# Real-World On-Label Treatment Persistence Through 24 Months in Biologic-Naïve and Biologic-Experienced Patients With Psoriatic Arthritis: Comparison of Guselkumab versus Subcutaneous Interleukin-17A Inhibitors



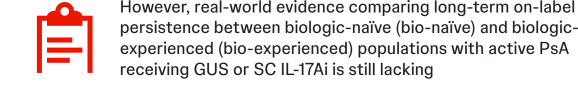
Philip J Mease<sup>1,2</sup>, Jessica Walsh<sup>3,4</sup>, Timothy P Fitzgerald<sup>5</sup>, Soumya D Chakravarty<sup>5,6</sup>, Elizabeth Adamson<sup>5</sup>, Bruno Emond<sup>7</sup>, Carmine Rossi<sup>7</sup>, Samuel Schwartzbein<sup>7</sup>, Kana Yokoji<sup>7</sup>, Yuxi Wang<sup>7</sup>, Patrick Lefebvre<sup>7</sup>, Dominic Pilon<sup>7</sup>, Shikha Singla<sup>8</sup>, Joseph F Merola<sup>9</sup> ¹Rheumatology Research, Providence Swedish Medical Center, Seattle, WA, USA; ²University of Washington School of Medicine, Salt Lake City, UT, USA; ⁴University of Utah Health, Salt Lake City, UT, USA; ⁵Janssen Scientific Affairs, LLC, a Johnson & Johnson company, Horsham, PA, USA; <sup>6</sup>Drexel University College of Medicine, Philadelphia, PA, USA; <sup>9</sup>Department of Dermatology, and Department of Medicine, Division of Rheumatology, UT Southwestern Medical Center, Dallas, TX, USA

## Background

(FDA) for the treatment of active psoriatic arthritis (PsA) in - FDA-approved dosing regimen<sup>1</sup> (on-label): GUS 100 mg at

> week 0, week 4, then every 8 weeks persistence for patients (pts) with PsA initiating on-label

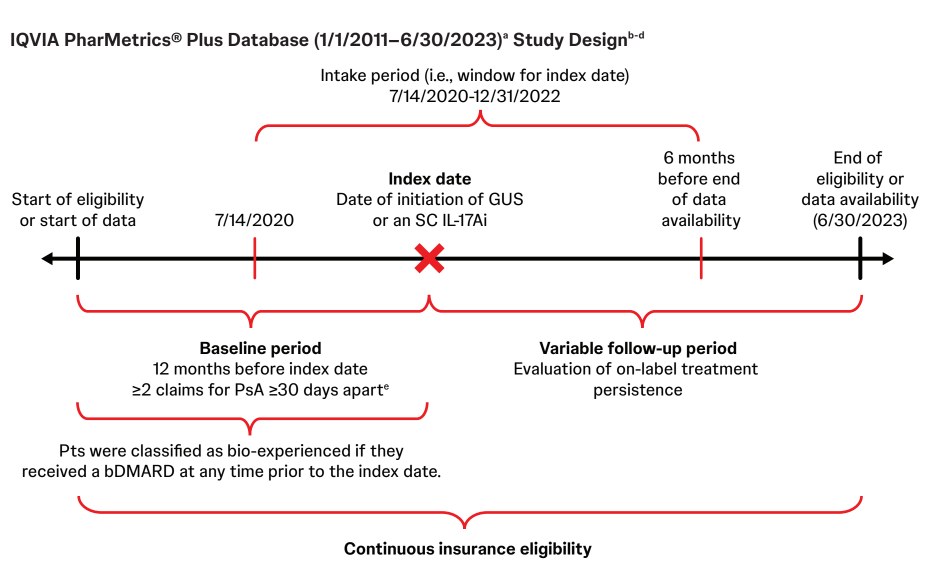
 Pts receiving GUS were significantly (~1.5x) more likely to remain persistent through 24 months



**Objectives** 

treatment persistence through 24 months in bio-naïve and bio-experienced pts with active PsA newly initiating on-label therapy with either GUS or an initial SC IL-17Ai

## Methods



employers, payers, providers, and therapy areas. <sup>b</sup>A validated algorithm for identifying pts with PsA in US claims data was used: ≥2 claims with a PsA diagnosis (ICD-10-CM: L40.5x) ≥30 days apart and

≥1 prescription claim for a PsA-related medication (i.e., GUS or SC IL-17Ai). °Pts could be bio-naïve or bio-experienced during baseline but were naïve to treatment with GUS or SC IL-17Ai agents. ⁴Pts in the

