Intravenous Golimumab in Patients with Polyarticular Juvenile Idiopathic Arthritis and Juvenile Psoriatic Arthritis and Subcutaneous Ustekinumab in Patients with Juvenile Psoriatic Arthritis: Extrapolation of Data from Studies in Adults and Adjacent Pediatric Populations

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Abstract

Objectives To describe the extrapolation approaches used to support intravenous (IV) golimumab for polyarticular juvenile idiopathic arthritis (pJIA) and juvenile psoriatic arthritis (jPsA) and subcutaneous (SC) ustekinumab for jPsA.

Methods Pharmacokinetic, clinical response, and safety data from trials of IV golimumab and SC ustekinumab in polyarticular-course JIA (pc-JIA) (GO-VIVA) or pediatric psoriasis (PsO) (CADMUS and CADMUS Jr) and data from pivotal, phase 3 trials of these agents in adults with similar diseases were used to support extrapolation in pJIA and jPsA. In the phase 3 GO-VIVA trial, patients with pc-JIA aged 2 to < 18 years received IV golimumab 80 mg/m² at weeks 0, 4, then every 8 weeks (Q8W). In the phase 3, randomized, placebo-controlled CADMUS trial, patients with PsO aged ≥ 12 to < 18 years received ustekinumab at weeks 0, 4, then Q12W. In the phase 3 CADMUS Jr trial, patients with PsO aged ≥ 6 to < 12 years received ustekinumab at weeks 0, 4, then Q12W. The ustekinumab analyses used data only from patients who received the standard ustekinumab dosing regimen (≤ 60 kg: 0.75 mg/kg; > 60 to ≤ 100 kg: 45 mg; > 100 kg: 90 mg).

Results In the 127 patients with pc-JIA treated with IV golimumab (GO-VIVA), pharmacokinetic and exposure-response results were similar to those in adults with rheumatoid arthritis treated with IV golimumab. Additionally, pharmacokinetic and clinical response data from five patients with jPsA in GO-VIVA were comparable to those in adults with PsA treated with IV golimumab. No new safety signals were observed in GO-VIVA. Pharmacokinetic and clinical response data observed in the four pediatric patients with PsO and jPsA treated with ustekinumab in CADMUS and CADMUS Jr were similar to those in the 91 pediatric patients with PsO without jPsA in these trials and to those in adults with PsA treated with ustekinumab. Safety was extrapolated from CADMUS or CADMUS Jr; no new signals were observed.

Conclusions These three sets of analyses corroborate similar exposure and efficacy of IV golimumab in pediatric patients with pc-JIA or jPsA and SC ustekinumab in patients with jPsA to support extrapolation of established adult efficacy. The overall safety profiles of IV golimumab in pediatric patients with pc-JIA or jPsA and SC ustekinumab in pediatric patients with PsO with or without jPsA were consistent with the safety profiles of these agents in the context of their clinical programs and cumulative use. Based on these analyses, the US Food and Drug Administration approved IV golimumab for polyarticular JIA and active PsA in patients 2 years and older and SC ustekinumab for pediatric PsA in patients 6 years and older, highlighting how use of an extrapolation approach can help streamline drug development for pediatric patient populations in whom larger clinical trials are not feasible.

Clinical Trial Registration GO-VIVA (NCT02277444) was registered at clinicaltrials.gov on 29 October 2014; CADMUS (NCT01090427) was registered on 22 March 2010; and CADMUS Jr (NCT02698475) was registered on 3 March 2016.

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1 Introduction

Under the United States (US) Pediatric Research Equity Act (PREA), first passed in 2003, the US Food and Drug Administration (FDA) requires all applications for new treatments to contain an assessment of the product in the relevant pediatric population unless the applicant has obtained a waiver or deferral [1]. Pediatric drug studies pose unique challenges including small patient numbers and ethical restrictions for placebo-controlled trial designs. Extrapolation can address these challenges by utilizing the premise that the diseases and treatment responses are sufficiently similar in the populations being considered [2]. If this can be established, descriptive comparisons between observed pharmacokinetic (PK) and/or model-based exposure data from pediatric patients and data from adults can be conducted to support extrapolation. Exposure-response (ER) and clinical response data can also be compared, typically for a single-dose regimen in pediatric patients with data observed over a range of doses evaluated in adults.

Juvenile idiopathic arthritis (JIA) is an umbrella term encompassing several inflammatory arthritides of unknown etiology with symptom onset prior to 16 years of age and lasting for ≥ 6 weeks [3]. It has an estimated prevalence of 0.04–4 per 1000 children worldwide [4–7]. Available treatments for JIA include nonsteroidal anti-inflammatory drugs, systemic and intra-articular glucocorticoids, and conventional synthetic and biologic disease-modifying antirheumatic drugs (csDMARDs and bDMARDs, respectively) [4]. Registry data show that 45–52% of patients treated with ≥ two bDMARDs continue to have chronically uncontrolled JIA [8], highlighting the continued unmet need in these patients.

