**Bench-to-bedside translation of chimeric antigen receptor (CAR) T cells using a multiscale systems pharmacokinetic-pharmacodynamic model: A case study with anti-BCMA CAR-T**

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Abstract

Despite tremendous success of chimeric antigen receptor (CAR) T cell therapy in clinical oncology, the dose-exposure-response relationship of CAR-T cells in patients is poorly understood. Moreover, the key drug-specific and system-specific determinants leading to favorable clinical outcomes are also unknown. Here we have developed a multiscale mechanistic pharmacokinetic (PK)-pharmacodynamic (PD) model for anti-B-cell maturation antigen (BCMA) CAR-T cell therapy (bb2121) to characterize (i) *in vitro* target cell killing in multiple BCMA expressing tumor cell lines at varying effector to target cell ratios, (ii) preclinical *in vivo* tumor growth inhibition and blood CAR-T cell expansion in xenograft mice, and (iii) clinical PK and PD biomarkers in patients with multiple myeloma. Our translational PK-PD relationship was able to effectively describe the commonly observed multiphasic CAR-T cell PK profile in the clinic, consisting of the rapid distribution, expansion, contraction, and persistent phases, and accounted for the categorical individual responses in multiple myeloma to effectively calculate progression-free survival rates. Preclinical and clinical data analysis revealed comparable parameter estimates pertaining to CAR-T cell functionality and suggested that patient baseline tumor burden could be more sensitive than dose levels toward overall extent of exposure after CAR-T cell infusion. Virtual patient simulations also suggested a very steep dose-exposure-response relationship with CAR-T cell therapy and indicated the presence of a “threshold” dose, beyond which a flat dose-response curve could be observed. Our simulations were concordant with multiple clinical observations discussed in this article. Moving forward, this framework could be leveraged *a priori* to explore multiple infusions and support the preclinical/clinical development of future CAR-T cell therapies.
INTRODUCTION

Chimeric antigen receptor (CAR)-engineered T cells have recently demonstrated unprecedented efficacy in multiple haematological malignancies and have garnered tremendous recognition in the field of cancer immunotherapy.1 A typical CAR construct is composed of extracellular single-chain variable fragment of antibody, a transmembrane domain, and intracellular signalling domain containing CD3ζ linked with 0–2 costimulatory domains.2 Upon CAR engagement with the tumor-associated antigen (TAA), a nonclassical immune synapse is formed, which eventually leads to antitumoral effects by releasing perforins and granzymes. With the regulatory approvals of Yescarta (axicabtagene ciloleucel), Kymriah (tisagenlecleucel), and Tecartus (brexucabtagene autoleucel),3 the clinical landscape of CAR-T cells is burgeoning with ≥ 500 CAR candidates currently in clinical development.4 However, the pharmacokinetic (PK)-pharmacodynamic (PD) characterization of these constructs presents many challenges and unique opportunities because of the self-proliferating and long-term persistence capabilities in vivo.5,6

CAR-T cells exhibit a very unique clinical PK profile, which is often discerned by the rapid distribution, expansion, contraction, and persistence phases.7,8 Although mathematical models have been developed recently9,10 to empirically describe the slopes associated with multiphasic PK profiles for CAR-T cells, they have limited capability toward extrapolation to predict the PK and PD behavior of alternative CAR constructs and dose levels. The cellular kinetic behavior of cell therapies is dependent on several (i) drug-specific attributes, such as CAR-affinity, CAR-density, effector cell type (αβ T cell, γδ T cell, natural killer cells), costimulatory domains (CD28, 4-1BB); (ii) system-specific attributes such as disease type, tumor accessibility (solid tumor or hem malignancies), and initial tumor burden; as well as (iii) product-specific attributes such as CD4:CD8 ratios, phenotypic composition, transduction efficiency, in vitro effector doubling time, etc. Therefore, it is imperative to adopt a more mechanistic PK-PD approach5,6 while quantitatively describing CAR-T cell functionalities.

In our prior work,7 we presented a multiscale mechanistic modeling approach that incorporated key drug-specific (CAR-affinity, CAR-density) and system-specific (e.g., tumor burden) parameters to characterize preclinical CAR-T cell activity. The model exhibited the capability of characterizing the distribution of CAR-T cells to pertinent tissues and site of action,5 where upon interaction with tumor cells, the CAR-Target engagement drives the simultaneous killing of tumor cells and the expansion of CAR-T cells. The developed model was able to well characterize the multiphasic CAR-T cellular kinetic profile, and some of the model simulations suggested a very steep dose-exposure-response relationship.

