Drug Interaction Potential as a Result of Cytokine Release Syndrome Using a Physiologically Based Pharmacokinetic Model: Case Study of Teclistamab

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INTRODUCTION

- The ability of a physiologically based pharmacokinetic (PBPK) model to capture drug-drug interactions (DDIs) was first verified using data from the literature^{1,2}
- Teclistamab is the first off-the-shelf B-cell maturation antigen (BCMA) × CD3 bispecific antibody approved, and is a new standard of care (overall response rate, 63%) for patients with heavily pretreated relapsed/refractory multiple myeloma³⁻⁵
- Cytokine release syndrome (CRS) has been associated with teclistamab, with most events being low grade and reported during step-up doses and the initial treatment dose at the recommended phase 2 dose (RP2D)
- Among the different cytokines released during CRS, interleukin 6 (IL-6) is a potent suppressor of cytochrome P450 (CYP450) enzyme activity
- A PBPK model was developed to evaluate the impact of IL-6 following teclistamab (administered at the RP2D in MajesTEC-1) on the exposure to co-administered CYP450 substrates

METHODS

- A PBPK model was developed for IL-6 based on the literature⁶
- DDIs toward different CYP450 substrates (caffeine [CYP1A2], s-warfarin [CYP2C9], omeprazole [CYP2C19], midazolam and cyclosporine [CYP3A4 and CYP3A5], and simvastatin [CYP3A4]) were evaluated where substrates were administered as a single dose when minimum (or maximum for CYP1A2) enzymatic activity was reached
- Inactivation of CYP2C9, CYP2C19, CYP3A4, and CYP3A5, but induction of CYP1A2, has been observed in vitro and in vivo^{1,7}
- Prospective simulations were performed using observed IL-6 profiles from 112 patients following teclistamab administered at the RP2D without or prior to administration of tocilizumab (an anti-IL-6 agent) in MajesTEC-1 as of the March 16, 2022 cut-off
- Observed data were recovered by adjusting the dosing regimens of IL-6, modeled as an intravenous infusion. Two scenarios were envisioned:
- The observed mean serum IL-6 kinetic profile
- The worst-case scenario (in terms of risk of a DDI) corresponding to the IL-6 profile observed in the patient presenting the highest IL-6 maximum concentration (C_{max}) value
- In addition, time to reach maximum change in CYP450 activity due to IL-6 and return to 80% of the baseline enzymatic activity was evaluated, with the start of cycle 1 as reference
- 80% was considered as a reasonable cut-off at which DDI liability would be low

MULTIPLE MYELOMA

RESULTS

Model verification

- Transient peak IL-6 concentration and IL-6 concentration at steady state (50 pg/mL) were successfully predicted
- Observed DDIs from literature with CYP1A2, CYP2C9, CYP2C19, CYP3A4, and CYP3A5 substrates in the presence of steady state concentrations of IL-6 at 50 pg/mL were well predicted, providing confidence in the application of the model to assess IL-6 as a perpetrator of these CYP substrates

Model application to teclistamab RP2D

- The mean IL-6 profile following teclistamab at RP2D (mean C_{max}= 21 pg/mL) was predicted to result in a limited change in exposure of CYP1A2, CYP2C9, CYP2C19, CYP3A4, and CYP3A5 substrates (0.87 ≤ area under the curve [AUC] ratio ≤ 1.20) (Table)
- The IL-6 kinetic profile with the highest C_{max} (288 pg/mL) was predicted to result in:
- A mild-to-moderate inhibition on exposure of CYP2C19, CYP3A4, and CYP3A5 substrates (1.90 ≤ AUC ratio ≤2.23) (Table)
- Minimal impact on exposure of CYP1A2 and CYP2C9 substrates (AUC ratio=0.82 and 1.25, respectively) (Table)
- For both scenarios, the maximum change in exposure for the studied substrates occurred 3–4 days after the first treatment dose (Figure 1A)
- For the patient with the highest IL-6 C_{max}, return to 80% of the baseline enzymatic activity was observed ~7 days after the first treatment dose (Figure 1B)
- The highest AUC ratio (at 80% of enzymatic activity) reached was ≤1.47 (CYP2C19), which corresponds to a weak interaction