JIA categories were developed to identify clinically distinct groups for research purposes, and do not reflect emerging understanding of the disease [9, 10]; validation of an updated JIA classification system is underway [11]. The current International League of Associations for Rheumatology JIA classification system identifies seven JIA categories: systemic JIA, polyarticular JIA (pJIA) rheumatoid factor (RF)-negative, pJIA RF-positive, oligoarticular (persistent: ≤ four joints; extended: > four joints after the first 6 months of disease), juvenile psoriatic arthritis (jPsA), enthesitis-related arthritis, and undifferentiated arthritis [3]. Polyarticular-course JIA (p-cJIA) affects ≥ five joints cumulatively throughout the course of the disease and can occur within each JIA category. Many of the underlying pathological immune mechanisms that have been identified in rheumatoid arthritis (RA) have also been identified in pJIA (Fig. 1) [5, 6, 8, 10, 12]. Thus, PREA requires that bDMARDs approved for RA also be studied in JIA [8]. Given the relative rarity of JIA and the jPsA JIA subtype (5% of patients with JIA) [13, 14], the FDA previously granted waivers for studies of jPsA following approvals of treatments for adult psoriatic arthritis (PsA).

In the context of the aforementioned challenges in pediatric drug development and substantial unmet need in JIA, the use of extrapolation approaches for well-established classes of drugs has been considered, including utilization of open-label PK studies to demonstrate similarity to adult exposures and extrapolation of efficacy and safety observed in adults [15–17]. In addition, after a 2019 FDA/Center for Excellence in Regulatory Science and Innovation workshop entitled “Accelerating Drug Development for Polyarticular Juvenile Idiopathic Arthritis (pJIA),” the FDA advised that PREA requirements for pJIA can be addressed with PK studies to support the extrapolation of efficacy data from adults with RA [15, 16]. Subsequently, the FDA advised that, as the underlying mechanisms, comorbidities, and clinical presentations for PsA may overlap with RA and psoriasis (PsO), data from studies conducted in p-cJIA and RA and in pediatric and adult PsO and adult PsA populations can be used to support extrapolation for jPsA without a PK study [18]. The assessment of safety could also be supplemented by results from other relevant pediatric and adult populations.

Here, we describe three sets of analyses that support pediatric extrapolation for the use of two bDMARDs, approved
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in adult rheumatologic and psoriatic indications (i.e., RA, PsA, and PsO), in similar pediatric indications (i.e., pJIA and jPsA). First, intravenous (IV) golimumab, a tumor necrosis factor-α inhibitor (TNFi) FDA approved for adults with active RA, PsA, or ankylosing spondylitis, was studied for use in pc-JIA. Per FDA request, an initial set of PK exposure and ER analyses was conducted using observed data from an open-label, phase 3 study in pediatric patients with active pc-JIA [19] to determine if extrapolation of data from pivotal, phase 3 studies of IV golimumab in adults with RA was appropriate [20, 21]. As several patients in this study also had jPsA, a second set of analyses assessed data from the jPsA and pc-JIA populations as well as data from adults with PsA [22]. These analyses were the basis of FDA approval of IV golimumab for pJIA and active PsA in patients 2 years of age and older. As a result of these analyses, an extrapolation approach was applied to support the use of ustekinumab for jPsA. Ustekinumab is an interleukin (IL)-12/IL-23 inhibitor that is FDA approved for adults with active PsA, PsO, Crohn’s disease, or ulcerative colitis and for pediatric patients ≥6 years of age with moderate-to-severe PsO. This third set of analyses evaluated data from studies of ustekinumab in pediatric patients with PsO, including a small number of patients with PsO and jPsA [23, 24], and in adults with PsA [25]. These analyses were the basis of FDA approval of SC ustekinumab for pediatric PsA in patients 6 years and older.

2 Methods

2.1 Clinical Trials

As detailed in Online Supplemental Material (OSM) Table S1 and OSM Table S2, data employed in these analyses were derived from several clinical trials. Studies of IV golimumab were: (1) GO-VIVA (NCT02277444; phase 3, open-label, PK, efficacy, and safety trial), which evaluated IV golimumab (80 mg/m² at weeks 0, 4, then every 8 weeks (Q8W)) in 127 patients aged 2 to <18 years with pc-JIA, including patients with jPsA [19]; (2) GO-FURTHER (NCT00973479; phase 3, randomized, placebo-controlled trial), which evaluated IV golimumab (2 mg/kg at weeks 0, 4, then Q8W) in 592 adults with RA [20]; (3) GO-LIVE (NCT00361335; phase 3, randomized, placebo-controlled trial), which evaluated IV golimumab (2 or 4 mg/kg Q12W) in 643 adults with RA [21]; and (4) GO-VIBRANT (NCT02181673; phase 3, randomized, placebo-controlled trial), which evaluated IV golimumab (2 or 4 mg/kg Q12W) in 480 adults with PsA [22]. The analyses for IV golimumab utilized data only from patients who received concomitant methotrexate and had available PK samples at the required timepoints.