In this article, a similar modeling paradigm has been extended to characterize clinical PK-PD data sets for CAR-T cells. A unified mechanism-based PK-PD model is developed to characterize the preclinical (in vitro and in vivo)11

Study Highlights

WHAT IS THE CURRENT KNOWLEDGE ON THE TOPIC?
Although a promising platform in immune-oncology, the drug-specific and system-specific determinants governing the observed dose-exposure-response relationship of chimeric antigen receptor (CAR)-T cell therapy in the clinic are poorly understood.

WHAT QUESTION DID THIS STUDY ADDRESS?
Using a multiscale translational pharmacokinetic (PK)-pharmacodynamic modeling approach, this study integrates key drug-specific and system-specific parameters responsible for CAR-T cell functionality in preclinical (in vitro and in vivo) and clinical settings. These analyses also provide mechanistic insights toward the observed multiphasic CAR-T cell PK profile as well as the generally observed steep dose-exposure-response relationship in the clinic.

WHAT DOES THIS STUDY ADD TO OUR KNOWLEDGE?
This analysis suggests the presence of a “threshold” CAR-T dose, beyond which a flat dose-exposure-response curve could be observed. In addition, baseline patient tumor burden could be more sensitive than administered dose levels toward the overall extent of CAR-T cell exposure.

HOW MIGHT THIS CHANGE DRUG DISCOVERY, DEVELOPMENT, AND/OR THERAPEUTICS?
The multiscale modeling framework could be applied in forward translation to support CAR-T cell dose selection (starting, escalation, and expansion cohorts), simulating multiple infusion regimens and reverse translation to inform optimal drug-specific characteristics.
and clinical PK-PD data sets for anti-B-cell maturation antigen (BCMA) (bb2121, Idecabtagene vicleucel) CAR-T cells. Later, model simulations were performed to assess the impact of dose and patient tumor burden on cellular kinetics and clinical responses. Virtual patient simulations were also performed to understand and characterize the dose-response relationship for bb2121 CAR-T cell therapy in patients with multiple myeloma. Simulations were later compared with recently reported pivotal clinical trial results for bb2121. The translational PK-PD modeling framework described in this article can be used toward forward-translation and reverse-translation of cell therapies in the future.

### METHODS

#### Preclinical and clinical data sets used for PK-PD model development

**In vitro cytotoxicity experiments**

A single timepoint (4-hour) killing assay of six BCMA + cell lines was evaluated (Table 1). CAR-T (bb2121) cells derived from three different T cell donors was cocultured with BCMA + cell lines to obtain mean cytolysis at effector to target cell (E:T) ratios ranging from 1:1 to 10:1. In vitro cytotoxicity experiments describes the simultaneous model-fitted cytolysis percentage profiles overlaid with observed data (Figure 2a1–2a6) from six BCMA + cell lines after coculture with bb2121 CAR-T cells at different E:T ratios. Our previously proposed in vitro modeling framework accounts for simultaneous CAR-T cell expansion and tumor cell depletion. However, because of the lack of an in vitro expansion data set of bb2121 CAR-T cells within the underlying experiment, the activated bb2121 CAR-T cell doubling time (DT<sub>CART</sub>) parameter was fixed (Table 1) to known values. In addition, while performing data analysis, the system-specific parameters such as CAR-affinity, BCMA receptor densities, and CAR densities were fixed to the known values (Table 1), whereas the parameters associated with the efficacy of CAR-T cells, i.e., maximum killing rate (K<sub>Kill</sub><sup>Max</sup>) and killing potency (K<sub>CART</sub><sup>50</sup>) were estimated. The estimated K<sub>CART</sub><sup>50</sup> revealed that ~ 2.24 CAR-Target complexes per tumor cells were required to achieve 50% of the maximum killing rate (K<sub>Kill</sub><sup>Max</sup>). The extent of the cytolsis percentage was not correlated with the antigen densities on six cell lines. Hence, ~ 60% interindividual variability (IVV) was estimated on the mean maximum killing rate (K<sub>Kill</sub><sup>Max</sup>) parameter. In the subsequent modeling analysis, the cell-level in vitro potency estimates (K<sub>CART</sub><sup>50</sup>) were fixed while characterizing in vivo CAR-T cell–induced tumor growth inhibition (TGI).

