On-Label Persistence Through 24 Months Among Patients With Psoriatic Arthritis Initiating Guselkumab or Subcutaneous TNF Inhibitors

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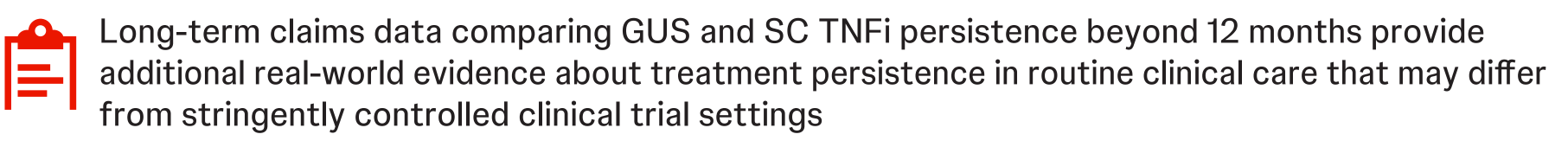
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Background

- Suselkumab (GUS), a fully human IL-23 p19-subunit inhibitor, was approved by the US Food and Drug Administration (FDA) for the treatment of active psoriatic arthritis (PsA) in July 2020
 - FDA-approved dosing regimen¹ (on-label): GUS 100 mg at week 0, week 4, then every 8 weeks

A previous claims-based analysis compared on-label persistence for patients (pts) with PsA initiating treatment with on-label GUS or their first subcutaneous (SC) tumor necrosis factor inhibitor (TNFi)²

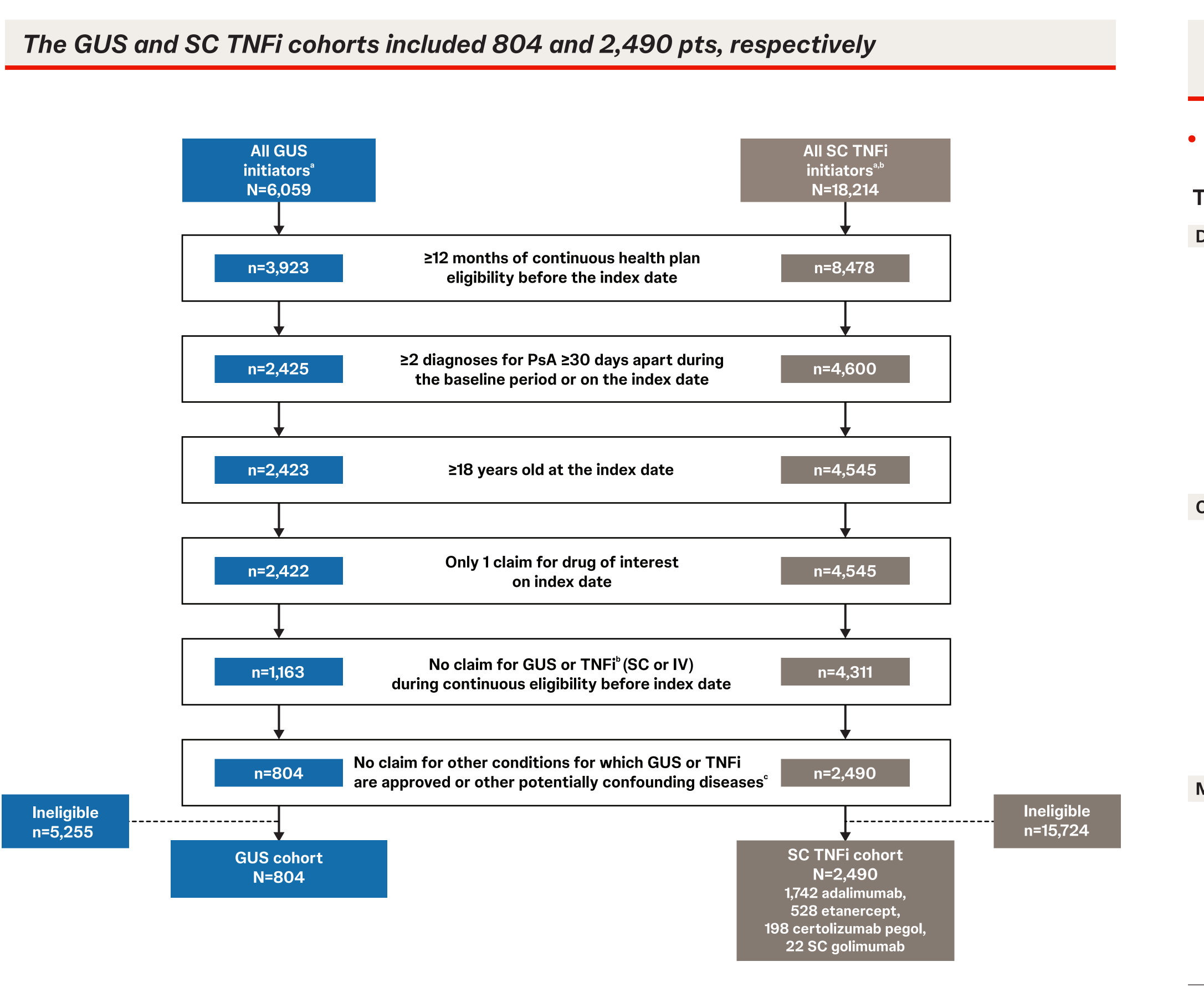
- Pts receiving GUS were significantly (~3x) more likely to remain persistent through 12 months