Ustekinumab trials were: (1) CADMUS (NCT01090427; phase 3, randomized, placebo-controlled), which evaluated ustekinumab (standard or half-standard dosing regimens subcutaneously at weeks 0, 4, then Q12W) in 110 patients ≥12 to <18 years of age with PsO, including patients with jPsA [23]; (2) CADMUS Jr (NCT02698475; phase 3, open-label, PK, efficacy, and safety trial), which evaluated IV golimumab (2 mg/kg at weeks 0, 4, then Q8W) in 241 adults with PsA; and (3) PSTELLAR (NCT01550744; phase 3, randomized, placebo-controlled trials that evaluated ustekinumab (45 or 90 mg subcutaneously at weeks 0, 4, then Q12W for both) in 615 and 312 adults, respectively, with PsA; and (5) PSTELLAR (NCT01550744; phase 3, randomized,
controlled trial), which evaluated ustekinumab (45 or 90 mg subcutaneously at weeks 0, 4, 16, then Q12W up to Q24W) in 478 adults with PsO [27]. The analyses for ustekinumab used data only from pediatric patients who received the standard ustekinumab dosing regimen (≤ 60 kg: 0.75 mg/kg; > 60 to ≤ 100 kg: 45 mg; > 100 kg: 90 mg), which demonstrated drug exposure in pediatric patients was comparable to that in adult patients who received ustekinumab 45 mg [23, 24].

All trials included in these analyses were conducted and reported in accordance with the ethical principles originating in the Declaration of Helsinki [28], International Council for Harmonisation (ICH) Good Clinical Practice guidelines [29], and applicable regulatory requirements and in compliance with the respective protocols. Data from relevant dose regimens that were analyzed with the same bioanalytical assays were included to assess the similarity of PK exposure and/or clinical response analyses among the different populations.

2.2 IV Golimumab for Polyarticular JIA

To support IV golimumab for pJIA, PK and ER data observed in 127 children and adolescents with pc-JIA (including all categories of JIA) from the overall GO-VIVA population were descriptively compared with data from adults with RA (GO-FURTHER, GO-LIVE) (Fig. 2a).

2.2.1 Exposure Matching for IV Golimumab for Polyarticular JIA

Median observed steady-state trough golimumab concentrations ($C_{\text{trough,ss}}$) and model-predicted, steady-state area under the concentration-time curves ($AUC_{\text{ss}}$) over the 8-week golimumab dosing interval at week 28 (overall and by age category) in patients with pc-JIA (GO-VIVA) were compared descriptively with golimumab $C_{\text{trough,ss}}$ and $AUC_{\text{ss}}$ assessed at weeks 20 and 36 in adults with RA (GO-FURTHER). Week 20 and week 36 data from GO-FURTHER were used in the absence of a week 28 visit. Model-predicted $AUC_{\text{ss}}$...
was generated from GO-VIVA and GO-FURTHER data using similar population PK models established for pJIA and RA, respectively (see OSM Methods).

### 2.2.2 Exposure-Response Analyses for IV Golimumab for Polyarticular JIA

The PK exposure metrics underpinning the ER analyses were the observed golimumab $C_{\text{trough,ss}}$, calculated average golimumab concentrations at steady state ($C_{\text{avg,ss}}$), and cumulative AUC. Observed golimumab $C_{\text{trough,ss}}$ values below the lower limit of quantitation were set to 0. $C_{\text{avg,ss}}$ was calculated as dose/clearance/tau, where tau was the dosing interval. Cumulative AUC was calculated using patientspecific PK parameters and their corresponding dosing and sampling timepoints per the final population PK models. The efficacy endpoints in these ER analyses were the proportions of patients achieving ≥ 20% improvement in the American College of Rheumatology response criteria (ACR20) [30] in GO-FURTHER and GO-LIVE or JIA ACR30 response (≥ 30% improvement from baseline in ≥ three components, without worsening of ≥ 30% in > one of the remaining JIA core measures) in GO-VIVA [31]. While numerically different, the ACR20 and JIA ACR30 responses are similar measures of efficacy in the adult RA and pJIA populations, respectively. The core components of the ACR and JIA ACR scores generally align, noting that the pediatric equivalent of the adult patient’s global assessment of disease activity is the parent/patient assessment of overall well-being [30–32]. Patients missing data for all components of the composite endpoint were considered nonresponders (i.e., nonresponder imputation (NRI)). Last observation carried forward was used to impute the composite endpoint when ≥ one of the component values was missing.