SC IL-17Ai cohort were newly initiated within the class. Diagnoses for PsA include claims on the index date. ICD-10-CM=International Classification of Disease, 10th revision, Clinical Modification.

eligibility before index date

ndex date: 1st GUS or SC IL-17Ai clair during intake period (7/14/2020-

• PsA pt identification: ≥2 PsA Dx (ICD-10-CM code L40.5x) ≥30 days apart within 12 months prior to or on the index date, and ≥1 claim for either GUS or first SC IL-17Ai<sup>4</sup> ≥12 months of continuous health insurance

 ≥18 years of age No claims for other conditions for which GUS or IL-17Ai are approved or other potentially confounding diseases<sup>b</sup>

Pts were classified as bio-experienced if they had ≥1 claim for a PsA-indicated biologic disease modifying antirheumatic drug (bDMARD) at any time prior to the index date, and bio-naïve

**Censoring and Imputations** Censoring: On earliest of first off-label claim or last

 Baseline demographic and disease characteristics (12 months pre-index): day of index agent supply preceding end of follow-up period if discontinuation was not observed Balanced between the GUS and SC IL-17Ai cohorts separately for bio-naïve and bio-experienced pts using propensity score weighting Medical Claims<sup>1,5</sup>

> No treatment discontinuation or dose modification relative to US FDA-approved labeling Proportion of pts determined using weighted GUS vs SC IL-17Ai cohorts compared using weighted Cox proportional hazard models

On-label persistence up to 24 months post-index:

administration per label after induction. Sensitivity analyses were conducted based on 1x the

## **Key Takeaways**

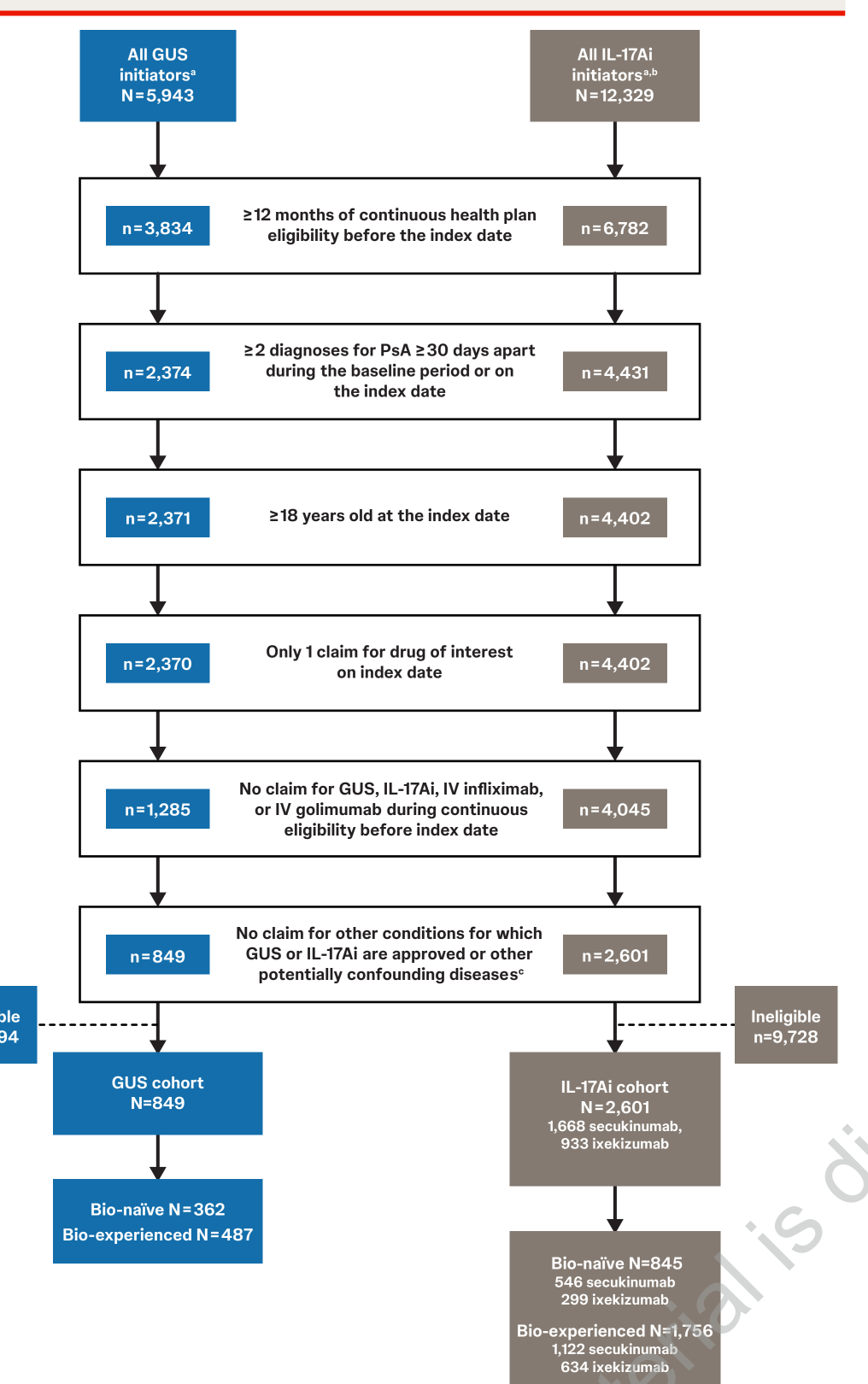
First real-world claims data analysis of on-label treatment persistence over 24 months in bio-naïve and bio-experienced pts with active PsA newly initiated on GUS vs initial SC IL-17Ai per US FDA-approved labeling

