

VISIBLE COHORT B: GUSELKUMAB DEMONSTRATED SCALP CLEARANCE AND IMPROVED HEALTH-RELATED QUALITY OF LIFE THROUGH WEEK 48 IN PARTICIPANTS WITH MODERATE-TO-SEVERE SCALP PSORIASIS ACROSS ALL SKIN TONES



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BACKGROUND/OBJECTIVE



VISIBLE is an ongoing Phase 3b, multicenter, randomized, double-blinded, placebo (PBO)-controlled study of guselkumab (GUS) for the treatment of participants with moderate-to-severe plaque psoriasis (PsO) across all skin tones



VISIBLE is comprised of 2 cohorts:



moderate-to-severe plaque PsO

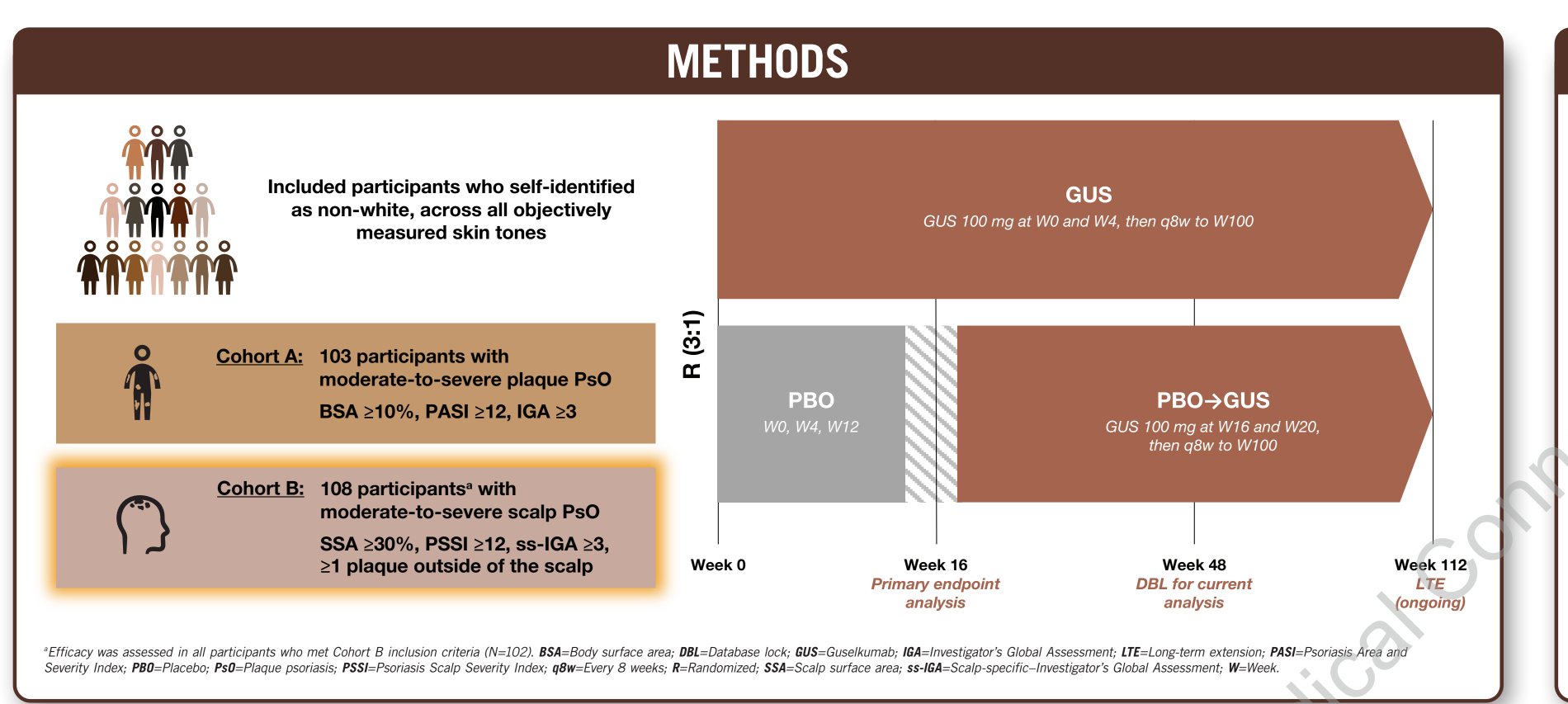
Cohort A: participants with

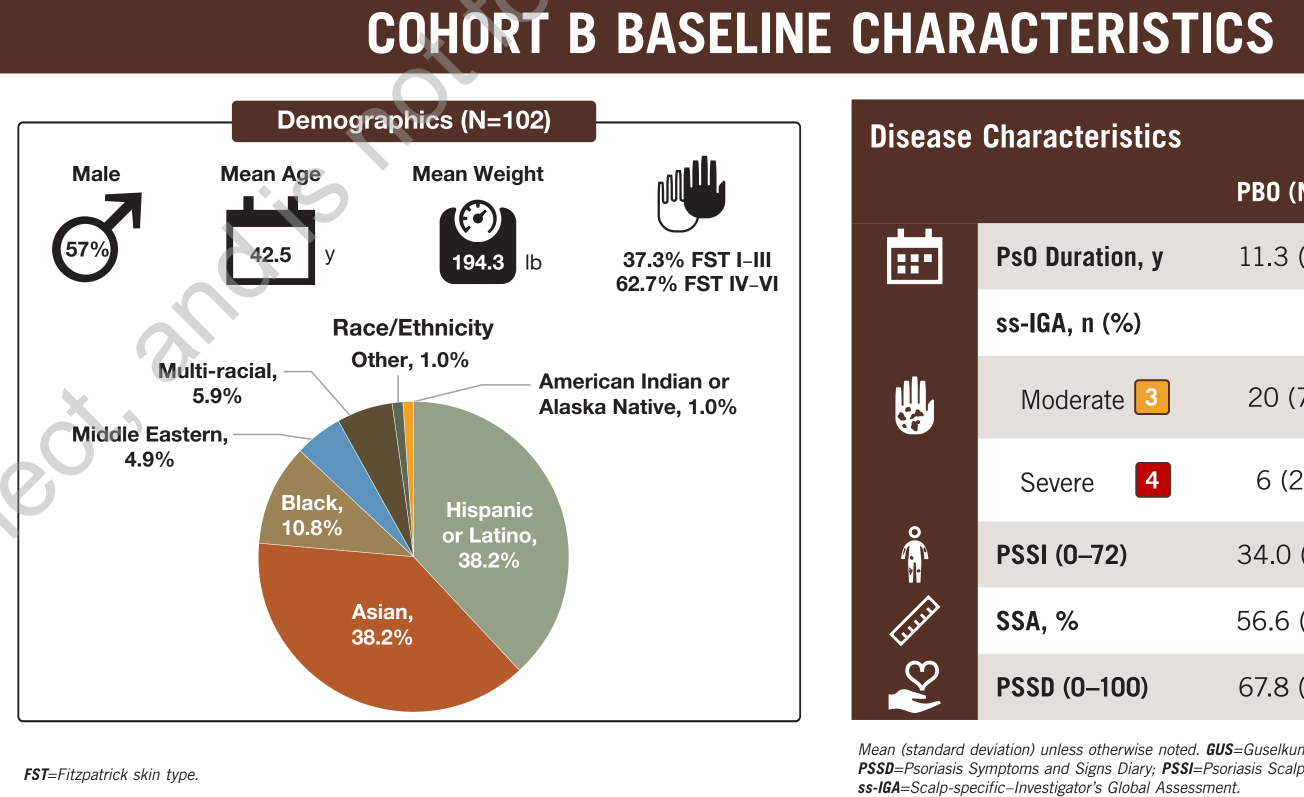


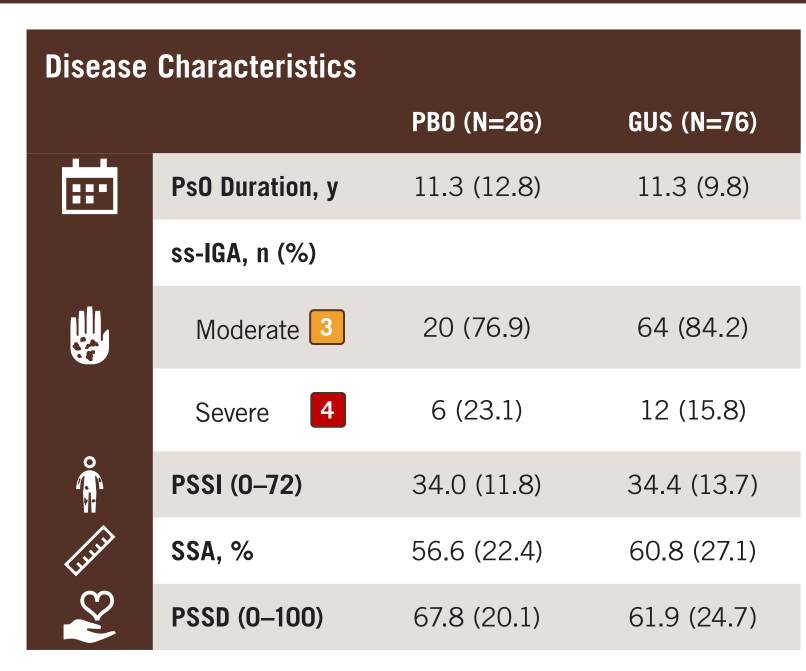
Cohort B: participants with moderate-to-severe scalp PsO



VISIBLE evaluated the efficacy and safety of GUS for treating scalp PsO in Cohort B participants through