Week 20 PK data from GO-VIVA and GO-FURTHER and week 24 PK data from GO-LIVE were descriptively compared as the closest steady-state trough concentrations representing approximately 6 months of exposure because it was hypothesized that efficacy would be similar at these timepoints. Likewise, week 52 efficacy data from GO-VIVA and GO-FURTHER and week 48 efficacy data from GO-LIVE were compared in the same analyses. The proportions of adult RA patients in GO-FURTHER and GO-LIVE achieving ACR20 at weeks 20 and 24, respectively, versus corresponding golimumab $C_{\text{trough,ss}}$ quartiles were compared with the proportion of pc-JIA patients in GO-VIVA achieving JIA ACR30 at week 20 versus $C_{\text{trough,ss}}$ quartiles at week 20. Percent changes from baseline in the shared components of the ACR and JIA ACR composite endpoints (i.e., tender joints, swollen joints, patient assessment of disease activity, physician assessment of disease activity, and patient assessment of physical function) [30, 31] from GO-VIVA (at week 20), GO-FURTHER (at week 20), and GO-LIVE (at week 24) versus $C_{\text{trough,ss}}$ quartiles at these timepoints were also compared.

### 2.2.3 Safety Analyses for IV Golimumab for Polyarticular JIA

As previously reported, GO-VIVA adverse events (AEs) were monitored through week 52 [19]. Safety data from the overall population in GO-VIVA were evaluated in the context of the established, well-characterized safety profile of IV golimumab in adults with RA or PsA, the extensive safety data that have been collected in clinical trials to support multiple indications [33, 34], and the SC golimumab safety data that have accrued since the first approval in 2009. In company-sponsored interventional trials, > 13,000 individuals have been exposed to IV golimumab, including 1213 adults with RA in GO-FURTHER and GO-LIVE [20, 21], 460 adults with PsA in GO-VIBRANT [22], and 204 adults with ankylosing spondylitis (GO-ALIVE) [35]. The cumulative clinical trial exposure also includes 173 patients with pc-JIA aged 2–17 years who received SC golimumab in the GO-KIDS placebo-controlled, randomized-withdrawal trial [36]. Owing to interstudy variations and differences in sample size and duration of follow-up, safety comparisons focused on the types and patterns of AEs rather than on direct comparisons of AE frequency. An estimated 918,867 patients have been exposed to SC golimumab worldwide cumulatively, and 121,713 patients have been exposed to IV golimumab since approval.

### 2.3 IV Golimumab for JPsA

The GO-VIVA pc-JIA population included five patients with JPsA [19]. To support extrapolation of IV golimumab for JPsA, PK and clinical response data observed in the five patients with JPsA in GO-VIVA were descriptively compared with data from adults with PsA (GO-VIBRANT) (Fig. 2a).

#### 2.3.1 Exposure Matching for IV Golimumab for JPsA

Golimumab $C_{\text{trough,ss}}$ at weeks 28 and 52 in patients with JPsA in GO-VIVA were descriptively compared with golimumab $C_{\text{trough,ss}}$ at weeks 28 and 52 in the overall GO-VIVA population and in the overall population excluding patients with JPsA, and golimumab $C_{\text{trough,ss}}$ at weeks 36 and 52 in adults with PsA in GO-VIBRANT.

#### 2.3.2 Response Analyses for IV Golimumab for JPsA

For these clinical response analyses, the proportions of patients with JPsA from GO-VIVA who achieved JIA ACR30/50/70/90 (≥ 30, ≥ 50, ≥ 70, and ≥ 90%
2.3.3 Safety Analyses for IV Golimumab for jPsA

Safety was assessed among the five patients with jPsA from GO-VIVA and evaluated in the context of the safety profile of the overall GO-VIVA population and the established IV golimumab and SC golimumab safety profile as described previously (see Sect. 2.2.3).

2.4 Ustekinumab for jPsA

In the CADMUS and CADMUS Jr trials of ustekinumab in adolescents and children, respectively, with PsO, seven patients also had jPsA [23, 24]. Given the availability of these data, and the successful prior extrapolation of adult PsA studies for IV golimumab for jPsA, extrapolation was then planned for ustekinumab in jPsA. To support this extrapolation, data from patients with pediatric PsO alone and from those with PsO and jPsA in CADMUS and CADMUS Jr were descriptively compared with data from adults with PsO (PSTELLAR) or PsA (PSUMMIT-1) (Fig. 2b).

2.4.1 Exposure Matching for Ustekinumab for jPsA

Ustekinumab $C_{\text{trough,ss}}$ through week 52 in adults with PsO from PSTELLAR were descriptively compared with ustekinumab $C_{\text{trough,ss}}$ through week 52 in adults with PsA from PSUMMIT-1 among patients who received the 45 mg dose, which is consistent with the standard approved adult dosage for PsA (45 mg at weeks 0, 4, then Q12W). In CADMUS and CADMUS Jr, four of the seven patients with jPsA received the standard dosage. The ustekinumab $C_{\text{trough,ss}}$ through week 52 in these four patients were compared with the overall population from CADMUS and CADMUS Jr who received the standard adult dosage, with the overall population excluding patients with jPsA, and with adults with PsA from PSUMMIT-1 who received the standard dosage.

