

# Bimekizumab safety and tolerability in moderate to severe plaque psoriasis: Pooled analysis from up to 4 years of treatment in 5 phase 3/3b clinical trials

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## Synopsis

- Bimekizumab (BKZ) is a monoclonal immunoglobulin G1 antibody that selectively inhibits interleukin (IL)-17F in addition to IL-17A.<sup>1</sup>
- Psoriasis is a chronic condition requiring long-term management, thus evaluating long-term safety of treatments is essential to informing decision-making for clinicians, while managing risk for patients.<sup>2</sup>
- We report the first 4-year safety data for BKZ in patients with moderate to severe psoriasis.

## Objective

To evaluate BKZ safety data up to 4 years in patients with moderate to severe plaque psoriasis, using the largest pool of phase 3/3b safety data at the time of this study.

To assess whether rates of treatment-emergent adverse events (TEAEs) changed with each year of BKZ treatment.

## Methods

- Data were pooled from the BE SURE, BE VIVID, and BE READY phase 3 trials, their open-label extension (OLE) BE BRIGHT, the BE RADIANT phase 3b trial, and the BE RADIANT OLE.<sup>3-7</sup> The BE RADIANT trial ran for 3 years; therefore, the overall total pooled exposure only included BE RADIANT data to Year 3, in addition to BE BRIGHT data to Year 4. Data were pooled for all patients who received  $\geq 1$  BKZ dose in the included studies (Figure 1).
- Included patients received BKZ 320 mg every 4 weeks (Q4W) or every 8 weeks (Q8W); all received Q8W from Week 64 (BE RADIANT)/OLE Week 48 (BE BRIGHT) or the next scheduled clinic visit. Patients who switched from adalimumab, ustekinumab, or secukinumab to BKZ in BE SURE, BE VIVID, and BE RADIANT, respectively, were also included following the switch to BKZ.
- TEAEs were reported over 4 years using exposure-adjusted incidence rates (EAIRs) per 100 patient-years (PY).
- TEAEs were evaluated separately for Years 1, 2, 3, and 4 (Weeks 0–52, 52–104, 104–156, and 156–208) of BKZ treatment.

## Results

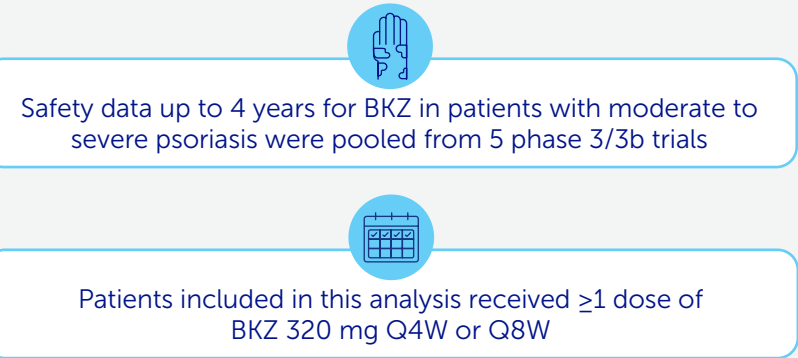
- Total BKZ exposure was 6,324.3 PY (N=2,186; Year 1, Year 2, Year 3, Year 4: 2,053.3 PY [n=2,186], 1,904.3 PY [n=2,013], 1,521.1 PY [n=1,803], 819.5 PY [n=1,309]; Table 1).
- TEAEs occurred at an EAIR of 170.5/100 PY (Year 1, Year 2, Year 3, Year 4: 230.9/100 PY, 137.7/100 PY, 107.1/100 PY, 99.9/100 PY), serious TEAEs at 5.5/100 PY (6.5/100 PY, 5.9/100 PY, 5.8/100 PY, 5.6/100 PY), and TEAEs leading to discontinuation at 2.9/100 PY (4.6/100 PY, 2.3/100 PY, 2.3/100 PY, 1.1/100 PY). Overall, the EAIR of TEAEs decreased with longer BKZ exposure over 4 years (Figure 2).
- The most common TEAEs were nasopharyngitis at 12.7/100 PY (Year 1, Year 2, Year 3, Year 4: 25.8/100 PY, 13.2/100 PY, 5.4/100 PY, 5.9/100 PY), oral candidiasis at 8.9/100 PY (18.9/100 PY, 10.7/100 PY, 6.8/100 PY, 5.4/100 PY), and upper respiratory tract infection at 5.7/100 PY (10.4/100 PY, 5.7/100 PY, 3.7/100 PY, 3.9/100 PY; Table 2).
- Fewer TEAEs over 4 years occurred with BKZ Q8W versus (vs.) Q4W (115.4/100 PY vs. 224.4/100 PY), including for oral candidiasis (6.5/100 PY vs. 16.7/100 PY).