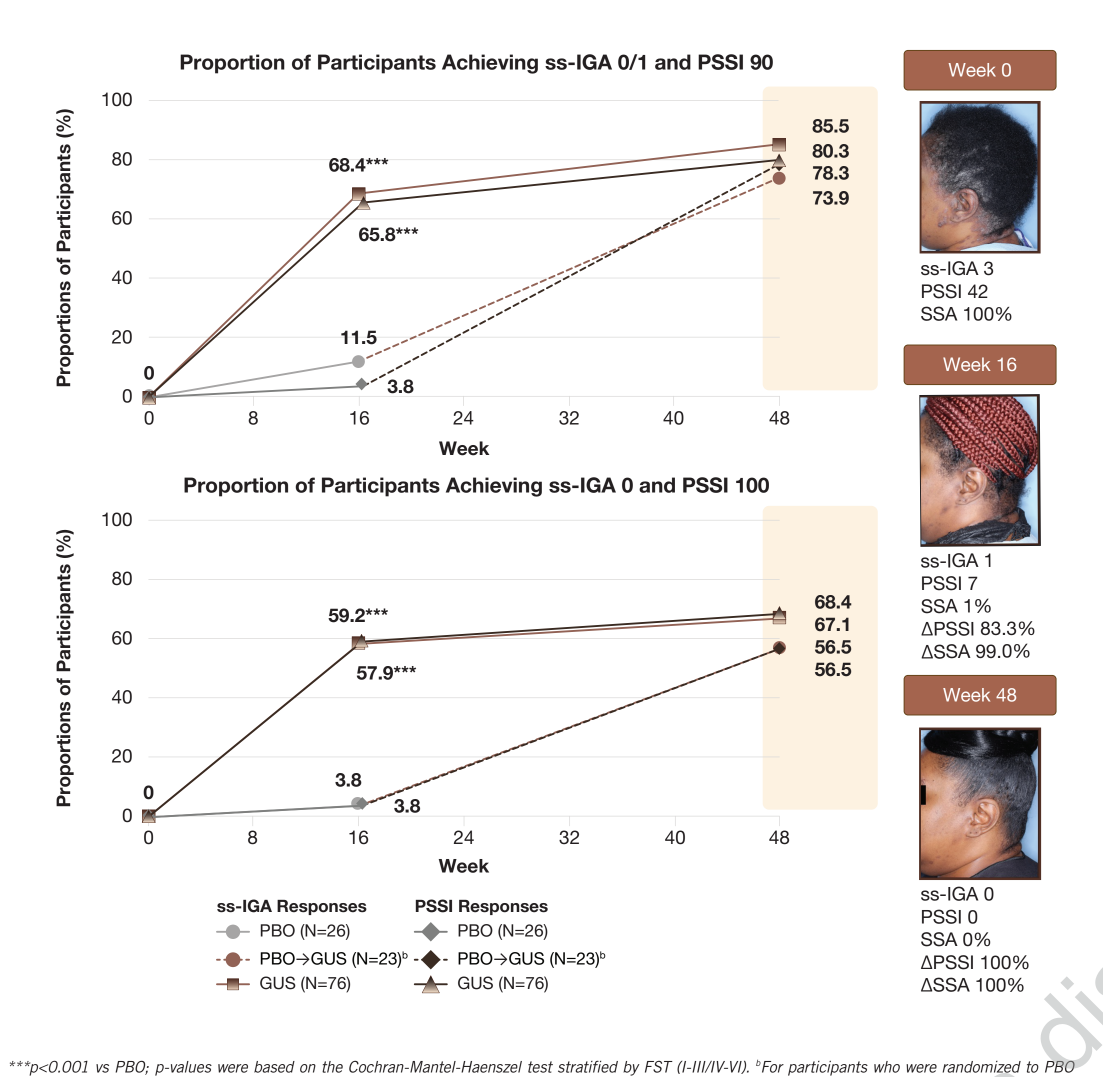




Mean (standard deviation) unless otherwise noted. GUS=Guselkumab; PBO=Placebo; PsO=Plaque psoriasis; **PSSD**=Psoriasis Symptoms and Signs Diary; **PSSI**=Psoriasis Scalp Severity Index; **SSA**=Scalp surface area;

RESULTS

Significantly greater proportions of GUS-randomized vs