**Preclinical xenograft experiments to assess efficacy and expansion of CAR-T cells**

NOD scid gamma mice subcutaneously inoculated with BCMA + RPMI-8226 tumors were randomized (at 151 mm<sup>3</sup> tumor volume) for treatment with vehicle control or ~ 5 million bb2121 CAR-T cells/mouse. Tumor volumes were measured twice a week, and CD3+ CAR+ T cells were measured (using flow cytometry) to assess PK in blood.

**Phase 1 clinical trial in patients with relapsed or refractory (r/r) multiple myeloma**

In this clinical study (n = 33), anti-BCMA (bb2121) CAR-T cells were administered as a single infusion at flat dose levels of 50, 150, 450 and 800 million CAR+ cells per patient. PK was evaluated by measuring median vector transgene copies per µg of genomic DNA. Mean longitudinal data for PD biomarkers, i.e., soluble BCMA and serum M-protein, was reported as surrogates for antitumor activity. In addition, swimmer plots associated with individual patient categorical response data for 33 patients were also reported based on the International Myeloma Working Group (IMWG) Uniform Response Criteria for Multiple Myeloma.

### RESULTS

#### Model-based characterization of in vitro functional activity of bb2121

Figure 2a describes the simultaneous model-fitted cytolysis percentage profiles overlaid with observed data (Figure 2a1–2a6) from six BCMA + cell lines after coculture with bb2121 CAR-T cells at different E:T ratios. Our previously proposed in vitro modeling framework accounts for simultaneous CAR-T cell expansion and tumor cell depletion. However, because of the lack of an in vitro expansion data set of bb2121 CAR-T cells within the underlying experiment, the activated bb2121 CAR-T cell doubling time (DT<sub>CART</sub>) parameter was fixed (Table 1) to known values. In addition, while performing data analysis, the system-specific parameters such as CAR-affinity, BCMA receptor densities, and CAR densities were fixed to the known values (Table 1), whereas the parameters associated with the efficacy of CAR-T cells, i.e., maximum killing rate (K<sub>Kill</sub><sup>Max</sup>) and killing potency (K<sub>CART</sub><sup>50</sup>) were estimated. The estimated K<sub>CART</sub><sup>50</sup> revealed that ~ 2.24 CAR-Target complexes per tumor cells were required to achieve 50% of the maximum killing rate (K<sub>Kill</sub><sup>Max</sup>). The extent of the cytolsis percentage was not correlated with the antigen densities on six cell lines. Hence, ~ 60% interindividual variability (IVV) was estimated on the mean maximum killing rate (K<sub>Kill</sub><sup>Max</sup>) parameter. In the subsequent modeling analysis, the cell-level in vitro potency estimates (K<sub>CART</sub><sup>50</sup>) were fixed while characterizing in vivo CAR-T cell–induced tumor growth inhibition (TGI).

#### Model-based characterization of in vivo expansion and bb2121 CAR-T cell–induced TGI in preclinical xenograft mice

Figure 2b describes the simultaneous model-fitted profiles overlaid with the observed data for bb2121 induced TGI (Figure 2b1) and bb2121 CAR-T cell expansion in blood (Figure 2b2) in a RPMI-8226 tumor-bearing...
### Table 1

The list of parameters, either fixed or estimated, that were used to build the preclinical (*in vitro* and *in vivo*) and clinical PK-PD models for CAR-T cells