Pts in the GUS cohort were significantly more likely to remain persistent on treatment through 24 months in both the bio-naïve and bio-experienced populations

Higher long-term on-label persistence may improve disease management outcomes in pts with active PsA initiating GUS<sup>7</sup>, regardless of prior biologic treatment status

## Results

The GUS and SC IL-17Ai cohorts, respectively, included 362 and 845 bio-naïve pts and 487 and 1,756 bio-experienced pts



°f° GUS or SC IL-17Ai claim during intake period (7/14/2020-12/31/2022). The SC IL-17Ai cohort is defined as pts with an index claim for an SC IL-17Ai (ie, ixekinumab). Assessed during the 12-month baseline period. IV=intravenous.

Weighted baseline demographic and clinical characteristics were similar between the GUS and SC IL-17Ai cohorts among bio-naïve and bio-experienced pts

Bio-naïve

		Bio-naive		Bio-expe	
Weighted Baseline Demographics and Clinical Characteristics <sup>a</sup>		GUS (N=362)	SC IL-17Ai (N=845)	GUS (N=487)	SC IL-17Ai (N=1,756)
Demographics					
	<b>Age at index date (years),</b> Mean ± SD [median]	49.5 ± 11.6 [50.8]	49.6 ± 11.9 [51.3]	49.7 ± 10.6 [50.7]	49.4 ± 11.0 [50.5
	Female	58.6	58.6	60.3	60.3
	Insurance type at index date				G
·Ππ.	Preferred provider organization	77.7	77.7	78.5	78.5
	Health maintenance organization	10.7	10.7	11.4	11.4
	Other <sup>b</sup>	11.6	11.6	10.1	10.1
	Year of index date				
	2020	12.2	12.2	11.0	11.0
	2021	40.2	40.2	39.2	39.2
	2022	47.7	47.7	49.8	49.8
haracteristics					
	Months between latest observed PsA diagnosis and index date, Mean ± SD [median]	1.4 ± 1.7 [0.8]	1.4 ± 1.6 [0.9]	1.2 ± 1.5 [0.7]	1.2 ± 1.3 [0.8]
W	<b>Quan-CCI,</b> Mean ± SD [median]	0.6 ± 1.2 [0.0]	0.6 ± 1.2 [0.0]	0.5 ± 1.2 [0.0]	0.6 ± 1.3 [0.0
~~	Comorbidities				
	Hyperlipidemia	37.8	37.5	33.6	33.6
20	Osteoarthritis	27.8	29.2	29.9	29.9
= %	Diabetes	15.7	14.3	13.6	14.6
	Peripheral vascular disease	2.0	2.3	2.4	2.4
	Psoriasis	89.2	89.2	81.3	81.3
	Smoking	12.1	12.7	9.0	9.0
ledication Use <sup>c</sup>					
9	<b>bDMARDs</b> <sup>d</sup>	0	0	85.0	85.1
	0	0	0	15.0	14.9
<del></del>	1	0	0	74.3	73.9
+	≥2	0	0	10.7	11.2
	csDMARDs <sup>e</sup>	24.2	24.2	27.4	27.0
	tsDMARDs <sup>f</sup>	29.4	29.2	16.7	16.3
	Corticosteroids	71.6	68.5	72.4	74.0

Bio-naïve and bio-experienced pts treated with GUS were significantly more likely to remain persistent with on-label treatment through 24 months vs. pts treated with SC IL-17Ai