PBO-randomized participants achieved ss-IGA and PSSI endpoints at Week 16, and response rates were sustained or increased at Week 48



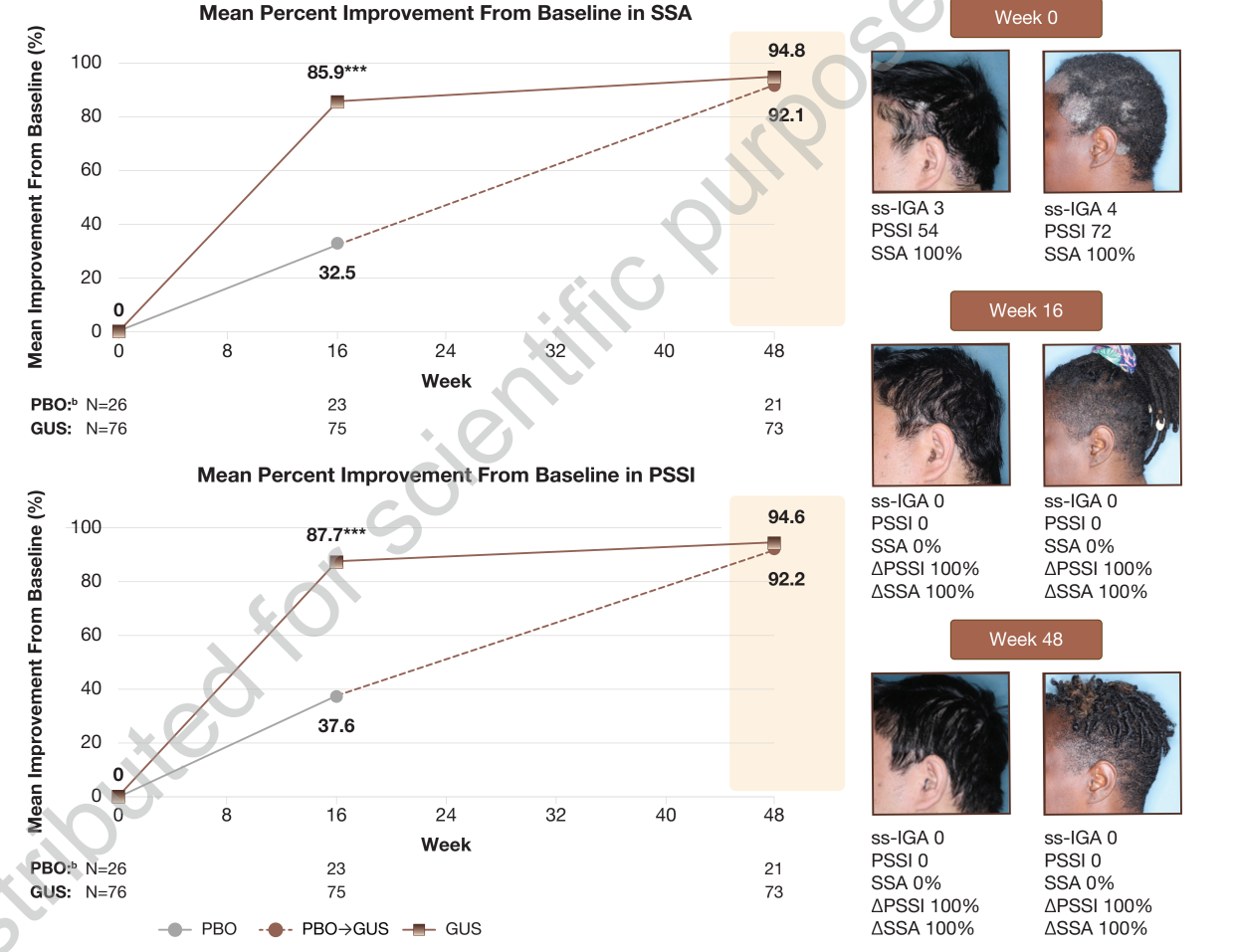
at Week O, only those who crossed over to GUS at or after Week 16 were included at Week 48. Participants who discontinued study agent due to lack of