| Parameter name | Description (units) | Estimate (mean/RSE%) | Estimate (of RSE%) | Source |
|----------------|----------------------|----------------------|--------------------|--------|
| **DT\textsuperscript{Tumor}** | The doubling time of tumor cells (hour) | Daudi = 24 hours | — | 43 |
| | | JeKo = 26 hours | — | |
| | | K562-bcma = 47 hours | — | |
| | | NCI-H929 = 50 hours | — | |
| | | RPMI-8226 = 60 hours | — | |
| | | U266-B1 = 108 hours | — | |
| **DT\textsuperscript{CAR-T}** | The doubling time of CAR-T cells (hour) | 24h | — | 11 |
| **Ag\textsuperscript{Tumor}** | Overall density of TAA on different tumor cell lines (numbers/cell) | JeKo = 222/cell | — | 11 |
| | | Daudi = 1173/cell | — | |
| | | U266-B1 = 2930/cell | — | |
| | | NCI-H929 = 10,000/cell | — | |
| | | RPMI-8226 = 12,590/cell | — | |
| | | K562-BCMA = 76,942/cell | — | |
| **Ag\textsuperscript{CAR}** | Overall density of CARs on CAR-T cells (numbers/CAR-T cell) | 15,000/cell | — | Internal data set |
| **K\textsubscript{Max Kill}** | The first-order maximum rate of killing of tumor cells by CAR-T cells (1/hour) | 0.353 (14) | 0.62 (17) | Estimated |
| | | | | |
| **K\textsubscript{CAR-T} 50** | The number of “CAR-Target complexes per tumor cell” required to achieve 50% of the maximum killing rate (number/cell) | 2.24 (2) | — | Estimated |
| **γ\textsubscript{Tumor Growth}** | The Sigmoidicity factor associated with the killing of tumor cells (unitless) | 1.07 (0.2) | — | Estimated |
| **K\textsubscript{on}** | The binding affinity of CAR to TAA (1/Molar/second) | 7.1E4 (Fixed) | — | 44 |
| **K\textsubscript{off}** | The dissociation rate of CAR to TAA (1/second) | 2.39E-3 (Fixed) | — | 44 |
| **Tumor\textsubscript{0}** | The initial tumor cells (number) | 1E5 (Fixed) | — | 44 |

### Parameters associated with the preclinical PK-PD model

| Parameter name | Description (units) | Estimate (mean/RSE%) | Estimate (of RSE%) | Source |
|----------------|----------------------|----------------------|--------------------|--------|
| **K\textsubscript{Exp max}** | The first-order maximum rate of CAR-T cells expansion (1/day) | 0.9168 (8.47) | — | Estimated |
| **Ec\textsubscript{50}** | The number of “CAR-Target complexes per effector cell” required to achieve 50% of the maximum rate of CAR-T cell expansion (number/cell) | 1.15 (30.9) | — | Estimated |
| **Rm** | The first-order conversion rate from effector cells to memory cells (1/day) | — | — | — |
| **K\textsubscript{el}e** | The first-order elimination rate of effector CAR-T cells (1/day) | 113 (Fixed) | — | Clinical model estimates |
| **K\textsubscript{el}m** | The first-order elimination rate of memory CAR-T cells (1/day) | — | — | — |
| **K\textsubscript{Max Kill}** | The first-order maximum rate of killing of tumor cells by CAR-T cells (1/day) | 0.612 (28.2) | — | Estimated |
| **KC\textsubscript{CAR-T} 50** | The number of “CAR-Target complexes per tumor cell” required to achieve 50% of the maximum killing rate (number/cell) | 2.24 (Fixed) | — | *In vitro* estimates |
| **K\textsubscript{12}** | The first-order distribution rate from blood compartment to bone marrow compartment (1/day) | 20,304 (20.9) | 0.1 (Fixed) | Estimated |
| **K\textsubscript{21}** | The first-order redistribution rate from bone marrow compartment to blood compartment (1/day) | 0.3288 (29.5) | 0.1 (Fixed) | Estimated |
| **K\textsubscript{g Tumor}** | The first-order rate of tumor growth (1/day) | 0.0888 (10.9) | 0.1 (Fixed) | Estimated |
| **V\textsubscript{b}** | The volume of blood compartment (mL) | 0.944 (Fixed) | — | 45 |
| **V\textsubscript{tumor}** | The volume of tumor compartment (mL) | 0.151 (Fixed) | — | 11 |

(Continues)
xenograft mouse model using the PK-PD model described in Figure 1b. While performing the preclinical modeling analysis, the memory differentiation of CAR-T cells at the site of action (as described in Figure 1b) was neglected due to the lack of observed data at later timepoints. Overall, the model was able to characterize the integrated CAR-T cell PK

| Parameter name | Description (units) | Estimate (mean/RSE%) | Estimate (co/ RSE%) | Source |
|----------------|----------------------|----------------------|---------------------|--------|
| $K_{on}$       | The binding affinity of CAR to TAA (1/Molar/second) | 7.1E4 (Fixed) | — | 44 |
| $K_{off}$      | The dissociation rate of CAR to TAA (1/second) | 2.39E-3 (Fixed) | — | 44 |
| $Ag^{CAR}$     | Overall density of CARs on CAR-T cells (number/CAR-T cell) | 15,000 (Fixed) | — | Internal data set |
| $Ag^{Tumor}$   | Overall density of TAA on different tumor cell lines (number/ cell) | 12,590 (Fixed) | — | 44 |
| $\tau$         | The transit time parameter associated with signal transduction of killing signal (hour) | 47.4 (36.7) | — | Estimated |

