

**Absence of Association Between Abatacept Exposure and Initial Infection in Patients With Juvenile Idiopathic Arthritis**

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**Running head:** Abatacept Exposure and Infection

## ABSTRACT

**Objective.** To assess the relationship between infection risk and abatacept exposure levels in patients with polyarticular-course juvenile idiopathic arthritis (pJIA) following treatment with subcutaneous and intravenous abatacept.

**Methods.** Data from two published studies (NCT01844518, NCT00095173) of abatacept treatment in pediatric patients were analyzed. One study treated patients aged 2–17 years with subcutaneous abatacept and the other treated patients aged 6–17 years with intravenous abatacept. Association between serum abatacept exposure measures and infection was evaluated using Kaplan–Meier plots of probability of first infection versus time on treatment by abatacept exposure quartiles and log-rank tests. Number of infections by abatacept exposure quartiles was investigated.

**Results.** Overall, 343 patients were included in this analysis: 219 patients received subcutaneous abatacept and 124 patients received intravenous abatacept. Overall, 237/343 (69.1%) patients had  $\geq 1$  infection over 24 months. No significant difference in time to first infection across four quartiles of abatacept exposure levels was observed in the pooled ( $p = 0.4458$ ), subcutaneous (2–5 years  $p = 0.9305$ ; 6–17 years  $p = 0.4787$ ), or intravenous ( $p = 0.4999$ ) analyses. Concomitant use of methotrexate and glucocorticoids (at baseline and throughout) with abatacept did not increase infection risk across the abatacept exposure quartiles. There was no evidence of association between number of infections and abatacept exposure quartiles. No opportunistic infections related to abatacept were reported.

**Conclusion.** In patients aged 2–17 years with pJIA, no evidence of association between higher levels of exposure to intravenous or subcutaneous abatacept and incidence of infection was observed.

Juvenile idiopathic arthritis (JIA) is a term encompassing seven clinically heterogeneous groups of arthritides of unknown cause in children (1). For patients with polyarticular-course (p)JIA (JIA of any category with  $\geq 5$  affected joints) (1), methotrexate (MTX) is the recommended first-line of disease-modifying antirheumatic drug (DMARD) therapy (2,3). Addition of a biologic (b) DMARD (tumor necrosis factor inhibitor [TNFi], anti-interleukin-6 inhibitors, or abatacept) is suggested if moderate or high disease activity persists after 3 months of treatment with MTX (2,3). Concomitant corticosteroid use is permitted with both MTX and bDMARDs, if required. Due to the chronic nature of pJIA, treatment agents are usually administered for a prolonged time, and blood concentrations achieved with bDMARDs may vary greatly between individual patients (4); thus, the safety of the treatment option is of utmost importance.

Infections have been shown to be the most frequent adverse event (AE) associated with some non-bDMARD and bDMARD treatments in both adult and pediatric patient populations (5-9). While information regarding the association between bDMARD treatment and infections in pediatric patients with JIA is limited, varied data exist for adult patients with rheumatoid arthritis (RA). A database study of 703 patients with RA indicated that high biologic drug levels (arbitrarily defined using concentration-effect curves for each drug), compared with low/normal levels, were associated with a higher risk of infection (10). In a multicenter retrospective cohort study of patients with RA in Japan, the risk of overall hospitalized infections did not correlate with the specific bDMARD, but the use of adalimumab was significantly associated with a greater risk of pulmonary hospitalized infections versus other agents (11). However, a recent study of data from administrative health databases suggested that among patients with RA treated with bDMARDs, abatacept was associated with the lowest risk of hospitalized infections across all studied biologic agents (12).

Abatacept is an immunomodulator that disrupts the continuous cycle of T-cell activation that characterizes rheumatic diseases, thereby inhibiting the production of B-cell-derived autoantibodies and pro-inflammatory cytokines (13-15). In an integrated data analysis of 9

RA clinical trials, no increased risk of infections, including opportunistic infections, with abatacept versus placebo was identified (16). Risk of infections is a particularly important consideration in pediatric patients due to their susceptibility to infections (17). Patients with JIA also have been reported to have an increased risk of hospitalized bacterial infections, compared with children without JIA, independent of treatment with MTX, corticosteroids or TNFi (18).

Intravenous (IV) abatacept has been proven to be effective and well tolerated in patients with pJIA in clinical trial and real-world settings (19-22). Furthermore, in patients with pJIA, weight-tiered subcutaneous (SC) abatacept has achieved the target therapeutic exposure threshold and has been effective and well tolerated (23). Results of clinical trials have shown that, compared with adult patients with RA treated with SC abatacept, patients aged 2–5 years with pJIA who received SC abatacept had a numerically higher rate of minor infections; no increase in infection rate was noted in patients aged 6–17 years (23,24).

It is not known whether infection risk is linked to the level of abatacept exposure in patients with pJIA following the approved SC or IV dosing regimen. As such, the aims of this analysis were to assess the relationship between the incidence of infection, regardless of seriousness, and levels of abatacept exposure in patients with pJIA following SC and IV abatacept treatment, and to compare the risk of infection between the IV and SC routes of administration.

## MATERIALS AND METHODS

*Data sources.* The data from two previously published studies of abatacept in pediatric patients were analyzed (19,23). One study included in this analysis was a Phase III, single-arm, open-label, international, multicenter, 2-part study comprised of 2 age cohorts of patients with pJIA (patients aged 6–17 years and patients aged 2–5 years; ClinicalTrials.gov identifier: NCT01844518) (23). Patients received SC abatacept based on body-weight tier (10–<25 kg [50 mg], 25–<50 kg [87.5 mg], and ≥50 kg [125 mg]) weekly for 4 months (23).