with missing data were considered non-responders at that time point. **FST**=Fitzpatrick skin type; **GUS**=Guselkumab; **PBO**=Placebo; **PsO**=Plaque psoriasis;

PSSI=Psoriasis Scalp Severity Index; **SSA**=Scalp surface area; **ss-IGA**=Scalp-specific–Investigator's Global Assessment; Δ =Mean improvement.

efficacy, worsening of PsO, or use of a prohibited PsO treatment prior to designated visit were considered non-responders from that point forward. Participants

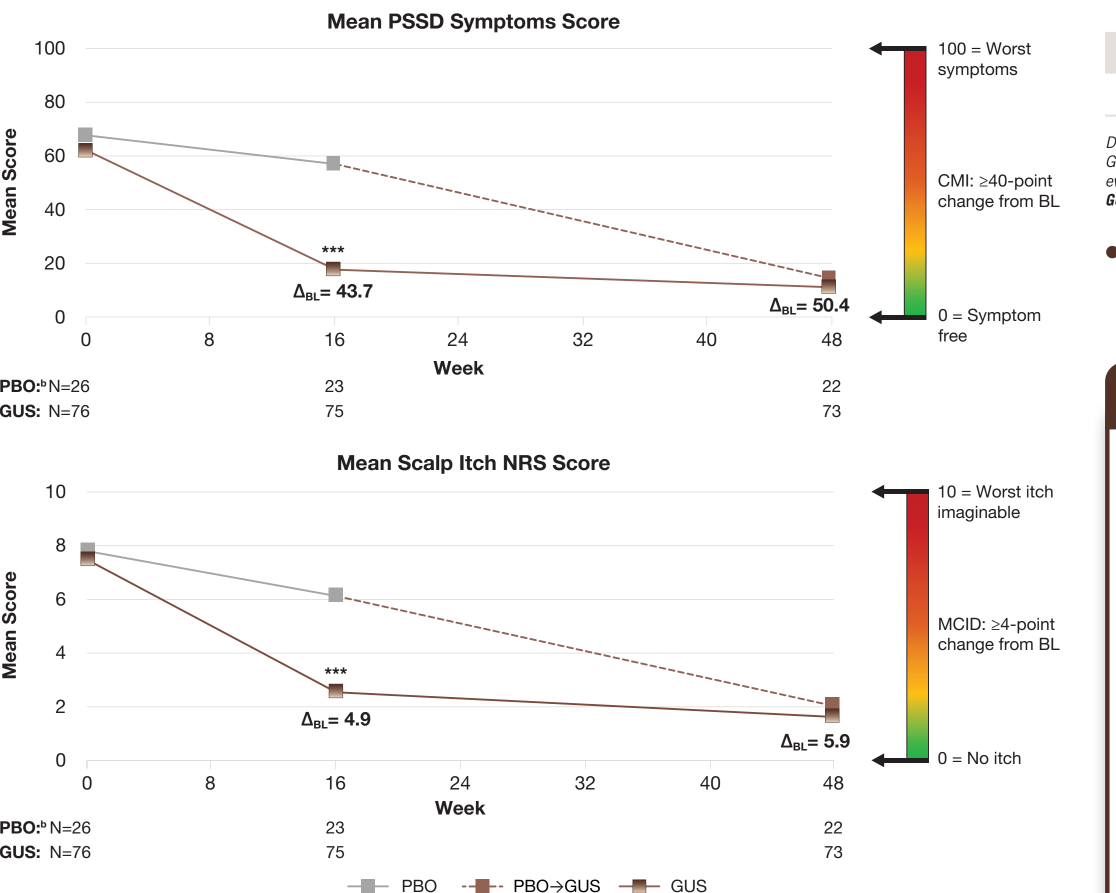
Among GUS-randomized participants, mean percent improvement in SSA and PSSI was >85% at Week 16 and increased to ~95% at Week 48