## Conclusions

Bimekizumab demonstrated good tolerability and a comparable safety profile over 4 years in patients with moderate to severe plaque psoriasis.

EAIRs of TEAEs remained consistent or decreased with longer bimekizumab exposure over 4 years, with no new safety signals observed.

## Summary



EAIR of TEAEs remained consistent or decreased with longer BKZ exposure over 4 years

Figure 1 Included studies

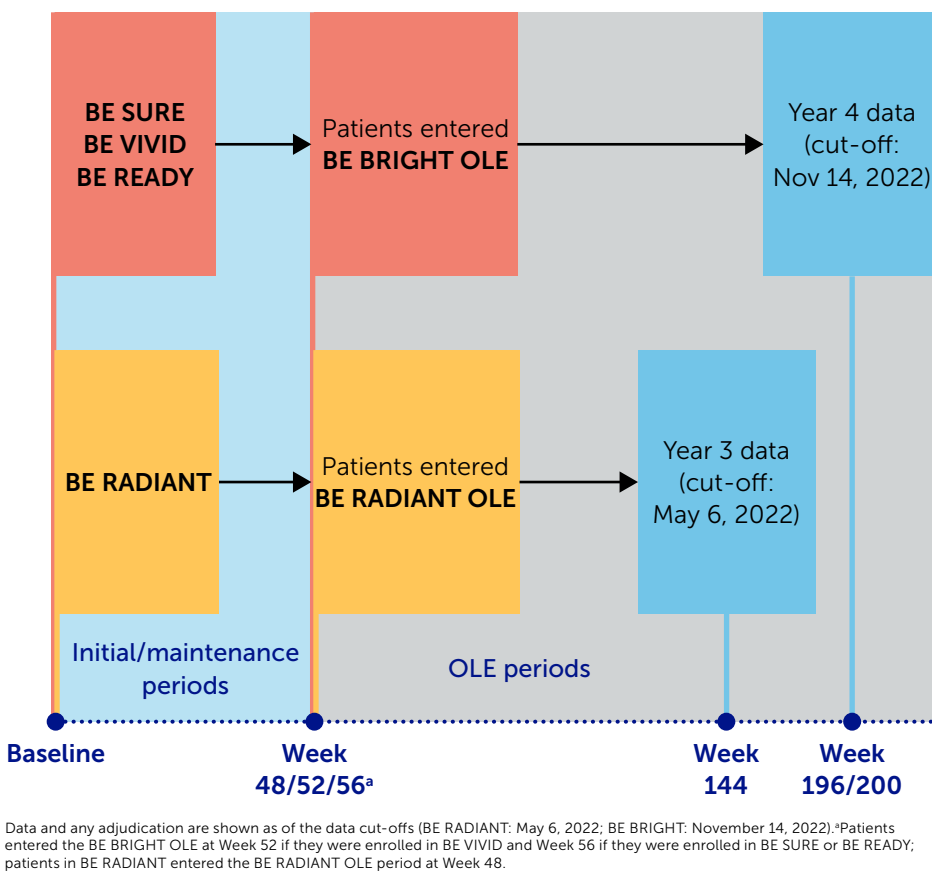
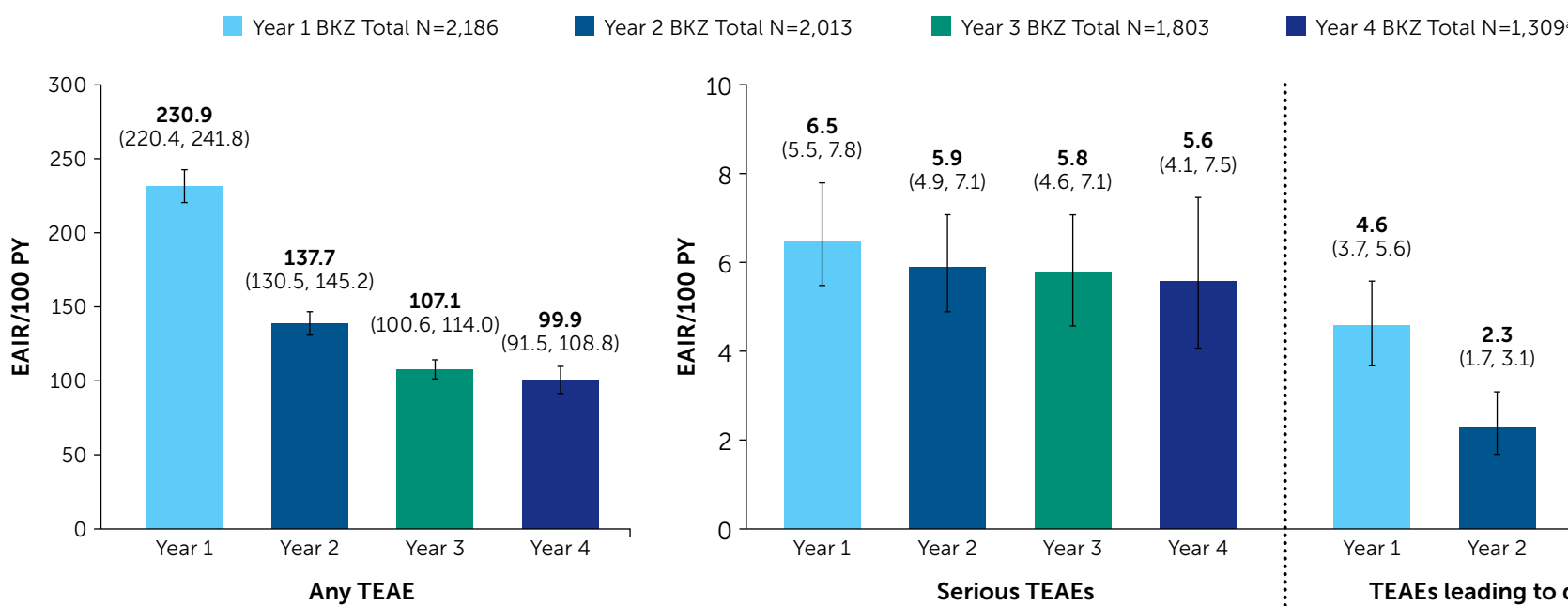