2.4.2 Response Analyses for Ustekinumab for jPsA

Response rates for achievement of $\geq 75$, $\geq 90$, and 100% improvement in Psoriasis Area and Severity Index (PASI) score (PASI75, PASI90, and PASI100, respectively) [37] in patients with jPsA from CADMUS and CADMUS Jr who received the standard adult ustekinumab dosage were descriptively compared with those of adults with PsA from PSUMMIT-1 who received the standard dosage. PASI data from CADMUS and CADMUS Jr were imputed using last observation carried forward for any missing components, and NRI if all components were missing.

2.4.3 Safety Analyses for Ustekinumab for jPsA

Safety for ustekinumab for jPsA was extrapolated from the established safety profile of ustekinumab in the 154 pediatric patients with PsO in CADMUS and CADMUS Jr, including seven patients who had PsO and jPsA [23, 24]. As previously reported, AEs were monitored through weeks 60 and 56, respectively, in these trials. Extensive safety data have also been collected in ustekinumab clinical trials for multiple indications in adults and pediatric patients [38]. In company-sponsored interventional studies, 12,228 participants have been exposed to ustekinumab, including 5750 adult and pediatric participants with PsO and 1045 adult participants with PsA. Ustekinumab clinical trial exposure also includes 44 pediatric patients with Crohn’s disease aged, 2 to <18 years, who received ustekinumab in a randomized, dose-ranging study [39]. As with IV golimumab, ustekinumab safety comparisons focused on the types and patterns of AEs rather than on direct comparisons of AE frequency due to differences in study populations and designs.
3 Results

3.1 IV Golimumab for Polyarticular JIA

3.1.1 PK Matching for IV Golimumab for Polyarticular JIA

The PK analysis set for patients with pc-JIA included 127 patients who received ≥ one golimumab infusion in GO-VIVA and had sufficient PK samples for analysis. Golimumab PK exposures ($C_{\text{trough,ss}}$ and $AUC_{\text{ss}}$) at week 28 in these pediatric patients were similar across pediatric age categories (Fig. 3). Median $C_{\text{trough,ss}}$ at week 28 in pediatric patients with pc-JIA (range 0.31–0.48 µg/mL) were similar to those at weeks 20 (0.21 µg/mL) and 36 (0.31 µg/mL) in adults with RA who received ≥ one IV golimumab infusion in GO-FURTHER (Fig. 3a). Model-predicted $AUC_{\text{ss}}$ at week 28 in patients with pc-JIA (range 387–424 µg·day/mL) was 50–64% higher than $AUC_{\text{ss}}$ in adults with RA (248 µg·day/mL) (Fig. 3b).

3.1.2 Exposure-Response for IV Golimumab for Polyarticular JIA

Week 20 JIA ACR30 response rates among pc-JIA patients in GO-VIVA were greater than ACR20 response rates at week 20 in GO-FURTHER (adults with RA) and at week 24 in GO-LIVE (adults with RA), across serum golimumab $C_{\text{trough,ss}}$ in adults (Fig. 4a). In addition, median improvements in JIA ACR and ACR component scores (i.e., tender joints, swollen joints, patient assessment of disease activity, physician assessment of disease activity, and patient assessment of physical function) at these timepoints were similar or greater in patients with pc-JIA versus adults with RA, regardless of $C_{\text{trough,ss}}$ in adults (Fig. 4b–f). Results were consistent when efficacy was evaluated by $C_{\text{avg,ss}}$ and cumulative $AUC$ (data not shown), although PK exposures were...
slightly higher in pediatric patients with pc-JIA versus adults with RA.

3.1.3 Safety for IV Golimumab for Polyarticular JIA

Detailed safety results through 1 year of IV golimumab treatment in GO-VIVA have been previously reported [19]. Briefly, through 1 year, 85% of pediatric patients with pc-JIA experienced ≥ one AE, with infections being the most common type; 7% experienced ≥ one serious AE, 65% experienced ≥ one infection, and 6% experienced ≥ one serious infection, including one patient with a serious opportunistic infection. The most commonly reported AEs were upper respiratory tract infection (21%) and nasopharyngitis (18%). Serious AEs were disseminated herpes zoster, infective exacerbation of bronchiectasis, sepsis, varicella, mycosis fungoides, suicidal ideation, cellulitis, pneumonia, streptococcal pneumonia, and pleural effusion (streptococcal pneumonia and pleural effusion were reported in the same patient). Mild, new-onset, anterior uveitis in both eyes was reported in one patient but did not require treatment. No cases of active tuberculosis, demyelination events, or anaphylactic or serum sickness reactions were reported. Systemic lupus erythematosus was reported in one patient, but was considered nonserious. No deaths were reported through week 52, but one death due to septic shock (likely due to constipation leading to bacterial translocation through the gut wall) was reported at week 78 (last IV golimumab dose at week 76) in this ongoing study.