***p<0.001 vs PBO; p-values were based on the mixed-effect model for repeated measures (MMRM). For participants who were randomized to PBO at Week 0, only those who crossed over to GUS at or after Week 16 were included. When participants who discontinued study agent due to lack of efficacy, worsening of PsO, or use of a prohibited PsO treatment, O change from baseline was assigned from that point onward. Missing data were not imputed. **GUS**=Guselkumab; **PBO**=Placebo; **Ps0**=Plaque psoriasis; **PSSI**=Psoriasis Scalp Severity Index; **SSA**=Scalp surface area; **ss-IGA**=Scalp-specific−Investigator's Global Assessment, **∆**=Mean improvement.

Among GUS-randomized participants, the overall PSSD Symptoms Score and the Scalp Itch Numeric Rating Scale (NRS) Score showed significant mean improvements from baseline at Week 16, which were maintained at Week 48

• The proportion of GUS-randomized participants achieving an overall PSSD Symptoms Score of O^c increased from 21% at Week 16 to 35% at Week 48, and the proportion achieving ≥4-point improvement from baseline in the Scalp Itch NRS Scored increased from 69% to 81%



***p<0.001 vs PBO; p-value was based on the MMRM for PSSD Symptoms Score and analysis of covariance for Scalp Itch NRS Score. When participants discontinued study agent due to lack of efficacy, worsening of PsO, or use of a prohibited PsO treatment, O change from baseline was assigned from that point onward. Missing data were not imputed. For participants who were randomized to PBO at Week 0, only those who crossed over to GUS at or after Week 16 were included. GUS-randomized participants with baseline PSSD Symptoms Score ≥1. Participants with baseline PSSD Itch Symptom Score ≥4. Participants who discontinued study agent due to lack of efficacy, worsening of PsO, or use of a prohibited PsO treatment prior to designated visit were considered non-responders from that point forward. Participants with missing data were considered non-responders at that time point. **BL**=Baseline; **CMI**=Clinically meaningful improvement; **GUS**=Guselkumab; **MCID**=Minimal clinically important difference; **NRS**=Numeric Rating Scale; **PBO**=Placebo; **Ps0**=Plaque psoriasis; **PSSD**=Psoriasis Symptoms and Signs Diary; Δ_{BL} =Mean improvement from baseline.

Key Safety Information		
	PBO→GUS ^e (Weeks 16-48)	GUS (Weeks 0–48)
Safety analysis set, N	24	81
Average duration of follow-up (weeks)	31.1	47.7
Participants with ≥1 AE	9 (37.5)	51 (63.0)
Participants with ≥1 AE leading to discontinuation of study agent	0	0
Participants with ≥1 SAE	0	2 (2.5) ^f
Participants with ≥1 injection-site reaction	O	1 (1.2)
Infections	4 (16.7)	27 (33.3)
Serious infections	0	0

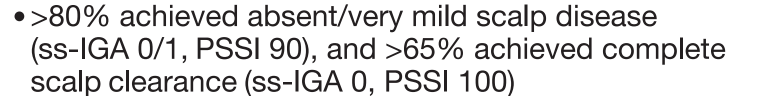
Data shown are n (%), unless otherwise indicated. "Includes only PBO participants who crossed over to receive GUS. 'The SAEs in GUS-treated participants were 1 event each of angina pectoris and pancreatitis. Participants were counted only once for any given event, regardless of the number of times they experienced the event. AEs were coded using MedDRA version 25.1. AE=Adverse event; GUS=Guselkumab; MedDRA=Medical dictionary for regulatory activities; PBO=Placebo; SAE=Serious adverse event.