Objectives

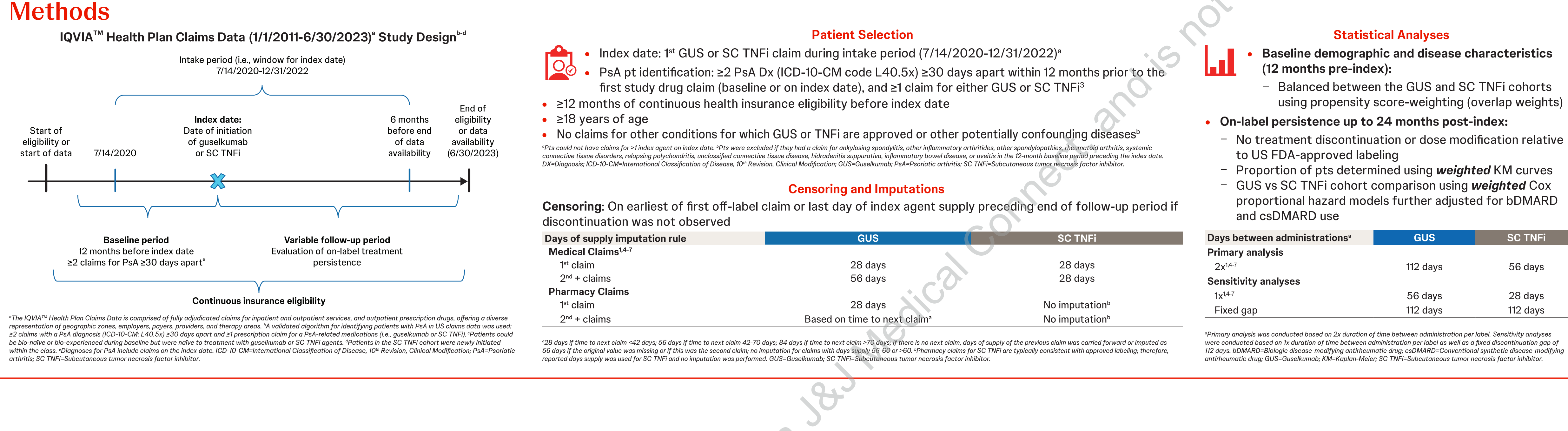
This study utilized health plan claims data to compare treatment persistence through 24 months between pts with active PsA newly initiating the on-label GUS dosing regimen and those starting an initial SC TNFi

