface area.4 Psoriasis has a profound negative impact on patients’ quality of life and increases the risk for early mortality and prevalence of comorbidities, including cardiovascular disease, hypertension, hyperlipidemia, diabetes, and depression.5–7 Complete or nearly complete clearance is now achievable with available biologic treatments targeting various cytokines involved in disease pathogenesis, namely interleukin (IL)-17, IL-23, and tumour necrosis factor (TNF)-α.8–11; however, the durability of response with many biologics is limited in clinical practice after 1–2 years due to loss of treatment effect over time.12,13

IL-23, a key regulatory cytokine, is essential for pathogenic T helper 17 (Th17) cell differentiation, activation, and survival.14 In psoriasis, the IL-23/Th17 pathway is activated, driving cutaneous plaque formation and chronic inflammation.14,15 Recent clinical trials demonstrated that selective inhibition of IL-23 through antibodies targeting the p19 subunit produced high and durable efficacy associated with reductions in inflammatory cytokine expression in skin.8,16–19 Specific inhibition of IL-23 may offer additional safety benefits over biologics that target IL-17 by preserving the function of IL-23-independent, IL-17–producing cells that are involved in mucocutaneous defence and barrier tissue integrity.20

Risankizumab is a humanised immunoglobulin G1 monoclonal antibody that binds with high affinity to the p19 subunit and specifically inhibits IL-23.21 Phase 3 clinical trials in patients with moderate-to-severe plaque psoriasis have demonstrated superior efficacy for risankizumab compared with placebo, adalimumab, ustekinumab, and secukinumab at week 16, which was sustained through weeks 44 (compared with adalimumab) and 52 (compared with ustekinumab and secukinumab).18,19,22

Relative improvements in the Psoriasis Area and Severity Index (PASI; eg, PASI 75 or PASI 90), often used for assessing the benefits of psoriasis therapies in clinical trials, have some limitations. Notably, calculation of relative PASI requires assessment of baseline disease status, which may not be available in routine clinical practice or other clinical settings in which patients may switch from one treatment to another, may have difficulty achieving PASI 90, given the known lack of precision in the PASI at lower levels of disease activity. Recent treat-to-target guidelines from the National Psoriasis Foundation recommend basing decisions to change therapy primarily on level of skin involvement and not on percent PASI reduction.23 Absolute PASI provides an additional tool to standardise measures of efficacy and may provide more clinically relevant information when evaluating the efficacy of biologics during the treatment maintenance phase, when changing therapy, and when baseline PASI information is lacking. Low absolute PASI values have been correlated with improvements in patient health-related quality of life (HRQoL),24–26 and treatment goals defined by absolute PASI targets have enabled a more standardised quality of care. Additionally, patients who achieve or almost-clear skin are more likely to report that psoriasis has no impact on their HRQoL and other measures of symptoms.27,28 In this report, we assess achievement of absolute PASI thresholds and improvements in HRQoL in patients with moderate-to-severe plaque psoriasis.
treated with risankizumab compared to ustekinumab using integrated data from 2 pivotal phase 3 randomised controlled trials (RCTs), UltIMMa-1 and UltIMMa-2, and assess the long-term (>52 to 172 weeks) efficacy of risankizumab using data from the ongoing open-label extension (OLE) study LIMMitless.

Methods

Study design and patients
This is a post hoc integrated analysis of the UltIMMa-1 and UltIMMa-2 replicate phase 3, randomised, double-blinded, placebo- and active comparator–controlled studies, comparing risankizumab and ustekinumab efficacy and safety, with long-term risankizumab data analysis from the LIMMitless ongoing, phase 3, single-arm, international, multicentre, OLE study. The eligibility criteria for UltIMMa-1 and UltIMMa-2 have been previously published.19 Briefly, the studies included adult patients (≥18 years) with stable (≥6 months) moderate-to-severe chronic plaque psoriasis (with or without psoriatic arthritis) with body surface area (BSA) involvement ≥10%, PASI ≥ 12, and static Physician’s Global Assessment (sPGA) ≥ 3. PASI was collected with 1 digit after the decimal, while BSA was up to integer in percentage scale.