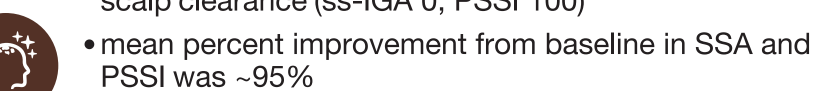
• Through Week 48, there were no cases of death, malignancy, active tuberculosis, major adverse cardiac event, inflammatory bowel disease, or serum-like sickness or anaphylaxis

CONCLUSIONS



At Week 48, among GUS-randomized participants in Cohort B of the VISIBLE study:





 clinically meaningful improvements in the mean overall PSSD Symptoms Score and the mean Scalp Itch NRS Score were achieved



No new safety signals were identified



These results demonstrate that GUS is a highly effective and durable treatment for moderate-to-severe scalp PsO participants across all objectively measured skin tones, with sustained or improved responses through Week 48

PRESENTED AT: 12th Annual Meeting of International Dermatology Outcomes Measures (IDEOM); Washington, D.C., USA; June 27–28, 2025. Acknowledgments: This presentation was supported by Johnson & Joh being sponsored by AbbVie, Castle, CorEvitas, Dermavant, Galderma, Mindera, and Pfizer, SUN Pharma, and UCB. SS: receives honoraria or research funding from Novartis, Pfizer, SUN Pharma, and UCB. SS: receives honoraria or research funding from Novartis, Pfizer, SUN Pharma, and Teoxane. TA, DC, 100 Pharma, and Teoxane. TA, DC, 101 Pharma, Almirall, Alumis, Amgen, Arcutis, Boehringer-Ingelheim, Bristol Myers Squibb, Eli Lilly, Ianssen, Hoberg, Nielsen, Pfizer, SUN Pharma, and Teoxane. TA, DC, 101 Pharma, Almirall, Alumis, Amgen, Arcutis, Pfizer, SUN Pharma, Indicate the search grants from AbbVie, Aclaris, Almirall, Alumis, Amgen, Arcutis, Pfizer, SUN Pharma, Indicate the search grants from AbbVie, Aclaris, Almirall, Alumis, Amgen, Arcutis, Pfizer, SUN Pharma, Indicate the search grants from AbbVie, Aclaris, Almirall, Alumis, Amgen, Arcutis, Pfizer, SUN Pharma, Indicate the search grants from AbbVie, Aclaris, Almirall, Alumis, Amgen, Arcutis, Pfizer, SUN Pharma, Indicate the search grants from AbbVie, Aclaris, Almirall, Alumis, Amgen, Arcutis, Pfizer, SUN Pharma, Indicate the search grants from AbbVie, Aclaris, Almirall, Alumis, Amgen, Arcutis, Pfizer, SUN Pharma, Indicate the search grants from AbbVie, Aclaris, Almirall, Alumis, Amgen, Arcutis, Pfizer, SUN Pharma, Indicate the search grants from AbbVie, Aclaris, Almirall, Alumis, Amgen, Amg TM, and S0: are employees and stockholders of Johnson & Centocor, Coherus, Dermira, Eli Lilly, Forward, Galderma, Eli Lilly, MC2, MedX, Novartis, Ortho Dermatologics, Pfizer, Regeneron/Sanofi, SUN Pharma, Takeda, and UCB. and Venon. GH: is a consultant, speaker, or received research support from AbbVie, Amgen, Athenex, Beiersdorf, Bristol Myers Squibb, Boehringer Ingelheim, Bond Avillion, Castle Biosciences, Celgene, CeraVe, Dermatologics, Pfizer, Regeneron/Sanofi, SUN Pharma, Takeda, and UCB. SCT: has received honoraria/stock options serving as an advisor/consultant and/or speaker for AbbVie, Arcutis, Beiersdorf, Biorez, Bristol Myers Squibb, Cara, Dior, Eli Lilly, EPI, Evolus, Galderma, GloGetter, Hugel America, Janssen, Johnson & Johnson & Johnson & Johnson & In Scrientis US, UCB, and Vichy; has received honoraria/stock options serving as an author/received honoraria/stock options serving as an advisor/consultant and/or speaker for AbbVie, Arcutis, Armis, Avita, Beiersdorf, Biorez, Bristol Myers Squibb, Cara, Dior, Eli Lilly, EPI, Evolus, Galderma, GloGetter, Hugel America, Janssen, Johnson & Jo Dermatologic Research, British Journal of Dermatology (peer reviewer), Cutis, and Practical Dermatology; served as an investigator for Concert Pharmaceuticals, Croma-Pharma, Eli Lilly, and Pfizer. PREVIOUSLY PRESENTED AT: American Academy of Dermatology (AAD) Annual Meeting; Orlando, FL, USA; March 7–11, 2025.