Responders with at least 30% improvement per American College of Rheumatology Criteria

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for JIA (JIA-ACR30) at Month 4 could receive SC abatacept for another 20 months; JIA-ACR30 non-responders at Month 4 were given the option to continue SC abatacept for an additional 3 months and discontinued treatment if a JIA-ACR30 response was not achieved by Month 7 (23). The other study included in this analysis was a Phase III, double-blind, randomized, placebo-controlled withdrawal trial, in which patients with pJIA aged 6–17 years received IV abatacept 10 mg/kg monthly (ClinicalTrials.gov identifier: NCT00095173) (19). JIA-ACR30 responders at Month 4 were randomized to receive either abatacept or placebo for 6 months, or until a flare occurred (19). Patients in this study also had the option of continuing abatacept in an open-label, long-term extension. In both studies, patients who were taking oral glucocorticoids at baseline were allowed to remain on a stable dose (0.2 mg/kg/day or 10 mg/day prednisone equivalent, whichever was lower) throughout the study. Adjustments to glucocorticoid doses were permitted in the IV study during the long-term extension phase, provided the total prednisone equivalent dose was  $\leq 10$  mg/day. Short courses ( $< 2$  weeks) of oral glucocorticoids ( $\geq 0.5$  mg/kg/day) were permitted in the SC study if clinically indicated.

Eligibility criteria have been described previously (19,23). Briefly, in both trials, patients had a history of  $\geq 5$  joints with active articular disease at baseline (defined as  $\geq 2$  active joints and  $\geq 2$  joints with limitation of motion), and were naïve to abatacept treatment, but may have had an inadequate response or prior intolerance to  $\geq 1$  non-biologic DMARD or bDMARD, including TNFi. Patients diagnosed with systemic JIA must have had an absence of systemic features for  $\geq 6$  months prior to enrollment. Patients were enrolled from 48 centers in 12 countries for the SC trial and from 43 centers in 11 countries for the IV trial, including the Paediatric Rheumatology International Trials Organisation (PRINTO) (25) and the Pediatric Rheumatology Collaborative Study Group (PRCSG) sites (26).

Both studies were conducted in accordance with the Declaration of Helsinki, the International Conference on Harmonisation Guidelines for Good Clinical Practice, and local regulations. At every study site, the protocol and amendments were reviewed and approved by the relevant independent review boards or ethics committees (19,23). For a full list of the



institutional review boards involved in these studies, please see the Supplementary Material for this article (note, the review boards did not provide ethics approval numbers).

*Assessments and statistical analyses.* The data from the two studies were analyzed both pooled and separately by route of abatacept administration. Data for SC abatacept were evaluated by age (6–17 years and 2–5 years) due to the previously noted numerically higher infection rate among 2–5-year-old patients with JIA treated with SC abatacept compared with patients in this study aged 6–17 years (23). The association between serum abatacept exposure measures and time to first infection (regardless of seriousness) was assessed. The relationship between levels of abatacept exposure measures and the occurrence or absence of infection was also investigated. The following serum abatacept exposure measures, estimated by population pharmacokinetic analysis, were used: steady-state maximum serum concentration ( $C_{\max ss}$ ), steady-state trough serum concentration ( $C_{\min ss}$ ), and steady-state average serum concentration ( $C_{av ss}$ ).

All patients who received at least one dose of study drug in the two studies were evaluated for safety, including infections. In the SC trial, infections were recorded as events of special interest, whereas in the IV trial, infections were recorded as part of routine AE reporting and then re-classified as events of special interest thereafter. In the SC trial, opportunistic infections were defined according to recent consensus statements in adults and children (27,28). In the IV study, similar organisms to those reported in the SC study, including active cytomegalovirus, active *Pneumocystis carinii*, aspergillosis or atypical mycobacterium infection, were considered to be opportunistic infections. The potential effects of a concomitant stable dose of MTX and glucocorticoid treatment (at baseline and throughout) on the association between abatacept exposure measures and time to first infection (regardless of seriousness) were evaluated.

Kaplan–Meier (KM) plots of probability of first infection versus time on treatment by abatacept exposure quartiles were created over time to Month 24 (pooled and separately by SC and IV abatacept administration). Quartiles of abatacept exposure were generated based

on drug concentrations at steady-state. Patients who experienced an infection had their exposure censored at the time of the first onset of the event. Log-rank tests were performed to evaluate the differences in probability of first infection at any time point across abatacept exposure quartiles. Exposure–response analysis in the safety analysis dataset confirmed a high degree of correlation among exposure parameters (Supplementary Figure 1); when combining the SC and IV data, the analysis between level of abatacept exposure measures suggested that  $C_{minss}$  was correlated with  $C_{avss}$  and  $C_{maxss}$  ( $r > 0.66$  or  $r < -0.66$ ). However,  $C_{maxss}$  and  $C_{avss}$  were not highly correlated ( $r = -0.19$ ), which was expected due to differences in SC and IV routes of administration.  $C_{maxss}$  was chosen as the main exposure measure in this analysis. The incidence of multiple infections was also investigated by quartiles of abatacept exposure and age categories (2–17 years, 2–5 years, and 6–17 years). Exploratory graphical analyses (box plots) of the relationship between level of abatacept exposure measures and the occurrence or absence of infection to Month 24 were performed. Quartiles of abatacept exposure in which infectious serious AEs occurred were also examined. The proportions of patients with infections deemed related to abatacept exposure were reported.