The comparator analysis is focused on patients initially randomised to risankizumab or ustekinumab, receiving either 150 mg risankizumab or 45/90 mg ustekinumab based on weight per label (45 mg for patients with body weight ≤100 kg or 90 mg for patients with body weight >100 kg) subcutaneously at weeks 0 and 4 and then every 12 weeks (Q12W) through week 40 (Fig. 1a). All patients who completed UltIMMa-1 or UltIMMa-2 and were candidates for long-term risankizumab treatment were eligible to enrol in the ongoing, phase 3, single-arm, international, multicentre, OLE study LIMMitless (NCT03047395). There was no requirement for response thresholds to enrol in LIMMitless, and additional inclusion and exclusion criteria are listed in Table S1 (Supporting Information). Upon enrollment in LIMMitless, patients receive 150 mg open-label risankizumab subcutaneously Q12W with no washout period (Fig. 1b). The long-term analysis, herein, is focused on patients initially randomised to risankizumab in UltIMMa-1 or UltIMMa-2 who elected to enrol in LIMMitless and continued to receive risankizumab as 150 mg open-label Q12W.

The studies were conducted in accordance with the protocol, International Council for Harmonisation of Technical Requirements for Pharmaceuticals for Human Use guidelines, applicable local regulations and Good Clinical Practice guidelines governing clinical study conduct, and ethical principles outlined in the Declaration of Helsinki. All study-related documents (including study protocols) were approved by an institutional review board or independent ethics committee at each study site; all patients provided written informed consent prior to study participation.

Outcomes
In this analysis, efficacy was assessed using measures of disease activity, including the proportion of patients achieving absolute PASI ≤ 3, PASI ≤ 1, and PASI 0 through 52 weeks in UltIMMa-1 or UltIMMa-2 and through 172 weeks in LIMMitless (interim snapshot on March 26, 2020). To gain further insight into the distribution of responses at each time point, absolute PASI response data were regrouped into 6 non-overlapping categories representing the proportion of patients reaching absolute PASI > 10, >5–10, >3–5, >1–3, >0–1, and 0–0, assessed through week 52 in UltIMMa-1 and UltIMMa-2 and through week 172 in LIMMitless. Time to achievement of absolute PASI thresholds (PASI ≤ 3, PASI ≤ 1, and PASI = 0) was assessed for patients treated with either risankizumab or ustekinumab in UltIMMa-1 and UltIMMa-2. To understand the relationship between absolute PASI and HRQoL, 2 distinct combined endpoints were assessed as (i) achievement of (PASI ≤ 3 and Dermatology Life Quality Index [DLQI] ≤ 5) or (ii) achievement of (PASI ≤ 1 and DLQI 0/1) at weeks 16 and 52 in UltIMMa-1 and UltIMMa-2. For the long-term risankizumab population, mean absolute DLQI scores and proportion of patients achieving DLQI 0/1 were assessed every 24 weeks starting after week 52 through week 172 in LIMMitless. Safety was assessed as treatment-emergent adverse events (TEAEs) for all patients receiving at least 1 dose of study medication.

Statistical analysis
For the UltIMMa-1 and UltIMMa-2 integrated post hoc analysis, patients with missing efficacy data for categorical variables were handled by non-responder imputation (NRI) and for continuous variables with last observation carried forward (LOCF). Categorical variables were tested using the Cochran-Mantel-Haenszel risk difference estimate. For the LIMMitless post hoc analysis, missing efficacy data were imputed using modified NRI (mNRI; non-response is imputed only for treatment failures, defined as patients who have worsening of psoriasis, then mixed-effect model was used on the imputed dataset), LOCF, and observed cases (OC). Nominal P values are reported. All statistical analyses were conducted using SAS® version 9.4 (SAS Institute, Inc., Cary, NC, USA) or higher.

Results
In UltIMMa-1 and UltIMMa-2, 797 patients were randomised to risankizumab (150 mg, n = 598) or ustekinumab (45 or 90 mg, n = 199); efficacy and safety for these pivotal trials have been previously published.19 Among patients initially randomised to risankizumab, 525 enrolled in LIMMitless and continued to receive open-label 150 mg risankizumab (Fig. S1, Supporting Information). At the time of this interim analysis, 465/525 patients receiving long-term, continuous risankizumab had completed 172 weeks of treatment in LIMMitless. Baseline patient demographics and disease characteristics upon entry to
UltIMMa-1 and UltIMMa-2 were generally balanced between treatment groups and were mostly consistent with those of recent trials in patients with psoriasis (Table 1). Reasons for discontinuation in UltIMMa-1, UltIMMa-2, and LIMMitless are reported in Fig. S1 (Supporting Information).