Comparison of safety data from GO-VIVA to the established safety profile for golimumab from studies conducted in adults did not identify any new AEs. In addition, the types of AEs observed in GO-VIVA through week 52 were comparable to those observed with IV golimumab in adults with rheumatic disease [20–22, 35] and with SC golimumab in patients with pc-JIA [36]. Furthermore, no new safety concerns were identified relative to the safety profile observed across the TNFi class in either adult or pediatric indications [40–43].

3.2 IV Golimumab for jPsA

3.2.1 PK Matching for IV Golimumab for jPsA

Among the five patients with pc-JIA in GO-VIVA with jPsA, four had evaluable PK samples at weeks 28 and 52 and were included in the PK analysis set. The observed serum golimumab $C_{\text{trough,ss}}$ values were similar among patients with jPsA, pc-JIA (excluding jPsA), and pc-JIA (overall) at weeks 28 and 52 (Fig. 5a). In addition, golimumab $C_{\text{trough,ss}}$ at week 28 in patients with jPsA was similar to $C_{\text{trough,ss}}$ at week 36 in adults with PsA from GO-VIBRANT with overlapping steady-state trough serum golimumab concentration ranges (Fig. 5b). Similar results were observed at week 52.

3.2.2 Response Analyses for IV Golimumab for jPsA

The proportions of patients with jPsA achieving JIA ACR30 responses through week 52 were consistent with the proportions of adults with PsA achieving ACR20 through week 52 (Fig. 6a). Efficacy as assessed by JIA ACR50/70/90 and ACR50/70/90 response rates was consistently seen in both pediatric and adult patients, respectively, through week 52 (Fig. 6b–d).

3.2.3 Safety for IV Golimumab for jPsA

The AEs observed in the five patients with jPsA who received IV golimumab in GO-VIVA were similar to those in the overall GO-VIVA population and to those in adults with PsA in GO-VIBRANT. Seven AEs occurred in four patients with jPsA through week 52 (pyoderma, gastroenteritis, nausea, upper respiratory tract infection, increased blood pressure, furuncle, and double-stranded DNA antibody positive). No serious AEs were reported. All AEs were singular events, and none resulted in discontinuation of golimumab. Furthermore, the overall safety profile of IV golimumab in patients with pc-JIA in GO-VIVA, which included patients with jPsA, was consistent with the safety profile of SC golimumab in patients with pc-JIA in GO-KIDS, which also included patients with jPsA [36].

3.3 Ustekinumab for jPsA

3.3.1 PK Matching for Ustekinumab for jPsA

The PK analysis set for ustekinumab in jPsA included four pediatric patients with PsO and jPsA and 91 pediatric patients with PsO without jPsA who received standard ustekinumab dosing in CADMUS and CADMUS Jr. Serum ustekinumab concentrations over time were comparable in adults with PsO (PSTELLAR) and adults with PsA (PSUMMIT-1) who received similar ustekinumab dosages, suggesting that extrapolating data between PsO and PsA is feasible (Fig. 7a). Serum ustekinumab concentrations over time were also comparable between pediatric patients with PsO with or without jPsA (CADMUS and CADMUS Jr) who received similar ustekinumab dosages, confirming that extrapolation between these pediatric patient populations was appropriate (Fig. 7b). In addition, PK exposures over time were comparable between pediatric patients with PsO with and without jPsA (CADMUS and CADMUS Jr) and adults with PsA (PSUMMIT-1), supporting the conclusion that extrapolation between these patient groups is appropriate.
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**Fig. 4** Efficacy responses by golimumab steady-state trough concentration quartiles in adults with RA from GO-FURTHER (week 20) and GO-LIVE (week 24) and pediatric patients with pc-JIA from GO-VIVA (week 20) a ACR20 and JIA ACR30 response rates, and percent change from baseline in b tender joint count, c swollen joint count, d patient assessment of disease activity, e physician assessment of disease activity, and f patient assessment of physical function. Circle within box = median; lower edge of box = first quartile; upper edge of box = third quartile; ends of whiskers represent ± 1.5 × IQR. ACR20 ≥ 20% improvement from baseline in tender and swollen joint counts and ≥ 20% improvement in three of the five other core set measures. JIA ACR30 = ≥ 30% improvement from baseline in three components, without worsening of ≥ 30% in > one of the remaining JIA core measures. Patients missing data for all components of the composite endpoint were considered nonresponders. Last observation carried forward was used to impute the composite endpoint when ≥ one of the component values was missing. ACR American College of Rheumatology, C<sub>trough,ss</sub> steady-state trough concentration, JIA juvenile idiopathic arthritis, pc-JIA polyarticular-course juvenile idiopathic arthritis, Q quartile, RA rheumatoid arthritis
3.3.2 Response Analyses for Ustekinumab for jPsA

PASI75 response rates in four pediatric patients with PsO and jPsA who received ustekinumab standard dosing were comparable with those in adults with PsA (PSUMMIT-1) (Fig. 8). PASI90 and PASI100 response rates were generally similar between the patient populations (OSM Fig. S1a–b).