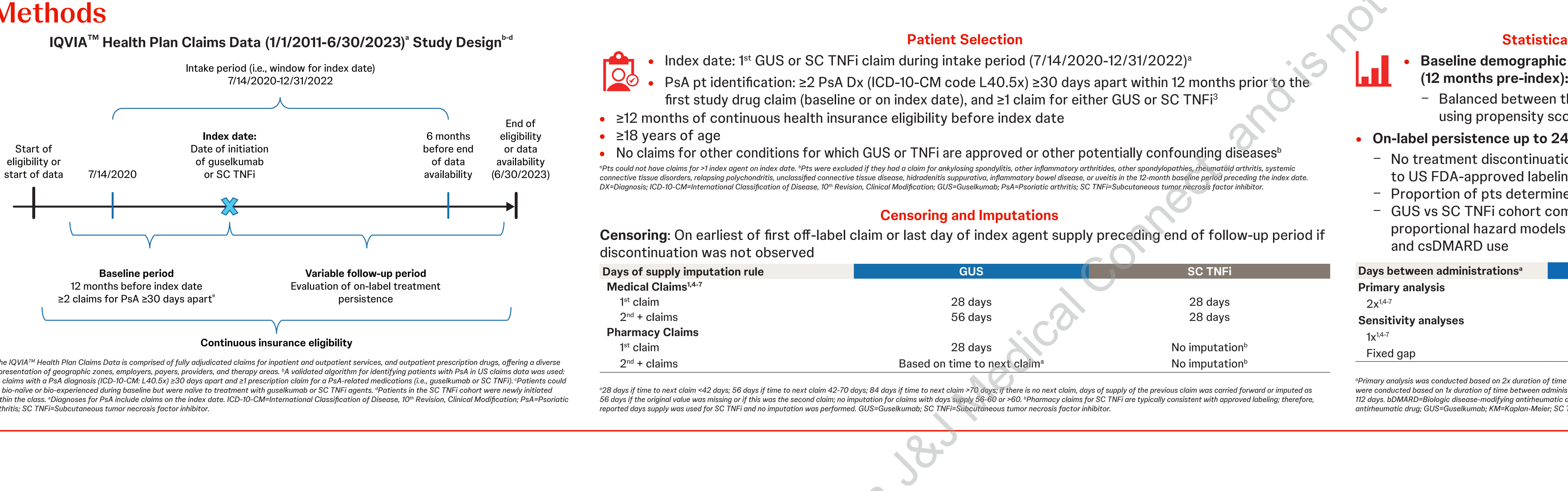
Results



^a1st GUS or SC TNFi claim during intake period (7/14/2020-12/31/2022). ^bThe SC TNFi cohort is defined as pts with an index claim for an SC TNFi (i.e., adalimumab, certolizumab pegol, etanercept, or SC golimumab). ^cAssessed during the 12-month baseline period. GUS=Guselkumab; IV=Intravenous; PsA=Psoriatic arthritis; SC TNFi=Subcutaneous tumor necrosis factor inhibitor.







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≥2 claims with a PsA dia
be bio-naïve or bio-exper
within the class. ^e Diagno
arthritis; SC TNFi=Subcu

modifying antirheumatic drug.

Weighted baseline demographic and clinical characteristics were similar between cohorts, except for prior bDMARD and csDMARD use

• 55.1% in the GUS cohort and 12.8% in the SC TNFi cohort had received ≥1 bDMARD at any time before the index date^a