#### Bio-naïve pts:

Coding System code for SC IL-17Ai in medical claims. Pharmacy claims for SC IL-17Ai are typically

consistent with approved labeling; therefore, reported days supply was used for SC IL-17Ai and no

**Pharmacy Claims** 

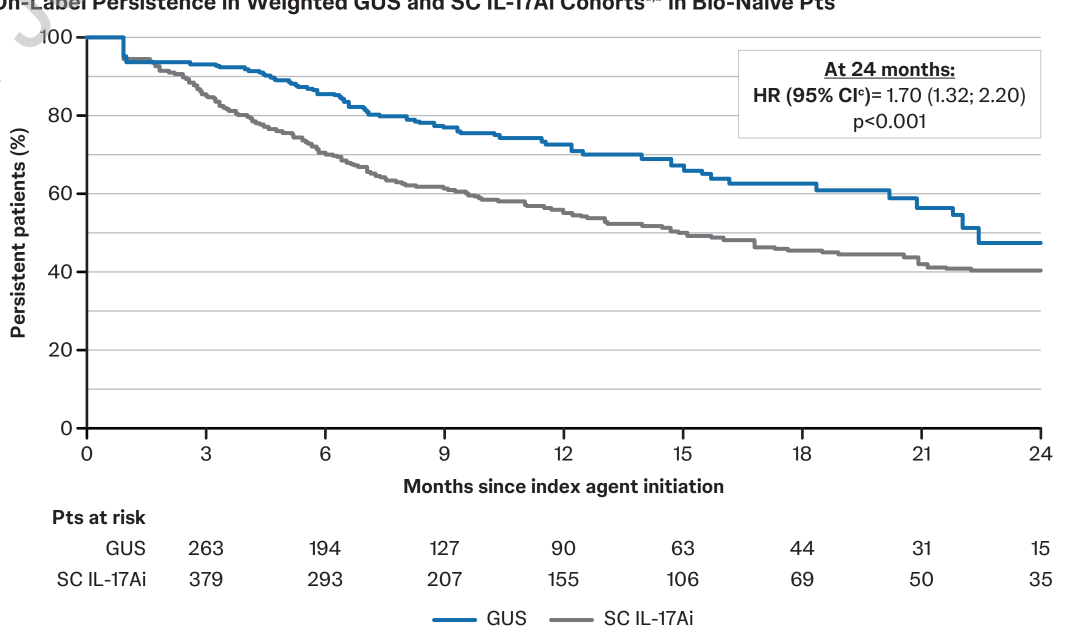
**Bio-experienced** 

- On-label persistence at 24 months: 47.5% with GUS vs 40.3% with SC IL-17Ai - Sensitivity analyses for bio-naive pts demonstrated similar trends

discontinuation gap of 112 days. **KM**=Kaplan-Meier.

- Median time to discontinuation: 22.4 months with GUS vs 14.9 months with SC IL-17Ai

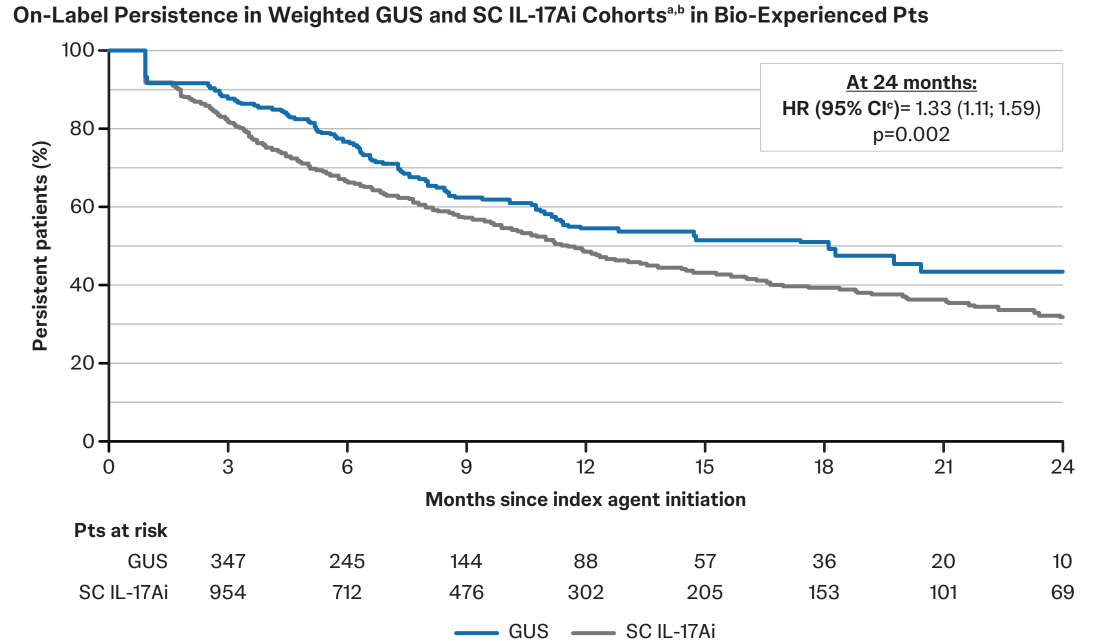
On-Label Persistence in Weighted GUS and SC IL-17Ai Cohorts<sup>a,b</sup> in Bio-Naïve Pts