PASI and ACR response rates were similar between adults with PsA from PSUMMIT-1 and PSUMMIT-2 who were diagnosed with PsA at < 18 years of age (pediatric onset of PsA) versus > 18 years of age (adult onset of PsA) when adjusted by PsO (OSM Fig. S2a) or PsA (OSM Fig. S2b) disease duration. Although wider confidence intervals were observed in the pediatric-onset group due to a small sample size, odds ratios for PASI and ACR responses for ustekinumab versus placebo were similar between the pediatric and adult-onset groups. When analyzed as a spidergram, adult and pediatric responses mirrored each other, with wider confidence intervals again observed in the pediatric-onset group due to the small sample size (OSM Fig. S2c).

3.3.3 Safety for Ustekinumab for jPsA

Through 1 year, no new safety signals were observed in the 154 pediatric patients with PsO treated with ustekinumab in CADMUS and CADMUS Jr [23, 24]. Nasopharyngitis, pharyngitis, and upper respiratory tract infection were among the most commonly reported AEs in both studies. Tonsillitis, gastroenteritis, and otitis media occurred more frequently in CADMUS Jr; however, these events are known to be more commonly reported in younger patients [44–46] and do not appear to represent newly identified safety concerns in this younger population. The incidence and types of AEs in the CADMUS Jr long-term extension were also consistent with...
those in the main study. In addition, the overall safety profile of ustekinumab in CADMUS and CADMUS Jr was consistent with the safety profile of ustekinumab in pivotal clinical trials of adult PsO and adult PsA [38].

In addition, no new safety signals were observed in the seven patients with PsO and jPsA who received standard or half-standard ustekinumab dosing in CADMUS or CADMUS Jr, which was not surprising given the small number of patients with PsO and jPsA. Six of these seven patients reported ≥ 1 AE, and none reported serious AEs. The similarity in safety profiles between pediatric PsO and jPsA is supported by the comparable safety profiles observed in phase 3 clinical trials in adults with PsO [47, 48] and adults with PsA who received similar ustekinumab dose regimens [25, 26]. In these studies, there was no clear ustekinumab dose-response relationship in AEs or difference in safety profiles between patients who did or did not receive concomitant methotrexate.

While postmarketing safety data for ustekinumab in pediatric patients with PsO are limited, no safety concerns have been identified to date in patients aged ≥ 6 to < 18 years who received ustekinumab, and the safety profile is similar to that observed for ustekinumab in the adult population.

4 Discussion

With the advent of PREA, in the absence of a waiver, products for which an application is being made to the FDA require pediatric studies. Though this requirement benefits pediatric
Fig. 7 Observed serum ustekinumab $C_{\text{trough, ss}}$ through week 52 in a adults with PsO (PSTELLAR) or PsA (PSUMMIT-1) and b adults with PsA (PSUMMIT-1) and pediatric patients with PsO with or without jPsA (CADMUS and CADMUS Jr). Horizontal line within box = median; diamond = mean; lower edge of box = first quartile; upper edge of box = third quartile; ends of whiskers represent ±1.5 $\times$ IQR. Adult PsO and PsA populations included only adults who received the ustekinumab 45 mg dose regimen. Pediatric PsO with or without jPsA populations included only pediatric patients assigned to the standard dose treatment group and who received the standard dose regimen at 0.75 mg/kg or 45 mg. $C_{\text{trough, ss}}$ steady-state trough concentration, jPsA juvenile psoriatic arthritis, PsA psoriatic arthritis, PsO psoriasis, w/o without
Extrapolation for IV Golimumab and SC Ustekinumab in Pediatric Rheumatology

patients, such studies are fraught with difficulties. Extrapolation provides an elegant solution to address practicalities (such as small sample size) and ethical challenges (including pediatric patients with a progressive disease receiving a placebo) that are barriers to conducting pediatric clinical trials. Indeed, these were the barriers faced when considering IV golimumab for pJIA and jPsA and SC ustekinumab for jPsA.

With a completed IV golimumab PK, efficacy, and safety study in patients with pc-JIA, including jPsA, data in pediatric patients and knowledge about PK exposure, efficacy, and safety were determined by the FDA to be adequate to support extrapolation. IV golimumab PK, efficacy, and safety data in adults with RA or PsA were also determined by the FDA to be suitable for extrapolation to patients with pc-JIA, including pJIA and active PsA. Similarly, with two completed ustekinumab pediatric PsO trials, including patients with jPsA, as well as one completed pediatric Crohn’s disease trial, data in pediatric patients and knowledge about PK exposure, efficacy, and safety were determined by the FDA to be adequate to support extrapolation without enrolling any children into a pediatric PsA-specific clinical trial. Ustekinumab data in adults with PsO or PsA were also determined by the FDA to be adequate for extrapolation to patients with pediatric PsA.