Parameters associated with the clinical PK-PD model

| Parameter name | Description (units) | Estimate (mean/RSE%) | Estimate (co/ RSE%) | Source |
|----------------|----------------------|----------------------|---------------------|--------|
| $K_{Exp}^{max}$| The first-order maximum rate of CAR-T cells expansion (1/day) | 1.73 (10) | 0.22 (38) | Estimated |
| $EC_{50}^{Exp}$| The number of “CAR-Target complexes per effector cell” required to achieve 50% of the maximum rate of CAR-T cell expansion (number/cell) | 10 (18) | — | Estimated |
| $Rm$           | The “net” first-order conversion rate from effector cells to memory cells (1/day) | 0.00002 (66) | 0.00004 (Fixed) | Estimated |
| $Ke_{e}$       | The first-order elimination rate of effector CAR-T cells (1/day) | 113 (19) | — | Estimated |
| $Ke_{m}$       | The first-order elimination rate of memory CAR-T cells (1/day) | 0.219 (13) | — | Estimated |
| $K_{12}$       | The first-order distribution rate from blood compartment to bone marrow compartment (1/day) | 1.71(11) | — | Estimated |
| $K_{21}$       | The first-order redistribution rate from bone marrow compartment to blood compartment (1/day) | 0.176(14) | — | Estimated |
| $V_{b}$        | The volume of blood compartment (L) | 5 (Fixed) | — | 46 |
| $V_{bm}$       | The volume of bone marrow compartment (L) | 3.65 (Fixed) | — | 46 |
| $Ag^{CAR}$     | Overall density of CARs on CAR-T cells (number/CAR-T cell) | 15,000 (Fixed) | — | Internal data set |
| $Ag^{Tumor}$   | Overall density of TAA on different tumor cell lines (number/ cell) | 12,590 (Fixed) | — | 11 |
| $K_{on}$       | The binding affinity of CAR to TAA (1/M/s) | 7.1E4 (Fixed) | — | 44 |
| $K_{off}$      | The dissociation rate of CAR to TAA (1/second) | 2.39E-3 (Fixed) | — | 44 |
| $TransC$       | The conversion factor from CAR-T cells to transgene copies (unitless) | 0.002 (Fixed) | — | 46 |
| $K_{Tumor}^{Max}$ | The first-order rate of tumor growth (1/day) | 0.008 (Fixed) | — | 47 |
| $K_{Kill}^{Max}$ | The first-order maximum rate of killing of tumor cells by CAR-T cells (1/day) | 0.343 (21) | 0.50 (37) | Estimated |
| $K_{CAR-T50}^{-}$ | The number of “CAR-Target complexes per tumor cell” required to achieve 50% of the maximum killing rate (number/cell) | 2.24 (Fixed) | — | In vitro estimates |
| $K_{m}$        | The degradation rate of serum M-protein (1/day) | 0.117 (Fixed) | — | 48,49 |
| $P_{m}$        | The production rate of serum M-protein (picogram/cell/day) | 12.1 (Fixed) | — | 48,49 |
| $K_{b}$        | The degradation rate of soluble BCMA (1/day) | 0.7 (Fixed) | — | 50 |
| $P_{b}$        | The production rate of soluble BCMA (picogram/cell/day) | 0.175 (Fixed) | — | Calculated |
| $\gamma_{m}$  | The factor associated with the production of M-protein in response to the tumor change (unitless) | 0.215 (5) | 0.21 (44) | Estimated |
| $\gamma_{b}$  | The factor associated with the production of sBCMA in response to the tumor change (unitless) | 1 (Fixed) | 1.0 (31) | Estimated |

BCMA, B-cell maturation antigen; CAR, chimeric antigen receptor; NSG, NOD scid gamma; PD, pharmacodynamic; PK, pharmacokinetic; RPMI, Roswell Park Memorial Institute; sBCMA, soluble BCMA; TAA, tumor-associated antigen.
and TGI data set, where all of the known drug-specific and system-specific parameters were fixed (as listed in Table 1), and the parameters associated with CAR-T cell expansion and CAR-T cell–induced tumor killing were estimated.