## RESULTS

*Patient disposition and baseline characteristics.* Overall, there were 219 patients in the SC abatacept trial (age 6–17 years,  $n = 173$ ; age 2–5 years,  $n = 46$ ) and 190 patients in the IV abatacept trial. A total of 219 patients treated with SC abatacept and 124 of those treated with IV abatacept were included in the current analysis. From the IV abatacept study, 62 patients who did not receive continuous abatacept treatment (patients randomized to receive placebo at Month 4) and 4 patients who did not have abatacept exposure measures were excluded from this analysis. Overall baseline demographics and disease characteristics have been reported previously (19,23). Briefly, in the SC and IV abatacept studies, respectively, median (minimum, maximum) age was 11.0 (2.0, 17.0) and 13.0 (5.0, 17.0) years, disease duration was 1.0 (0.0, 15.0) and 3.0 (0.0, 14.0) years, Childhood Health

Assessment Questionnaire-Disability Index was 1.0 (0.0, 2.9) and 1.2 (0.0, 2.9), and C-reactive protein was 0.2 (0.1, 21.1) and 1.3 (0.0, 26.0) mg/dL ( $\leq 0.6$  mg/dL defined as normal by the central laboratory), respectively.

In the SC and IV abatacept studies, a total of 172/219 (78.5%) and 91/124 (73.4%) patients, respectively, received concomitant MTX treatment at baseline, at mean (standard deviation [SD]) doses of 12.3 (4.1) and 13.6 (4.6) mg/m<sup>2</sup>/week. Over the 24-month period, 177/219 (80.8%) patients in the SC study received MTX treatment at a mean (SD) dose of 12.1 (4.0) mg/m<sup>2</sup>/week; the median (Q1–Q3) duration of MTX treatment among these patients was 717 (414–722) days. In the IV study, 92/124 (74.2%) received MTX in the 24-month period at a mean (SD) dose of 13.4 (5.1) mg/m<sup>2</sup>/week for a median (Q1–Q3) of 729 (211–729) days. Overall, 60/219 (27.4%) patients in the SC abatacept study and 60/124 (48.4%) patients in the IV abatacept study received oral glucocorticoids at baseline. The mean (SD) doses of oral glucocorticoids (prednisone equivalents) at baseline were 0.15 (0.08) and 0.15 (0.07) mg/kg/day in the SC and IV abatacept groups, respectively. Over the 24-month period, 82/219 (37.4%) patients in the SC abatacept group and 65/124 (52.4%) patients in the IV abatacept group received oral glucocorticoids for a median (Q1–Q3) of 359 (78–722) and 393 (85–729) days; mean (SD) doses were 0.19 (0.16) and 0.17 (0.14) mg/kg/day, respectively. A total of 58/219 (26.5%) patients in the SC abatacept study and 49/124 (39.5%) in the IV abatacept study received both concomitant MTX and glucocorticoids at baseline; over 24 months, 71/219 (32.4%) patients in the SC abatacept group and 52/124 (41.9%) patients in the IV abatacept group received both concomitant MTX and glucocorticoids for a median (Q1–Q3) of 361 (108–722) and 449 (119–729) days, respectively.

*Abatacept exposure and infections: pooled SC and IV analysis.* Overall, 237/343 (69.1%) patients treated with either SC or IV abatacept had at least 1 infection over 24 months. No statistically significant ( $p = 0.4458$ ) difference in probability of first infection at any time point across the 4 quartiles of abatacept exposure ( $C_{\text{maxss}}$ ) in the 2–17-year-old patient population was observed (Figure 1; Table 1). Similar findings in infection probability across the 4

quartiles were also observed for  $C_{\min ss}$  and  $C_{av ss}$  (Supplementary Figure 2). The median (95% confidence interval [CI]) time to first infection in the lowest to highest quartiles of abatacept exposure ( $C_{\max ss}$ ) was 244 (129, 462), 152 (106, 216), 162 (91, 246), and 186 (82, 224) days, respectively. There was no apparent relationship between the numbers of infections per patient seen with increasing exposure to abatacept (Supplementary Table 1). Median level of exposure measures by infection status (occurrence or absence) did not show an association between abatacept exposure and occurrence of infection (Figure 2). Median values and distributions of level of abatacept exposure measures were similar in patients who experienced infections versus those who did not. A total of 3 infectious serious AEs were reported: varicella in a 6-year-old patient treated with IV abatacept ( $C_{\max ss} = 213$   $\mu\text{g/mL}$ ; quartile 2), cellulitis in a 2-year-old patient treated with SC abatacept ( $C_{\max ss} = 49.7$   $\mu\text{g/mL}$ ; quartile 1), and appendicitis in a 17-year-old patient treated with SC abatacept ( $C_{\max ss} = 70.5$   $\mu\text{g/mL}$ ; quartile 4). All 3 serious infections were resolved and deemed unrelated to abatacept treatment. No opportunistic infections related to abatacept were reported (29,30), including no cases of herpes zoster in either study during the 24-month period.

*Abatacept exposure and infections: analysis by route of administration.* A total of 156/219 (71.2%) and 81/124 (65.3%) patients receiving SC and IV abatacept, respectively, had at least 1 infection over 24 months. No statistically significant difference in probability of first infection at any time point was observed among the exposure quartiles for  $C_{\max ss}$  following SC (Figure 3; Table 1) or IV (Figure 4; Table 1) abatacept. Likewise, KM curves were similar across the quartiles for  $C_{\min ss}$  and  $C_{av ss}$  (Supplementary Figure 3; Supplementary Figure 4). Median level of exposure measures and distribution by occurrence or absence of infection did not show an association between abatacept exposure and occurrence of infection in either the SC or IV populations (data not shown). Infections considered related to abatacept exposure were reported in 60/219 (27.4%) and 18/124 (14.5%) patients receiving SC and IV abatacept, respectively.