Figure 2 Incidence rates of TEAEs: Any, serious, and discontinuations over time (BKZ Total)



Data are reported as EAIRs; error bars represent 95% CI. Data are presented for the BKZ Total for the full pooled trial period, and separately for Years 1 (Weeks 0–52), 2 (Weeks 52–104), 3 (Weeks 104–156), and 4 (Weeks 156–208). Data were pooled for all patients who received  $\geq 1$  BKZ dose in each of the periods examined (BKZ Total). aBE RADIANT patients were not included after Year 3.

ALT: alanine aminotransferase; AST: aspartate aminotransferase; BKZ: bimekizumab; CI: confidence interval; COVID-19: Coronavirus disease 2019; EAIR: exposure-adjusted incidence rate; IL: interleukin; OLE: open-label extension; PY: patient-years; Q4W: every 4 weeks; Q8W: every 8 weeks; SD: standard deviation; TEAE: treatment-emergent adverse event; ULN: upper limit of normal; vs.: versus.

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**References:** Adams R et al. Front Immunol 2020;11:1894-1894. <sup>1</sup>Al-Jarrah A & Yu ZN. Psoriasis [Auckl] 2022;12:1-14; <sup>2</sup>Warren RB et al. N Engl J Med 2021;385(2):130-41. NCT03412747; <sup>3</sup>Reich K et al. Lancet 2021;397(10273):475-486. NCT03370133; <sup>4</sup>Gordon KB et al. JAMA Dermatol 2022;158(7):735-744. NCT03598790; <sup>5</sup>Reich K et al. N Engl J Med 2021;385(2):142-152. NCT03536884. **Author Contributions:** Substantial contributions to study conception/design, or acquisition/analysis/interpretation of data: KBG, DT, MG, YO, BS, LP, DD, JMLP, PG. Drafting of the publication, or reviewing it critically for important intellectual content: KBG, DT, MG, YO, BS, LP, DD, JMLP, PG. **Author Disclosures:** KBG: Received consulting fees from AbbVie, Amgen, Boehringer Ingelheim, Bristol Myers Squibb, Celgene, Dermira, Eli Lilly and Company, Janssen, Novartis, UCB Pharma; research support from AbbVie, Bristol Myers Squibb, Celgene, Eli Lilly and Company, Janssen, Novartis, and UCB Pharma. DT: Investigator and/or consultant/advisor for AbbVie, Amgen, Boehringer Ingelheim, Bristol Myers Squibb, Celgene, Dermira, Eli Lilly and Company, Janssen, Novartis, UCB Pharma, Galderma, Janssen-Cilag, Kyowa Kirin, LEO Pharma, L'Oréal, New Bridge, Novartis, Pfizer, Regeneron, Sanofi, Target-RWE, UCB Pharma, and Vichy; received grants from AbbVie, LEO Pharma, and Novartis. MG: Investigator, speaker, consultant or advisory board member for AbbVie, Amgen, Anapsylbio, Arcutis, Arista, Astria, Bausch Health, Boehringer Ingelheim, Bristol Myers Squibb, Dermavant, Dermira, Eli Lilly and