Efficacy of risankizumab versus ustekinumab was assessed as the achievement of absolute PASI thresholds for patients who participated in UltIMMa-1 and UltIMMa-2. Significantly greater proportions of patients receiving risankizumab compared with those receiving ustekinumab achieved PASI $\leq 3$ as early as week 4 ($P < 0.01$), and this difference was increased and maintained through week 52 ($P < 0.001$; Fig. 2a). Similarly, greater proportions of risankizumab-treated patients achieved PASI $\leq 1$ (Fig. 2b) and PASI $= 0$ (Fig. 2c) compared with ustekinumab-treated patients as early as week 8 and through week 52 of treatment. Among risankizumab-treated patients, 87.1%, 69.9%, and 57.9% achieved PASI $\leq 3$, $\leq 1$, and $= 0$ at week 52, respectively (Fig. 2). Regrouping absolute PASI response data into the achievement of absolute PASI tiers allows for visualisation of individual improvement. Typical to the enrollment criteria of clinical trials, all patients entered UltIMMa-1 and UltIMMa-2 with absolute PASI $\geq 12$. Greater proportions of risankizumab-treated patients achieved lower absolute PASI thresholds (PASI $\leq 5$, PASI $\leq 3$, PASI $\leq 1$, and PASI $= 0$) than ustekinumab-treated patients as early as week 4 and through week 52 (Fig. 3 and Video S1, Supporting Information). The rapidity of absolute PASI response was also assessed. The median time to achieve PASI $\leq 3$ (58 vs 84 days), PASI $\leq 1$ (94 vs 199 days), and PASI $= 0$ (148 vs 370 days) was lower for risankizumab-treated patients than for ustekinumab-treated patients, respectively, $P < 0.001$ for all (Table 2).

To better understand the level of HRQoL improvement that patients experience with skin clearance, we assessed the achievement of combined PASI and DLQI targets for patients treated with risankizumab compared to those treated with ustekinumab. A significantly greater proportion of risankizumab-treated patients achieved combined absolute PASI $\leq 3$ and DLQI $\leq 5$ at week 16 and 52 compared with ustekinumab patients (Fig. 4a); the adjusted difference between the 2 treatments was 26.2% [18.6, 33.8 (95% CI)] and 27.2% [19.8, 34.6 (95% CI)] for week 16 and week 52, respectively. Similarly, combined absolute PASI $\leq 1$ and DLQI 0/1 was achieved by a greater proportion of risankizumab-treated patients than ustekinumab-treated patients (Fig. 4b), with adjusted differences between the 2

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Figure 1  Study design. Study design for (a) UltIMMa-1 and UltIMMa-2 post hoc analysis, full study design included a placebo arm$^{19}$ and (b) LIMMitless; OLE, open-label extension; RCT, randomised controlled trial; RZB, risankizumab; UST, ustekinumab. $^a$Patients weighing $\leq 100$ kg received 45 mg UST, while patients weighing $>100$ kg received 90 mg UST, per label instructions.
treatments of 22.1% [15.2, 29.1 (95% CI)] and 30.3% [23.1, 37.6 (95% CI)] for week 16 and week 52, respectively.

To assess the long-term efficacy of risankizumab treatment, we evaluated the achievement of absolute PASI thresholds for patients who were initially randomised to risankizumab in UltIMMa-1 or UltIMMa-2, completed the base study, and enrolled in the ongoing LIMMitless OLE study (n = 525). At week 172, with more than 3 years of continuous risankizumab treatment, the proportion of patients achieving an absolute PASI ≤ 3 remained above 90% [92.8% (LOCF) and 93.0% (mNRI), Fig. 5a, 94.2% (OC) (N = 465), Table S2, Supporting Information]. Similar durability is demonstrated by the proportion of patients achieving absolute PASI ≤ 1 [75.8% (LOCF) and 74.9% (mNRI), Fig. 5b, 77.2% (OC) (N = 465), Table S2, Supporting Information] and PASI = 0 [64.8% (LOCF) and 58.3% (mNRI), Fig. 5c, 66.0% (OC) (N = 465), Table S2, Supporting Information]. Additionally, these patients achieved high rates of traditional measures PASI 90 [88.6% (LOCF), 88.4% (mNRI), and 90.5% (OC)] and PASI 100 [64.6% (LOCF), 62.9% (mNRI), and 66.0% (OC)], as well as 95.9% (LOCF) and 96.6% (OC) mean percent PASI improvement from baseline after 3 years of continuous risankizumab treatment (Fig. S2a,b and Table S2, Supporting Information). Long-term risankizumab treatment also resulted in achievement of DLQI 0/1 in a high proportion of patients through week 172 [83.0% (LOCF) and 82.5% (mNRI), Fig. 6a, 84.3% (OC) (N = 471), Table S3, Supporting Information]. Grouping of absolute PASI response levels demonstrates high rates of skin clearance for risankizumab-treated patients through week 172 (Fig. 6b). Long-term risankizumab treatment also resulted in achievement of mean absolute DLQI < 2 by week 16 and mean DLQI = 1.0 at week 124 through week 172 (Fig. 6b).