*Abatacept exposure and infections: analysis of SC abatacept by age.* Among patients aged 6–17 years, a total of 116/173 (67.1%) patients experienced an infection within the 24-month cumulative period; no statistically significant ( $p = 0.4787$ ) difference in probability of first infection at any time point across abatacept exposure quartiles ( $C_{\max ss}$ ) was evident (Figure 3; Table 1). In these patients, the median (95% CI) time to first infection in the lowest to highest quartiles of abatacept exposure ( $C_{\max ss}$ ) was 430 (156, 577), 108 (63, 291), 204 (113, 466), and 266 (152, 448) days, respectively. A higher proportion of patients aged 2–5 years, compared with patients aged 6–17 years, experienced an infection within the 24-month cumulative period (40/46; 87.0%); however, no statistically significant ( $p = 0.9305$ ) difference in probability of first infection at any time point across abatacept exposure quartiles was observed (Figure 3; Table 1). Among the patients aged 2–5 years, the median (95% CI) time to first infection in the lowest to highest quartiles of abatacept exposure ( $C_{\max ss}$ ) was 77 (10, 511), 77 (27, 439), 87 (36, 200), and 67 (25, 245) days, respectively. The number of infections per patient did not show an association with abatacept exposure quartiles (Supplementary Table 1).

*Abatacept exposure and infections: analysis of pooled SC and IV abatacept by concomitant MTX and glucocorticoid use at baseline and throughout.* Overall, an infection was reported within the 24-month period in 73/107 (68.2%) patients receiving triple immunosuppression with abatacept, concomitant MTX and glucocorticoids in this analysis. No difference in probability of first infection at any time point across the 4 quartiles of abatacept exposure measures was observed (Figure 5; Supplementary Figure 5). Similarly, no difference in probability of first infection at any time point across the 4 quartiles of abatacept exposure measures was seen in patients receiving SC or IV abatacept monotherapy (Supplementary Figure 6; Supplementary Figure 7). Some differences in the shapes of KM curves for abatacept monotherapy analyses (Supplementary Figure 7), compared with all other analyses, could be attributed to very small sample sizes.

## DISCUSSION

Among patients aged 2–17 years with pJIA who received the approved SC or IV abatacept dose, with the possibility of MTX and/or glucocorticoids, no association of level of abatacept exposure with risk of infections over a 24-month period was seen. While patients aged 2–5 years had a numerically greater rate of infections than patients aged 6–17 years (23), as one might expect to observe in these two age groups among the general population, the level of abatacept exposure was not associated with time to first infection or occurrence of multiple infections in either population. The median values and distributions of abatacept exposure measures were similar between patients with pJIA, in whom infection occurred and those in whom infections were not reported. There was no consistent association with infectious serious AEs with abatacept level of exposure.

Infections and infestations are the most frequently reported AEs associated with abatacept treatment in pediatric patients, with nasopharyngitis and upper respiratory tract infections being the most common (19,23). No new cases of opportunistic infections, including herpes zoster and tuberculosis, were reported in either SC or IV studies, despite the presence of some study sites in tuberculosis-endemic, or high tuberculosis incidence, locations (19,23). In addition, an integrated data analysis of 9 clinical trials identified no increased risk of herpes zoster infection in abatacept-treated patients with RA, compared with patients receiving placebo (16). A large analysis combining data from clinical trials and registries revealed that in pediatric patients, abatacept had a similar infection profile to both adalimumab and etanercept; all 3 of these agents had favorable infection profiles compared with golimumab, infliximab, and tocilizumab (31). Consistent with the results seen with abatacept, IV and SC tocilizumab treatment in patients with RA resulted in similarly low rates of serious infections (32).

Due to a potential immunosuppressive effect, MTX and glucocorticoids may lead to a possible increase in infection risk, particularly in a susceptible population (33,34); as such, it is important to investigate their use concomitantly with bDMARDs. A systematic literature

review and meta-analysis of randomized controlled trials demonstrated that combination therapy with bDMARDs and MTX did not increase the risk of serious infections versus bDMARD monotherapy (35), in line with the results observed in this analysis (Supplementary Figures 5 and 6). In addition, a systematic literature review of 88 studies showed that long-term ( $\geq 2$  years) use of MTX monotherapy was not a risk factor for serious infections, including herpes zoster (36). This is potentially due to the fact that the mechanism of action of MTX in RA may be mediated by its anti-inflammatory rather than immunosuppressive properties (37). Conversely, the use of glucocorticoids has been associated with increased risk of infections in patients with RA (31,38-40) and JIA (18). In a longitudinal study of complete patient medical records, a dose-dependent association of use of glucocorticoids with risk for serious infections was observed in patients with RA (41); in addition to this, an association between herpes zoster infection and glucocorticoid use was also seen (42). In this study, concomitant use of MTX, glucocorticoids and abatacept did not markedly increase risk of infection across the level of abatacept exposure quartiles, which may indicate that even a relatively high degree of immunosuppression is well tolerated in this population (Supplementary Figure 5).

The limitations of this study should be considered. First, these were *post hoc* analyses and neither study was designed or powered specifically to investigate time to first infection. Due to the relatively small sample size in each quartile of the patients aged 2–5 years, these data must be interpreted with caution. In addition, due to the sample size, analysis of specific infections was not conducted. It should also be considered that due to limited patient exposure to triple immunosuppression with concomitant MTX and glucocorticoids in this analysis, detection of immunosuppressive effects may have also been limited. In addition, patients were restricted to relatively low doses of glucocorticoids (IV pulse steroids were not permitted); as a result, the effects of high-dose glucocorticoids could not be studied. However, long-term treatment with high doses of glucocorticoids is not desirable in children due to its negative impact on growth and development and well-known AEs.