The “CAR-Target complexes per tumor cell” based potency parameter (KCART50) was fixed to the value estimated from the in vitro model fitting (Figure 2a, Table 1). In addition, a brief delay was observed between the formation of the CAR-Target complexes to the initiation of TGI, which was estimated as the transit time of 7.84 days. The model-estimated CAR- T cell expansion parameters revealed that ~1.15 CAR-Target complexes per CAR-T cells were required (EC Exp50) to achieve 50% of the maximum expansion rate (KE Exp max), which was estimated to be 0.92 1/day (~18-hour half-life). Estimation of the distributional rate constants (K12, K21) between the blood and tumor compartments revealed a very rapid distribution of CAR-T cells to the peripheral tissues on systemic administration.

Model-based characterization of in vivo expansion and bb2121 CAR-T cell–induced TGI in patients with r/r multiple myeloma

Figure 3 describes the simultaneous model-fitted profiles overlaid with the mean observed data (with standard deviations) for bb2121 blood PK (Figure 3a), change in soluble BCMA levels compared with baseline (Figure 3b), and change in serum M-protein levels compared with baseline.
**Figure 2** Observed and model-fitted profiles for the preclinical data sets for anti B-cell maturation antigen (BCMA) chimeric antigen receptor (CAR) T cells (bb2121). (a) Tumor cell killing: Observed (dots) and model-generated (solid lines) curves of viability of BCMA-expressing tumor cell lines with varying antigen-densities (Ag-tumor: 222–76,942 receptors/cell) upon incubation with CAR-T cells with varying effector to target (E:T) ratios for 4 hours. (b) Tumor growth inhibition and CAR-T cell expansion: Observed (dots) and model-generated (solid lines) profiles of vehicle (blue) or 5 x 10^6 BB2121 CAR-T cells (red) induced (b1) tumor growth inhibition and (b2) blood CAR-T cell expansion in NOD scid gamma mice inoculated with RPMI-8226 tumors

(Figure 3c) at four different dose levels ranging from 50–800 million CAR-T cells/patient. The model was able to capture the overall dose-dependent multiphasic PK profile of bb2121, highlighted by the rapid distribution, expansion, contraction, and persistence phases.

It was observed that the lowest dose of 50 million CAR-T cell resulted in lower CAR-T peak blood concentration (C_{max}) (Figure 3a) and poor persistence in comparison to the rest of clinically investigated dose levels. The insert between within each plot (Figure 3a) shows the model characterization of the rapid expansion phase of bb2121, within the time frame of 12 days. The model estimated CAR-T cell expansion parameters revealed that ~ 10 CAR-Target complexes per CAR-T cells
MULTISCALE PK-PD MODELING FOR CAR-T CELL THERAPY

(EC_{Exp_{50}}) were required to achieve 50% of the maximum CAR-T cell expansion rate (K_{Exp_{max}}), which was estimated to be 1.73 1/day (~9.16-hour half-life). The estimated values for K_{max} and EC_{Exp} were < 2-fold and ~ 10-fold higher than the corresponding preclinical in vivo expansion parameter estimates. The model estimation also revealed that the blood elimination rate constant for CAR-T cells with effector phenotype (K_{e1e}) was much higher (shorter half-life) than that for the persistent memory phenotypic (K_{elm}) population. Global sensitivity analysis (Figure S1) revealed the relative sensitivities of PK parameters at different phases of blood PK, for example, the elimination parameter associated with the effector pool (K_{e1e}) was more sensitive in the expansion phase, whereas the memory differentiation parameter (K_{m}) and elimination parameter for memory pool (K_{elm}) was more sensitive in the persistent phases. Estimation of the distributional rate constants (K_{12}, K_{21}) between the blood and bone marrow compartments also revealed a very rapid distribution of CAR-T cells to peripheral tissues on systemic administration in concordance with our preclinical in vivo observations.