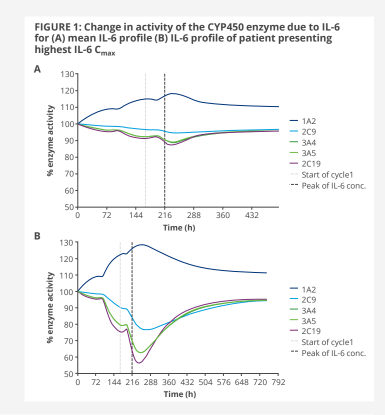


TABLE: Simulated change in CYP450 substrate exposure after single dose administration of the CYP450 substrate in presence of IL-6 kinetics profile observed in MajesTEC-1

With mean IL-6 kinetics profile ^a			With IL-6 kinetics profile of patient presenting highest IL-6 C _{max} b		
C _{max} ratio, mean	AUC ratio, mean	DDI liability	C _{max} ratio, mean	AUC ratio, mean	DDI liability
0.97	0.87	No interaction	0.95	0.82	No interaction
1.00	1.05	No inhibition	1.01	1.25	Weak inhibition
1.10	1.20	No inhibition	1.45	2.23	Moderate inhibition
1.11	1.17	No inhibition	1.46	1.90	Weak inhibition
1.17	1.20	No inhibition	1.86	2.09	Moderate inhibition
1.09	1.17	No inhibition	1.35	1.90	Weak inhibition
	nean 0.97 1.00 1.10 1.11 1.17 1.09	mean mean 0.97 0.87 1.00 1.05 1.10 1.20 1.11 1.17 1.17 1.20	mean mean Dol liability 0.97 0.87 No interaction 1.00 1.05 No inhibition 1.10 1.20 No inhibition 1.11 1.17 No inhibition 1.17 1.20 No inhibition 1.09 1.17 No inhibition	C _{max} ratio, mean AUC ratio, mean DDI liability C _{max} ratio, mean 0.97 0.87 No interaction 0.95 1.00 1.05 No inhibition 1.01 1.10 1.20 No inhibition 1.45 1.11 1.17 No inhibition 1.46 1.17 1.20 No inhibition 1.86 1.09 1.17 No inhibition 1.35	C _{max} ratio, mean AUC ratio, mean DDI liability C _{max} ratio, mean AUC ratio, mean 0.97 0.87 No interaction 0.95 0.82 1.00 1.05 No inhibition 1.01 1.25 1.10 1.20 No inhibition 1.45 2.23 1.11 1.17 No inhibition 1.46 1.90 1.17 1.20 No inhibition 1.86 2.09 1.09 1.17 No inhibition 1.35 1.90

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KEY TAKEAWAY



Interactions between substrates of CYP450 and IL-6 released when teclistamab is given at the RP2D are predicted to have limited clinical significance

CONCLUSIONS



The initial release of IL-6 during CRS following teclistamab treatment at the RP2D has minimal or moderate impact on exposure to CYP substrates



The highest risk of DDI is expected to occur from day 1 of the step-up dosing schedule to 7 days after the first treatment dose and during and after CRS



During this time period, for substrates with a narrow therapeutic index, such as warfarin and cyclosporine:

- Patients should be monitored for signs of toxicity
- Concentrations of these CYP substrates with a narrow therapeutic index should be assessed

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DISCLOSURES

M-EW is employed by Janssen. SXWL, JDG, and SG were employed by Janssen when this work was done and have stock/other ownership interests in Janssen. LDZ, LSW, XM, RV, and BL are employed by Janssen. AB, RK, and MQ are employed by and have stock/other ownership interests in Janssen. DO is employed by and has stock/other ownership interests in Janssen and has stock/other ownership interests in Janssen and has stock/other ownership interests in Janssen and has stock/other ownership interests.

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