In conclusion, no association of abatacept (SC or IV) exposure with risk of infections, including opportunistic infections, was found in pediatric patients aged 2–17 years with pJIA over a 24-month period, including with concomitant MTX and glucocorticoid treatment at baseline and throughout. These findings provide further support for the use of abatacept in patients as young as 2 years with pJIA.

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## AUTHORS' CONTRIBUTIONS

The study was designed jointly by academic authors and Bristol-Myers Squibb Company, with data collected by PRINTO/PRCSG investigators. All authors attest to the completeness and veracity of data and data analyses. Consistency in reporting the study data to healthcare authorities and institutional review boards was ensured by Bristol-Myers Squibb Company. All authors had full access to study data, reviewed and revised the manuscript, and approved the final version to be published. All authors were involved in the decision to submit the manuscript for publication and had the right to accept or reject comments or suggestions.

All authors were involved in drafting the article or revising it critically for important intellectual content, and all authors approved the final version to be published.

Study conception and design: NR, HIB, RW, AM, DJL

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Acquisition of data: NR, HIB, NT, GVC, IL, RC, JD, GE, EF, MFZ, VG, PQ, CAS, LWW, YG, AM, DJL

Analysis and interpretation of data: NR, HIB, NT, GVC, IL, RC, JD, GE, EF, MFZ, VG, PQ, CAS, LWW, YG, JP, MN, RW, AM, DJL

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## FIGURE LEGENDS

**Figure 1.** Pooled SC and IV analysis: Kaplan–Meier plots of probability of first infection,\* from first dose, by abatacept exposure quartiles over 24 months, 2–17-year-old patient population ( $C_{\max ss}$ ). For PK ranges, a square bracket indicates the respective endpoint is included in the interval. A regular bracket indicates the respective endpoint is not included in the interval. Q1 and Q4 represent the lowest and the highest quartile, respectively.

\*Regardless of seriousness.  $C_{\max ss}$ : steady-state maximum serum concentration; IV: intravenous; PK: pharmacokinetic; Q: quartile; SC: subcutaneous.

**Figure 2.** Pooled SC and IV analysis: boxplots of abatacept exposure measurements versus the occurrence of infections over 24 months (2–17-year-old patient population): A)  $C_{\min ss}$ , B)  $C_{\max ss}$ , C)  $C_{av ss}$ .

$C_{av ss}$ : steady-state time-averaged serum concentration;  $C_{\max ss}$ : steady-state maximum serum concentration;  $C_{\min ss}$ : steady-state trough serum concentration; IV: intravenous; SC: subcutaneous.

**Figure 3.** SC abatacept analysis: Kaplan–Meier plots of probability of first infection,\* from first dose, by abatacept exposure quartiles over 24 months ( $C_{\max ss}$ ): A) 2–5-year-old patient population, B) 6–17-year-old patient population. For PK ranges, a square bracket indicates respective endpoint is included in the interval. A regular bracket indicates respective endpoint is not included in the interval. Q1 and Q4 represent the lowest and the highest quartile, respectively. \*Regardless of seriousness.  $C_{\max ss}$ : steady-state maximum serum concentration; PK: pharmacokinetic; Q: quartile; SC: subcutaneous.

**Figure 4.** IV abatacept analysis: Kaplan–Meier plots of probability of first infection,\* from first dose, by abatacept exposure quartiles over 24 months, 6–17-year-old patient population ( $C_{\max ss}$ ). For PK ranges, a square bracket indicates respective endpoint is included in the interval. A regular bracket indicates respective endpoint is not included in the interval. Q1

and Q4 represent the lowest and the highest quartile, respectively. \*Regardless of seriousness.  $C_{\text{maxss}}$ : steady-state maximum serum concentration; IV: intravenous; PK: pharmacokinetic; Q: quartile.

**Figure 5.** Pooled SC and IV analysis: Kaplan–Meier plots of probability of first infection,\* from first dose, by abatacept exposure quartiles over 24 months, 2–17-year-old patient population with concomitant MTX and glucocorticoids at baseline and throughout ( $C_{\text{maxss}}$ ).

For PK ranges, a square bracket indicates respective endpoint is included in the interval. A regular bracket indicates respective endpoint is not included in the interval. Q1 and Q4 represent the lowest and the highest quartile, respectively. \*Regardless of seriousness.

$C_{\text{maxss}}$ : steady-state maximum serum concentration; IV: intravenous; MTX: methotrexate; PK: pharmacokinetic; Q: quartile; SC: subcutaneous.

Table 1. Median time to first infection for C<sub>maxss</sub> quartiles.