### Bio-experienced pts

- On-label persistence at 24 months: 43.3% with GUS vs 32.0% with SC IL-17Ai

 Sensitivity analyses for bio-experienced pts demonstrated similar trends - Median time to discontinuation: 18.1 months with GUS vs 11.6 months with SC IL-17Ai



Primary analysis: discontinuation was defined as having a gap in treatment of more than twice the duration of days of supply for a claim (i.e., 2 x 56 = 112 days for GUS or 2 x 28 = 56 days for IL-17Ai). Pts with dose changes inconsistent with

FDA-approved dosing were censored as of the first dose change. A weighted Cox proportional hazards model was used to compare on-label persistence between cohorts. CI=confidence interval; HR=hazard ratio.

In bio-naïve and bio-experienced pts, GUS was associated with significantly higher on-label persistence vs SC IL-17Ai at each time point assessed (6/12/18/24 months)

#### On-label persistence through 24 months in weighted GUS and SC IL-17Ai bio-naïve and bio-experienced cohorts<sup>a</sup> Primary analysis (2x duration)

Cox proportional hazards model <sup>b</sup>	6 months	12 months	18 months	24 months		
Bio-naïve cohorts						
Pts at risk, n (%)°						
GUS (N=362)	194 (53.6)	90 (25.0)	44 (12.1)	15 (4.3)		
SC IL-17Ai (N=845)	293 (34.7)	155 (18.3)	69 (8.2)	35 (4.2)		
Hazard ratios (95% CI)	2.18 (1.54; 3.09)	1.92 (1.44; 2.55)	1.83 (1.40; 2.38)	1.70 (1.32; 2.20)		
Chi-square p-value	<0.001	<0.001	<0.001	<0.001		
KM Persistence, % (95% CI)						
GUS	85.7 (76.1; 91.6)	72.6 (62.8; 80.3)	62.6 (50.1; 72.8)	47.5 (22.7; 68.7)		
SC IL-17Ai	70.6 (63.2; 76.8)	55.2 (46.0; 63.4)	45.2 (33.6; 56.1)	40.3 (26.2; 54.0)		
Log-rank test p-value	<0.001	<0.001	<0.001	<0.001		
Bio-experienced cohorts						
Pts at risk, n (%)°						
GUS (N=487)	245 (50.2)	88 (18.1)	36 (7.5)	10 (2.1)		
SC IL-17Ai (N=1,756)	712 (40.6)	302 (17.2)	153 (8.7)	69 (3.9)		
Hazard ratios (95% CI)	1.52 (1.21; 1.90)	1.28 (1.07; 1.54)	1.34 (1.12; 1.61)	1.33 (1.11; 1.59)		
Chi-square p-value	<0.001	0.007	0.001	0.002		
KM Persistence, % (95% CI)						

### **Strengths and Limitations**

SC IL-17Ai

Log-rank test p-value

A case-finding algorithm validated in US claims data was used to identify pts with active PsA<sup>4</sup>

Propensity score weights were used to obtain a balanced sample. Weights were estimated using a multivariable logistic regression model. Baseline covariates included several demographic and clinical characteristics. Weighted Cox proportional hazard models were used to compare risk of discontinuation between the GUS and SC II-17Ai cohorts "Pts at risk of having the event are pts who have not had the event and have not been lost to follow-up at that point in time

- Baseline demographic and disease characteristics between the GUS and SC IL-17Ai cohorts were balanced • Limitations:

- Claims data do not ensure treatments are taken as prescribed
- Claims data do not provide treatment effectiveness nor reasons for discontinuation

67.1 (62.5; 71.3)

54.4 (45.2; 62.7) 51.0 (40.6; 60.5) 43.3 (26.1; 59.3)

39.3 (31.9; 46.6)

32.0 (22.1; 42.3)

0.002