Rates of adverse events (AEs) ranged from 157.0–228.0 events/100 patient-years (E/100 PY), serious AEs from 6.3–9.4 E/100 PY, and serious infections from 1.1–1.8 E/100 PY in the UltIMMa-1/UltIMMa-2 and LIMMitless populations (Table 3).
In this post hoc analysis of the randomised, double-blinded, controlled clinical trials UltIMMa-1 and UltIMMa-2 and the long-term data from the OLE study LIMMitless, patients treated with risankizumab achieved clinically relevant absolute PASI thresholds and improved patient HRQoL. By these measures, risankizumab demonstrated superior efficacy to ustekinumab over 52 weeks and durable rates of efficacy over 172 weeks of continuous treatment.

Risankizumab treatment resulted in statistically superior (nominal P value) rates of patients achieving absolute PASI ≤0, ≤1, and ≤3 compared to ustekinumab by week 8. In

**Table 2** Time to achievement of absolute PASI thresholds (days)

| Absolute PASI | RZB (n = 598) | UST (n = 199) | RZB vs UST |
|---------------|---------------|---------------|------------|
|               | Median Days   | 25th Percentile | Median Days | 25th Percentile | P-value of log-rank Test, hazard ratio (95% CI) |
| PASI ≤ 3      | 58            | 56            | 85         | 84          | 57          | 153        | <0.001     |
|               |               |               |            |             |             | 1.664 (1.397, 1.982) |
| PASI ≤ 1      | 94            | 60            | 159        | 199         | 86          | N/A        | <0.001     |
|               |               |               |            |             |             | 2.273 (1.864, 2.773) |
| PASI = 0      | 148           | 85            | 280        | 370         | 145         | N/A        | <0.001     |
|               |               |               |            |             |             | 2.400 (1.926, 2.991) |

PASI, Psoriasis Area and Severity Index; RZB, risankizumab; UST, ustekinumab.

**Discussion**

In this post hoc analysis of the randomised, double-blinded, controlled clinical trials UltIMMa-1 and UltIMMa-2 and the long-term data from the OLE study LIMMitless, patients treated with risankizumab achieved clinically relevant absolute PASI thresholds and improved patient HRQoL. By these measures, risankizumab demonstrated superior efficacy to ustekinumab over 52 weeks and durable rates of efficacy over 172 weeks of continuous treatment.

Risankizumab treatment resulted in statistically superior (nominal P value) rates of patients achieving absolute PASI = 0, ≤1, and ≤3 compared to ustekinumab by week 8. In
risankizumab-treated patients, the proportion of patients treated with risankizumab who achieved an absolute PASI score of 0 continued to increase from week 16 to week 52. Risankizumab demonstrated numerically higher rates of achieving an absolute PASI = 0 and numerically lower rates of achieving absolute PASI scores >5 compared with ustekinumab at weeks 16 and 52. When assessed as rapidity of response, the median time to achieve PASI = 0, ≤1, and ≤3 was shorter for patients treated with risankizumab compared with ustekinumab. Long-term, continuous risankizumab dosed Q12W provided long-term maintenance of efficacy in terms of absolute PASI improvement through week 172, with most patients achieving clear skin (PASI = 0) for more than 3 years. Traditional measures of PASI 90, PASI 100, and mean percent PASI improvement also demonstrate consistently durable, high rates of skin clearance in this population. Safety outcomes for UltIMMa-1 and UltIMMa-2 have been previously reported,19 and safety for long-term risankizumab treatment was consistent with the known safety profile for risankizumab with no new safety signals.29

With the emergence of new, improved therapies for psoriasis, high levels of skin clearance are increasingly considered as a treatment target because of the impact of residual disease on patients’ quality of life.24-26,30,31 Patients who obtain clear or almost-clear skin are more likely to report that psoriasis has no impact on their HRQoL and other measures of disease signs and symptoms.27,28,32 In this analysis, achievement of low absolute PASI scores corresponded with an improvement in HRQoL as assessed by DLQI.