Exposure quartile	C <sub>maxss</sub> range (µg/mL)	Median (95% CI) time to first infection (days)	Log-rank p-value
Pooled SC and IV, 2–17-year-old population			
Q1	[19.1, 49.7]	244 (129, 462)	0.4458
Q2	(49.7, 65.8]	152 (106, 216)	
Q3	(65.8, 198.0]	162 (91, 246)	
Q4	(198.0, 307.0]	186 (82, 224)	
SC, 2–5-year-old population			
Q1	[37.7, 54.4]	77 (10, 511)	0.9305
Q2	(54.4, 61.1]	77 (27, 439)	
Q3	(61.1, 69.2]	87 (36, 200)	
Q4	(69.2, 119.0]	67 (25, 245)	
SC, 6–17-year-old population			
Q1	[19.1, 42.4]	430 (156, 577)	0.4787
Q2	(42.4, 52.8]	108 (63, 291)	
Q3	(52.8, 62.5]	204 (113, 466)	
Q4	(62.5, 101.0]	266 (152, 448)	
IV, 6–17-year-old population			
Q1	[119.0, 192.5]	188 (91, 473)	0.4999
Q2	(192.5, 214.5]	76 (51, 211)	
Q3	(214.5, 238.5]	186 (49, 684)	
Q4	(238.5, 307.0]	192 (81, 224)	

For PK ranges, a square bracket indicates respective endpoint is included in the interval. A regular bracket indicates respective endpoint is not included in the interval. Q1 and Q4

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represent the lowest and the highest quartile, respectively. CI: confidence interval;  $C_{\max ss}$ : steady-state maximum serum concentration; IV: intravenous; PK: pharmacokinetic; Q: quartile; SC; subcutaneous.

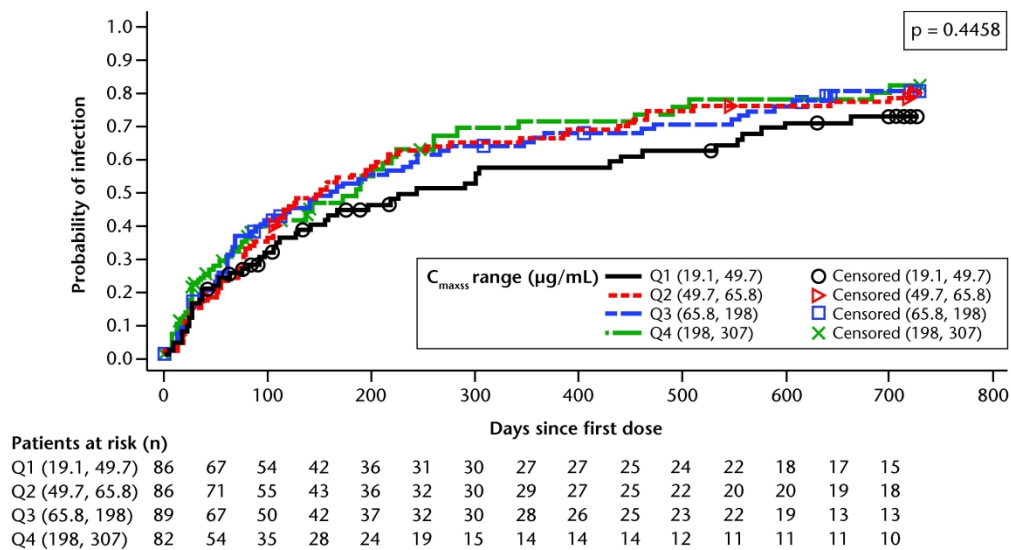


Figure 1. Pooled SC and IV analysis: Kaplan–Meier plots of probability of first infection,\* from first dose, by abatacept exposure quartiles over 24 months, 2–17-year-old patient population (Cmaxss). For PK ranges, a square bracket indicates the respective endpoint is included in the interval. A regular bracket indicates the respective endpoint is not included in the interval. Q1 and Q4 represent the lowest and the highest quartile, respectively. \*Regardless of seriousness. Cmaxss: steady-state maximum serum concentration; IV: intravenous; PK: pharmacokinetic; Q: quartile; SC: subcutaneous.

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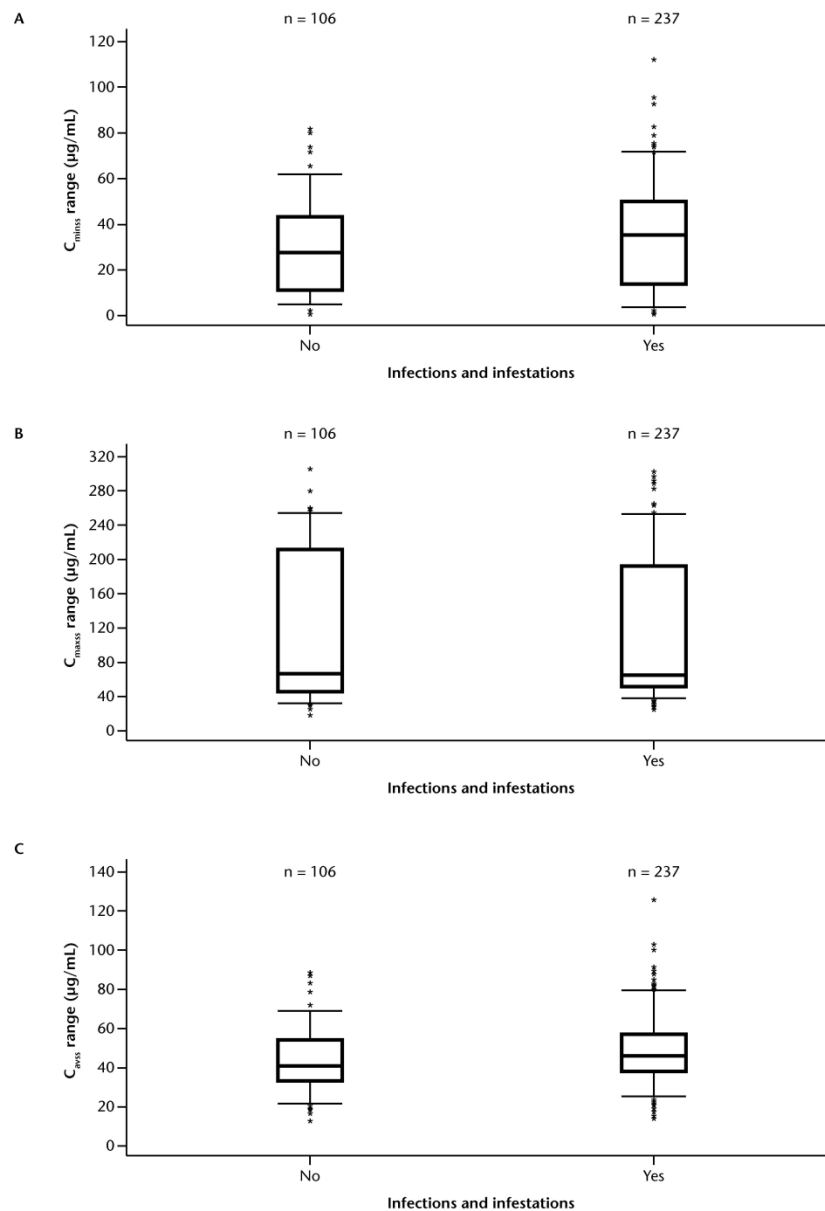