Figure 3b,c describes the decrease in two biomarkers, i.e., soluble BCMA (sBCMA) and serum M-protein levels, which serve as surrogates to the rate and extent of CAR-T cell–induced tumor cell depletion (Figure 1b). The extent of observed decrease in both biomarkers directly correlated with the observed blood PK of CAR-T cells, where lower C_{max} and poor persistence in the 50 million CAR-T dosing cohort (Figure 3, in blue) led to the limited decrease in mean sBCMA and serum M-protein profiles compared with baseline levels. While characterizing the data set, the production and degradation rates of the two biomarkers were obtained from literature sources as described in Table 1. The model estimated maximum rate constant of CAR-T induced tumor cell depletion rate (K_{max}^{Kill}) was estimated to be 0.343 1/day (2-day half-life), which was < 2-fold higher than the preclinical estimate (0.612 1/day), and the killing potency (K_{CART}^{50}) was fixed to the in vitro estimated value. Unlike preclinical TGI studies, the direct tumor cell depletion data set was unavailable in the clinical settings and hence no delay (transit time) was assumed between the formation of “CAR-Target complexes” and the induction of tumor cell depletion.
Impact of CAR-T dose level and patient tumor burden on clinical PK and PD end points

Figure 4 describes contour plots where the impact of patient initial tumor burden and CAR-T dose level was evaluated on $C_{\text{max}}$, time to reach maximum concentration ($T_{\text{max}}$), soluble BCMA ($s\text{BCMA}$) concentration decrease from baseline (%), and serum M-protein concentration decrease from baseline (%). Model simulations were performed after single intravenous administration of anti-BCMA (bb2121) CAR-T cells to patients with multiple myeloma at a dose range from $50$ to $800 \times 10^6$ CAR-T cells and a tumor burden from $5\%$ to $95\%$ of plasma cells.

A function of dose and tumor burden; however, this effect saturates after a threshold. The relative sensitivity of these PD biomarkers to initial tumor burden was also dependent on their relative degradation half-lives ($K_b, K_m$), where in sBCMA has a faster turnover rate in comparison to serum-M protein.

Model fittings for individual patient response data and calculation of progression-free survival (PFS)

Figure 5 describes the model-fitting results to the individual patient response data categorically scored based on percentage changes in serum-M protein measurements according to the IMWG Uniform Response Criteria for Multiple Myeloma. Figure 5a describes the individual model fits for representative subjects, whereas Figure S2 describes the model fittings for all 33 subjects. The error bars in Figures a1–45 represent the bandwidth of each categorical bin based on the response criteria. The model could simultaneously characterize individual patient responses, and the calculated PFS rates (dashed lines) were concordant with the reported values (solid lines) as shown.
While performing this modeling step, the IIV in the PD parameters was estimated (as shown in Table 1) while fixing the mean estimates for parameters from the previous step.

Virtual patient simulations and calculation of patient response rates

Figure 6a describes virtual patient simulations (250/1000) for percentage of change in serum-M protein after single-dose administration of CAR-T cells ranging from 50–800 million cells/patient. The Supporting Information describes the list of parameters and criteria for performing these Monte Carlo simulations, whereas Figure S3 describes all 1000 profiles at each dose level. The overall intensity of simulated longitudinal change in serum-M protein profiles in 1000 virtual patients describes the extent of variability in response. The simulations revealed that the 50 million dose level exhibited responses in fewer patients, with an overall higher probability of relapses. The subsequent dose level of 150 million exhibited a higher rate of response with fewer cases of relapse. The two higher dose levels (i.e., 450 and 800 million) exhibited a similar rate (c.f. 150 million), but marginally higher depth of response (Figure S3). In fact, the dose level of 450 million was determined to be most
To further elucidate the marginal differences in responses at various dose levels, categorical binning of simulated continuous profiles was performed at 6 months and 21 months post-CAR-T infusion based on the IMWG Uniform Response Criteria for Multiple Myeloma. Dose-response curves were later derived (Figure 6b) from Monte Carlo simulations, highlighting the percentage of subjects achieving progressive disease (PD), stable disease (SD), partial response (PR), and complete response (CR) compared with baseline at the end of 6 and 21 months. It was observed that the fraction of patients with PD increases by the end of 21 months in comparison to 6 months due to the reduction in CAR-T exposures at later timepoints and hence relapse in the overall tumor burden. In addition, dose-dependent (i) decrease in PD and (ii) increases in SD, PR, and CR were predicted until a threshold dose level ($150 \times 10^6$, dashed vertical line), beyond which a nearly flat dose-response curve was observed with no substantial improvement in exposure or efficacy with dose escalation.