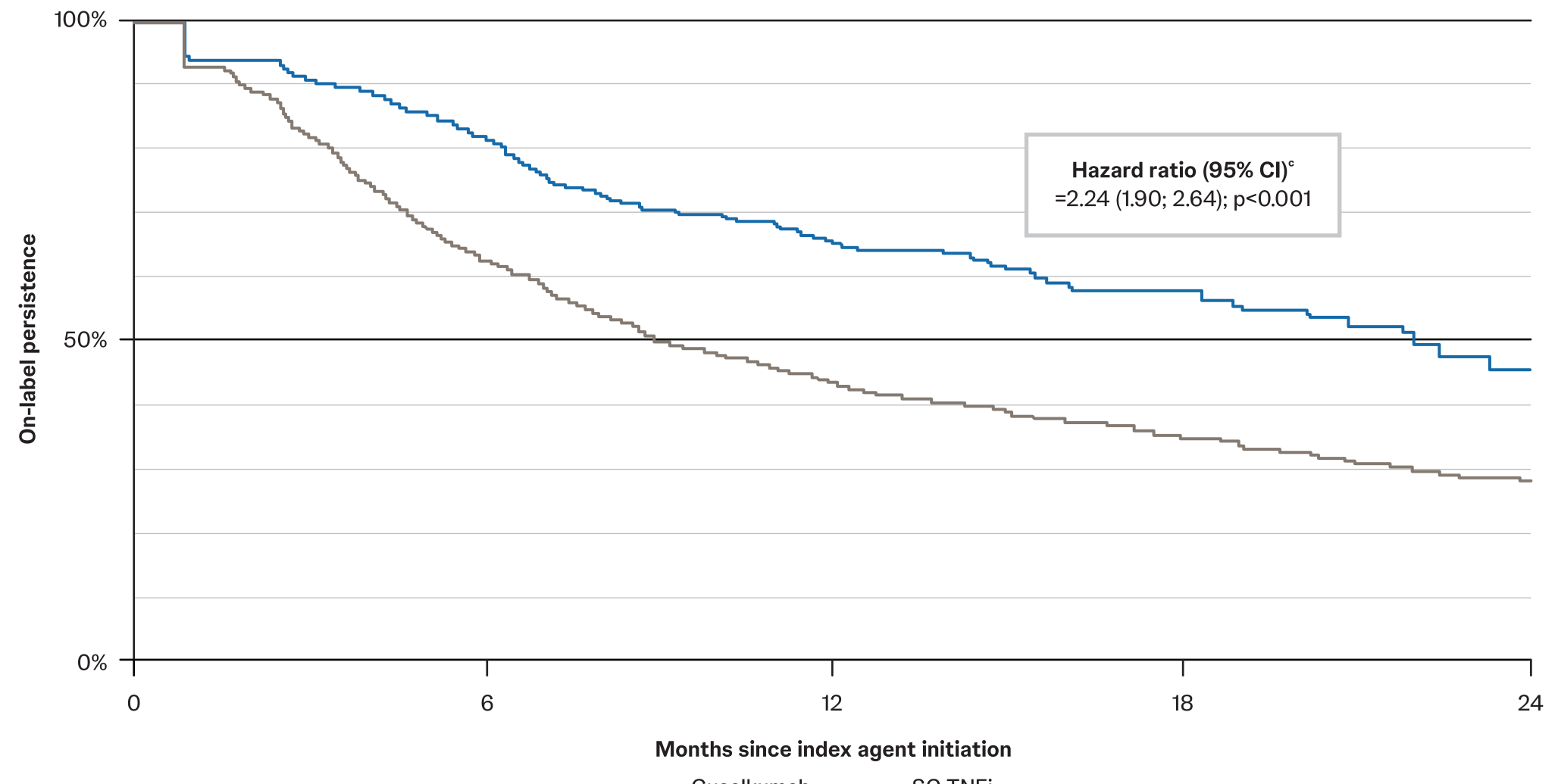
mographics			(N=2,490)
	Age at index date (years), Mean ± SD [median] Female Insurance type at index date	49.4 ± 11.2 [50.3] 60.3	49.5 ± 11.2 [51.0] 60.3
0 0	Preferred provider organization	76.2	76.2
	Health maintenance organization	12.4	13.1
	Other ^c	11.6	10.7
	Year of index date		
	2020	11.7	11.7
	2021	43.4	43.4
	2022	44.9	44.9
racteristics			
23	Months between latest observed PsA diagnosis and index date, Mean ± SD [median]	1.2 ± 1.4 [0.7]	1.2 ± 1.6 [0.7]
	Quan-CCI, Mean ± SD [median] Comorbidities	0.6 ± 1.3 [0.0]	0.6 ± 1.2 [0.0]
	Hyperlipidemia	34.7	34.7
	Osteoarthritis	30.3	30.3
- 414	Diabetes	15.3	15.5
	Peripheral vascular disease	1.4	2.3
	Psoriasis	86.3	86.3
	Smoking	11.6	11.2
ication Use ^d		47.0	14.0
	bDMARDs ^e	47.6	14.0
		52.4	86.0
F		41.2	12.6
	≥2	6.4	1.4
		22.4	48.3
	tsDMARDs ^g	21.1	23.4
	Corticosteroids	68.9	67.9



Pts in the GUS vs SC TNFi cohort were significantly (2.2x) more likely to remain persistent with on-label treatment through 24 months

- % pts with on-label persistence at 24 months: GUS (45.5%) vs SC TNFi (28.5%), despite a higher prevalence of biologic-experienced pts in the GUS cohort (47.6% vs 14.0% during 12-month baseline period)
- Median time to discontinuation: GUS (22.0 months) vs SC TNFi (9.2 months)
- In both sensitivity analyses, pts in the GUS cohort were significantly (~2x) more likely to remain persistent with on-label treatment at 24 months vs the SC TNFi cohort (1x: HR=1.90; fixed gap: HR=1.80; p<0.001 for both)