The extrapolation of the robust efficacy demonstrated by the well-controlled clinical trials in adults is supported by known clinical similarities between (1) pJIA and adult RA; (2) jPsA and adult PsA; and (3) pediatric and adult PsO (Fig. 1). The results of the analyses conducted, including overlapping golimumab and ustekinumab PK exposures and consistent efficacy between pediatric and adult patients, further support extrapolation. Although model-predicted AUC$_{ss}$ in patients with pc-JIA was substantially higher than AUC$_{ss}$ in adults with RA, the differences were within the wide range of expected variability that has been observed with monoclonal antibodies [49–51]. In the context of this moderate to high variability, it can be concluded that the PK data are similar enough to support extrapolation.

Similarly, the safety profiles of IV golimumab and SC ustekinumab in pediatric patients were consistent with the safety profiles of these drugs in adult patients with RA, PsO, or PsA. There were no new safety signals in the clinical studies of pediatric patients with pc-JIA [19] or PsO with or without jPsA [23, 24] treated with IV golimumab or SC ustekinumab, respectively. In addition, as expected due to the small patient numbers, no new signals were observed in pediatric patients with jPsA treated with either drug.

The number of pediatric patients was small, particularly for the jPsA analyses for both compounds, making true differences between populations difficult to discern due to the inherent uncertainty. However, as noted in the recently
published (April 2022) ICH guidelines for pediatric extrapolation [2], clinical judgment should be utilized to establish the tolerable level of uncertainty associated with the data supporting extrapolation to a target pediatric population. The overlapping serum steady-state drug concentrations over time corroborated similarity of PK exposure and consistent response patterns for pediatric and adult populations. Notably, for jPsA, these analyses of PK data from adjacent, closely related diseases helped to address uncertainty and to support extrapolation of safety and efficacy from the results of pediatric PsO, adult PsO, and adult PsA pivotal trials without the need for trials dedicated to jPsA, which are difficult to conduct due to the rarity of this category of JIA [13, 14]. In addition, the totality of comparable results between these populations supports a favorable benefit-risk profile for IV golimumab and SC ustekinumab in pediatric patients with pJIA and/or jPsA.

5 Conclusions

Altogether, overlapping clinical presentations across disease states, comparable PK exposure and clinical response, and consistent safety profiles between pediatric patients and adults support extrapolation of the robust efficacy of IV golimumab and SC ustekinumab demonstrated in well-controlled clinical trials in adults to pediatric patients. Based on these analyses, the FDA approved IV golimumab for pJIA and active PsA in patients 2 years and older in September 2020, and SC ustekinumab for pediatric PsA in patients 6 years and older in July 2022. These results highlight how use of exposure matching and clinical response analyses can help streamline drug development for pediatric patient populations in whom larger clinical trials are not feasible.

Supplementary Information

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Declarations

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Conflict of interest

Jocelyn H. Leu, Michael Clark, Karen Bensley, Kathleen G. Lomax, Katherine Berezny, and Zhenhua Xu are employees of Janssen Research & Development, LLC, a subsidiary of Johnson & Johnson, and may own stock in Johnson & Johnson. Honghui Zhou was an employee of Janssen Research & Development, LLC, a subsidiary of Johnson & Johnson, at the time the work was conducted and may own stock in Johnson & Johnson; Dr. Zhou is currently an employee at Elevare Therapeutics, Inc, Salt Lake City, UT, USA. Natalie J. Shiff is an employee of Janssen Scientific Affairs, LLC, a subsidiary of Johnson & Johnson; received salary support from the Childhood Arthritis and Rheumatology Research Alliance within the past 3 years; and owns or has owned stock in AbbVie, Gilead, Iovance, Johnson & Johnson, Novo-Nordisk, and Pfizer within the past 3 years. Robert M. Nelson is an employee of Johnson & Johnson and owns stock in Johnson & Johnson.

Ethics

All trials included in these analyses were conducted and reported in accordance with the ethical principles originating in the Declaration of Helsinki and in accordance with International Council for Harmonisation Good Clinical Practice guidelines, applicable regulatory requirements, and in compliance with the respective protocols.

Consent to participate

All participants or legal guardians of participants in the clinical trials included in these analyses provided written informed consent.

Consent to publication

Not applicable.

Availability of data and material (data transparency)

The data-sharing policy of Janssen Pharmaceutical Companies of Johnson & Johnson is available at https://www.janssen.com/clinical-trials/transparency. As noted on this site, requests for access to the trial data can be submitted through Yale Open Data Access (YODA) Project site at http://yoda.yale.edu.

Code availability

Not applicable.

Author contributions

Conceptualization: JHL, NJS, KBen, KGL, KBer, RMN, HZ, ZX. Methodology: JHL, MC, KBen, KGL, KBer, RMN, HZ, ZX. Data collection and/or analysis: JHL, NJS, MC, KBen, KGL, KBer, HZ, ZX. Data interpretation: JHL, NJS, MC, KBen, KGL, KBer, RMN, HZ, ZX. Manuscript preparation, review, and editing: All authors. All authors approved the final version of the manuscript for publication.

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