Figure 2. Pooled SC and IV analysis: boxplots of abatacept exposure measurements versus the occurrence of infections over 24 months (2–17-year-old patient population): A) C<sub>minss</sub>, B) C<sub>maxss</sub>, C) C<sub>avss</sub>.

C<sub>avss</sub>: steady-state time-averaged serum concentration; C<sub>maxss</sub>: steady-state maximum serum concentration; C<sub>minss</sub>: steady-state trough serum concentration; IV: intravenous; SC: subcutaneous.

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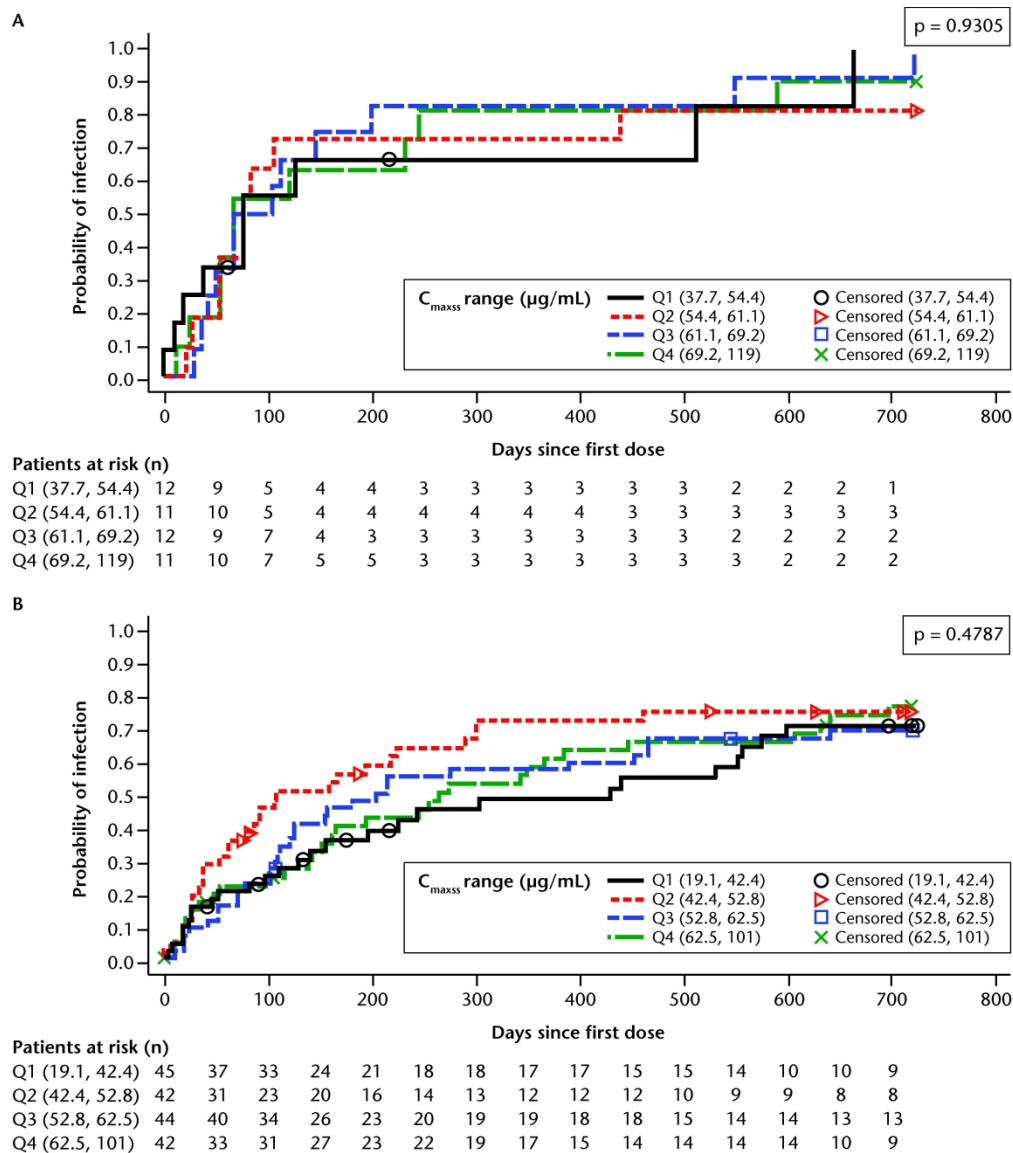


Figure 3. SC abatacept analysis: Kaplan–Meier plots of probability of first infection,\* from first dose, by abatacept exposure quartiles over 24 months ( $C_{maxss}$ ): A) 2–5-year-old patient population, B) 6–17-year-old patient population. For PK ranges, a square bracket indicates respective endpoint is included in the interval. A regular bracket indicates respective endpoint is not included in the interval. Q1 and Q4 represent the lowest and the highest quartile, respectively. \*Regardless of seriousness.  $C_{maxss}$ : steady-state maximum serum concentration; PK: pharmacokinetic; Q: quartile; SC: subcutaneous.