Figure 5 Long-term achievement of absolute PASI thresholds (mNRI and LOCF). Percentage of patients achieving absolute PASI scores of (a) ≤3, (b) ≤1, (c) or = 0 over long-term, continuous risankizumab treatment (OC, LOCF, and mNRI). *The drop-off in response rates (mNRI) is attributed to the lower number of observations at this timepoint, as many patients had not yet completed 172 weeks of treatment by the time of this analysis. Due to more missing data at this time point, the mixed-effect analysis results in lower response rates. Imputed patients who discontinued due to worsening of symptoms as non-responders; all other missing data handled using mixed-effects models. LOCF, last observation carried forward; mNRI, modified non-responder imputation; OC, observed cases; PASI, Psoriasis Area and Severity Index.
Figure 6  Long-term achievement of (a) DLQI 0/1 (mNRI and LOCF), (b) absolute PASI categories, and mean absolute DLQI (LOCF). (a) Percentage of patients achieving DLQI 0/1 over long-term, continuous risankizumab treatment (mNRI and LOCF). Error bars represent 95% confidence interval based on the normal approximation. *The drop-off in response rates (mNRI) is attributed to the lower number of observations at this timepoint, as many patients had not yet completed 172 weeks of treatment by the time of this analysis. Due to more missing data at this timepoint, the mixed-effect analysis results in lower response rates. (b) Percentage of patients within absolute PASI categories and mean DLQI achieved by patients with long-term, continuous risankizumab treatment (LOCF). Each circle represents the proportion of patients falling into each absolute PASI category; the whole numbers indicate that proportion rounded to the nearest percentage. All patients started the trials with a PASI > 10 since 1 of the inclusion criteria required patients to have a baseline PASI ≥ 12. DLQI, Dermatology Life Quality Index; LOCF, last observation carried forward; mNRI, modified non-responder Imputation (imputed patients who discontinued due to worsening of symptoms as non-responders; all other missing data handled using mixed-effects models). PASI, Psoriasis Area and Severity Index; RZB, risankizumab.
Table 3  Treatment-emergent adverse events

| Treatment-emergent adverse events, n (%) | Events (E/100 PY) |
|----------------------------------------|------------------|
|                                         | UltIMMa-1/2      | LIMMItless      |
|                                         | n = 598 (PY = 618) | n = 525 (PY = 1909.5) |
| Any AE                                  | 1409 (228.0)     | 2998 (157.0)    |
| Serious AE                              | 58 (9.4)         | 121 (6.3)       |
| Severe AE                               | 47 (7.6)         | 109 (5.7)       |
| AE leading to drug discontinuation      | 5 (0.8)          | 30 (1.6)        |
| Serious infection                       | 11 (1.8)         | 21 (1.1)        |
| Active tuberculosis                     | 0                | 0               |
| Adjudicated MACE                        | 2 (0.3)          | 1 (-0.1)        |
| Malignancies                            | 3 (0.5)          | 19 (1.0)        |
| Malignancies excluding NMSC             | 0                | 8 (0.4)         |
| Serious hypersensitivity                | 0                | 2 (0.1)         |
| Deaths (including non-TEAEs)            | 2 (0.3)          | 3 (0.2)         |

AE, adverse event; E/100 PY, events per 100 patient-years; MACE, major adverse cardiac event; NMSC, non-melanoma skin cancer; RZB, risankizumab; TEAEs, treatment-emergent adverse events.

1 Events with onset after the first dose of RZB until 20 weeks after last dose of RZB.
2 Cardiopulmonary arrest, unrelated to study drug.
3 Automobile accident, unrelated to study drug.
4 Sudden cardiac death, unrelated to study drug.

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Data availability statement

AbbVie is committed to responsible data sharing regarding the clinical trials we sponsor. This includes access to anonymised, individual, and trial-level data (analysis data sets), as well as other information (e.g., protocols and clinical study reports), as long as the trials are not part of an ongoing or planned regulatory submission. This includes requests for clinical trial data for unlicensed products and indications.

This clinical trial data can be requested by any qualified researchers who engage in rigorous, independent scientific research, and will be provided following review and approval of a research proposal and statistical analysis plan and execution of a data sharing agreement. Data requests can be submitted at any time and the data will be accessible for 12 months, with possible extensions considered. For more information on the process, or to submit a request, visit the following link: https://www.abbvie.com/our-science/clinical-trials/clinical-trials-data-and-information-sharing/data-and-information-sharing-with-qualified-researchers.html.

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Absolute PASI and HRQoL with risankizumab in psoriasis

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Supporting information

Additional Supporting Information may be found in the online version of this article:

Figure S1. Patient disposition for UltIMMa-1/2 and LIMMitless.
Figure S2. Long-term (a) PASI 90 and (b) PASI 100 achievement and (c) mean percent improvement in PASI from baseline.
Table S1. LIMMitless trial eligibility criteria.
Table S2. Long-term achievement of absolute PASI thresholds (OC).
Table S3. Long-term achievement of DLQI 0/1 (OC).
Table S4. Long-term achievement of PASI 90 and PASI 100 (OC).
Video S1. Percentage of patients within absolute PASI score categories over time (LOCF).