Company, Galderma, GSK, Incyte, Janssen, Kyowa Kirin, MedImmune, Meiji, Merck, Moonlake Immunotherapeutics, Nimbus, Novartis, Pfizer, Regeneron, Reistone, Sanofi Genzyme, Sun Pharma, Takeda, Tarsus, UCB Pharma, Union, and Vertex; <sup>YO</sup>: Received research grants from Eisai, Maruho, Shiseido, and Torii Pharmaceuticals; consulting and advisory board agreements from AbbVie, Amgen, Boehringer Ingelheim, Bristol Myers Squibb, Celgene, Eisai, Eli Lilly and Company, Janssen, Jimto, Kyowa Kirin, LEO Pharma, Maruho, Novartis, Pfizer, Sanofi, Sun Pharma, Taito, Tanabe-Mitsubishi, Torii Pharmaceutical, and UCB Pharma; clinical trials sponsored by AbbVie, Amgen, Boehringer Ingelheim, Bristol Myers Squibb, Celgene, Eli Lilly and Company, Janssen, LEO Pharma, Maruho, Pfizer, Sun Pharma, and UCB Pharma. BS: Consultant (honoraria) for AbbVie, Acelyrin, Almar, Almirall, Alumis, Amgen, Arcutis, Arena, Arista, Asana, Boehringer Ingelheim, Scientific Oncology, Capital One, Celtrion, CorEviTas, Dermavant, Eli Lilly and Company, ImagenBio, Janssen, Kangpu Pharmaceuticals, LEO Pharma, Maruho, Meiji Seika Pharma, Protagonist, Monte Carlo, Novartis, Pfizer, Rapt, Regeneron, Sanofi Genzyme, SG Cowen, Sun Pharma, Takeda, UCB Pharma, Union Therapeutics, Vertex, and VIV Therapeutics; stock options from Connect Biopharma, Mindera Health; speaker for AbbVie, Arcutis, Dermavant, Eli Lilly and Company, Incyte, Janssen, Regeneron, and Sanofi Genzyme; scientific co-director (consulting fee) for CorEviTas Psoriasis Registry; investigator for CorEviTas Psoriasis Registry; editor-in-chief (honoraria) of *Journal of Psoriasis and Psoriatic Arthritis*. LP, DD, JMLP: Employees and shareholders of UCB Pharma. PG: Consultant for AbbVie, Abogen, Almirall, Celgene, Eli Lilly and Company, Janssen, LEO Pharma, Merck, MSD, Novartis, Otsuka, Pfizer, Pierre Fabre, Sanofi, and UCB Pharma. **Acknowledgments:** These studies were funded by UCB Pharma. We would like to thank the patients and their caregivers in addition to the investigators and their teams who contributed to these studies. The authors acknowledge Susanne Wiegartz, MSc, UCB Pharma, Monheim am Rhein, Germany, and Joe Dixon, PhD, UCB Pharma, Slough, UK, for publication coordination; Sana Year, PhD, Costello Medical, Manchester, UK, for medical writing support and editorial assistance, and Danielle Hart of the Creative Team at Costello Medical, London, UK, for graphic design assistance. All costs associated with development of this poster were funded by UCB Pharma.



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