—— Guselkumab —— SC TNFi

^aDiscontinuation was defined as a gap in treatment of > twice the duration of days of supply for a claim (i.e., 2 x 56 = 112 days for guselkumab or 2 x 28 = 56 days for SC TNFi). ^bPatients with dose changes inconsistent with the FDA-approved dosing were censored as of the first dose change. ^cA weighted Cox proportional hazards model, further adjusted for baseline bDMARD and csDMARD use, was used to compare on-label persistence between cohorts. bDMARD=Biologic disease-modifying antirheumatic drug; CI=Confidence interval; csDMARD=Conventional synthetic disease-modifying antirheumatic drug; FDA=Food and drug administration; GUS=Guselkumab; *KM=Kaplan-Meier*; *SC TNFi=Subcutaneous tumor necrosis factor inhibitor*.



SC TNFi

GUS

112 days

56 days

l12 days

56 days

28 days 112 days

Key Takeaways

First real-world claims data analysis of on-label treatment persistence over 24 months in active PsA pts newly initiated on GUS vs initial SC TNFi



Pts in the GUS vs SC TNFi cohort were significantly (~2x) more likely to remain persistent on treatment through 24 months



Higher long-term on-label persistence may improve disease management outcomes, including functional status and quality of life, in pts with active PsA initiating GUS⁸

GUS was associated with significantly higher on-label persistence vs SC TNFi at each time point assessed (6/12/18/24 months)

Table 2. On-Label Persistence Through 24 months in Weighted GUS and SC TNFi Cohorts^a Primary Analysis (2x duration)

Cox proportional hazards model ^b	6 months	12 months	18 months	24 months			
Pts at risk, n (%)°							
GUS (N=804)	420 (52.2)	166 (20.6)	74 (9.2)	25 (3.1)			
SC TNFi (N=2,490)	1,068 (42.9)	479 (19.3)	234 (9.4)	114 (4.6)			
Hazard ratios (95% CI)	2.61 (2.10; 3.24)	2.34 (1.96; 2.79)	2.29 (1.94; 2.71)	2.24 (1.90; 2.64)			
Chi-square p-value	<0.001	<0.001	<0.001	<0.001			
KM persistence, % (95% CI)							
GUS	82.1 (76.3; 86.6)	65.9 (59.2; 71.8)	58.1 (49.5; 65.7)	45.5 (26.9; 62.1)			
SC TNFi	63.8 (60.1; 67.3)	43.8 (39.3; 48.2)	35.4 (30.0; 40.8)	28.5 (21.5; 35.9)			
Log-rank test p-value	<0.001	<0.001	<0.001	<0.001			

^aPropensity 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. ^bWeighted Cox proportional hazard models were used to compare risk of discontinuation between the GUS and SC TNFi cohorts. Models further adjusted for baseline use of bDMARDs and csDMARDs. ^cPts 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. bDMARD=Biologic disease-modifying antirheumatic drug; csDMARD=Conventional synthetic diseasemodifying antirheumatic drug; GUS=Guselkumab; KM=Kaplan-Meier; Pts=Patients; SC TNFi=Subcutaneous tumor necrosis factor inhibitor.

Strengths and Limitations

Strengths

- PsA pts were identified using a case finding algorithm validated in US claims data³
- After propensity score-weighting based on overlap weights, the GUS and SC TNFi cohorts were balanced for baseline demographic and disease characteristics, except for prior bDMARD or csDMARD use
- Given the claims database included a large sample of commercially insured PsA pts in the US, results are likely to be highly generalizable to that population

• Limitations

- Results may not be generalizable to non-commercially insured US pts or pts outside of the US
- Claims data do not ensure treatments are taken as prescribed
- Treatment effectiveness and reasons for discontinuation could not be assessed using claims data
- Days of supply in pharmacy claims data can be inaccurate due to coverage restrictions. Imputation is a valid approach commonly used for claims-based persistence analysis; however, it may occasionally lead to misclassifications

bDMARD=Biologic disease-modifying antirheumatic drug; csDMARD=Conventional synthetic disease-modifying antirheumatic drug; GUS=Guselkumab; PsA pts=Psoriatic arthritis patients; SC TNFi=Subcutaneous tumor necrosis factor inhibitor.