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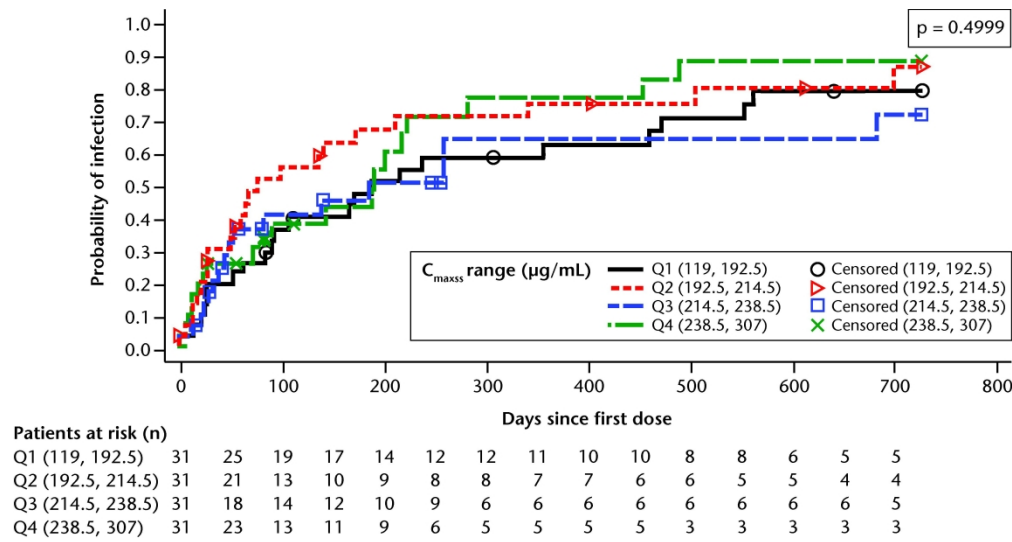


Figure 4. IV abatacept analysis: Kaplan-Meier plots of probability of first infection,\* from first dose, by abatacept exposure quartiles over 24 months, 6–17-year-old patient population ( $C_{maxss}$ ). For PK ranges, a square bracket indicates respective endpoint is included in the interval. A regular bracket indicates respective endpoint is not included in the interval. Q1 and Q4 represent the lowest and the highest quartile, respectively. \*Regardless of seriousness.  $C_{maxss}$ : steady-state maximum serum concentration; IV: intravenous; PK: pharmacokinetic; Q: quartile.

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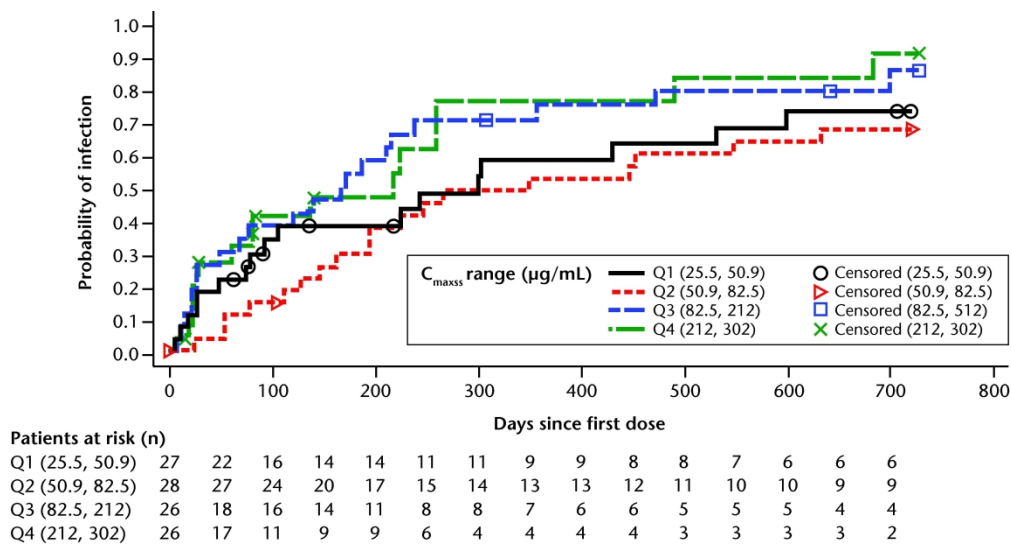


Figure 5. Pooled SC and IV analysis: Kaplan–Meier plots of probability of first infection,\* from first dose, by abatacept exposure quartiles over 24 months, 2–17-year-old patient population with concomitant MTX and corticosteroids (Cmaxss). For PK ranges, a square bracket indicates respective endpoint is included in the interval. A regular bracket indicates respective endpoint is not included in the interval. Q1 and Q4 represent the lowest and the highest quartile, respectively. \*Regardless of seriousness. Cmaxss: steady-state maximum serum concentration; IV: intravenous; MTX: methotrexate; PK: pharmacokinetic; Q: quartile; SC: subcutaneous.

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