**DISCUSSION**

The field of cancer immunotherapy has been revolutionized with the emergence of CAR-T cells. This “living” therapeutic modality possesses the potential for *in vivo* expansion, effective tumor cell killing, and long-term persistence. Although a promising new platform in oncology, the drug-specific and system-specific determinants influencing the PK and PD of these agents are poorly understood. Consequently, there are no established paradigms to facilitate preclinical to clinical translation of CAR-T cell therapies and to *a priori* predict safe and efficacious first-in-human dose levels. Moreover, due to the limited clinical experience with CAR-T cell therapy, the impact of dose escalation and baseline tumor burden on patient responses is still unclear. We believe that the utility of mechanism-based PK-PD models can be paramount in understanding the underlying dose-exposure-response relationship of CAR-T cell therapy and facilitate the discovery and development of these agents.
Here we have adopted a stepwise modeling approach that uses CAR-T cell *in vitro* and *in vivo* functional activity data sets from different drug development phases using anti-BCMA (bb2121) CAR-T cells as a case study. The first step toward this goal was to develop a cell-level PD model (Figure 1a). The model simultaneously characterized the bb2121 CAR-T cell-mediated killing of BCMA-expressing tumor cell lines (Figure 2a). Such model-based characterization of *in vitro* killing potential of CAR-T cells can help determine the cell-level potency parameter for CAR-T cells ($KC^{\text{CAR}}$), which could later be translated toward the development of *in vivo* (both preclinical and clinical) mathematical relationships. It is imperative that when designing such functional *in vitro* experiments in discovery settings, patient representative cell lines, CAR-T cells derived from multiple donors (healthy and diseased), relevant matrices (media vs. whole blood) and adequate E:T ratios should be used to characterize cytolysis, CAR-T cell expansion, and the release of relevant cytokines.

The second step involved the development of a mechanistic-based PK-PD model to characterize *in vivo* TGI and CAR-T cell expansion. A unified PK-PD model (Figure 1b) was used to adequately characterize both preclinical and clinical data sets for anti-BCMA (bb2121) CAR-T cells. Upon single infusion of ~5 million bb2121 CAR-T cells in RPMI-8226 xenograft mice, it was observed that CAR-T cells rapidly extravasate from the blood compartment to the site of action (solid tumor) and other peripheral tissues, followed by their expansion, which is later reflected in the blood pool with a delay (Figure 2b). When designing such *in vivo* xenograft studies, a range of dose levels should be explored to characterize dose dependency on TGI, blood/tumor CAR-T cell expansion, and the release of relevant cytokines. If feasible, the impact of donor-to-donor variability on *in vivo* CAR-T cell function should also be assessed.

Based on our sequential modeling analysis, there was ~10-fold difference in the expansion potency ($EC^{\text{ESP}}$) parameter estimate for bb2121 between preclinical xenograft studies and phase I clinical studies. However, mechanistic vigilance is required while directly translating this parameter estimate to predict the clinical expansion rate because it is dependent on donor-to-donor variability in the autologous CAR-T cell product, CAR-T cell fitness, and overall manufacturing process differences in the CAR-T cell product in preclinical and clinical settings. In addition, the host environment differences between immunocompromised mice and patients with t/r multiple myeloma along with the accessibility of the tumor site among two species should also be taken into consideration.

In the third step of our modeling analysis, the proposed PK-PD model (Figure 1b) was used to characterize the mean PK and two PD biomarkers (sBCMA, serum M-protein) data sets in patients with t/r multiple myeloma (Figure 3). The model was able to effectively capture the multiphasic PK profile of bb2121 CAR-T cells. The mean estimates revealed a very rapid extravasation of CAR-T cells from the blood compartment ($K_{12}$) in comparison to their circulation back to the systemic circulation ($K_{21}$). This observation was consistent with the known reports highlighting that only a fraction (~5%) of overall CAR-T cells reside in the peripheral circulation. The data analysis also revealed the differential elimination rates ($K_{el}$ vs. $K_{elm}$) of two CAR-T cell phenotypes, i.e., effector vs. memory cells (Table 1). This observation is also consistent with the expected half-lives of these two phenotypes, where effector phenotypes are known to be short lived compared with memory phenotypes. The shorter half-life of the effector pool could be explained by mechanisms such as activation-induced cell death, exhaustion, and loss of antigen stimulus as well as antibody and/or complement-driven lysis due to immunogenicity.

Although quiescent (memory) T cells (Figure 2b) could also differentiate back into short-lived, rapidly proliferating effector T cells (CAR-Te) upon CAR-Target engagement, a “net” unidirectional first-order rate constant of conversion (Rm) from short-lived effector to long-lived memory CAR-T cells was estimated. Due to the unavailability of measurements for different CAR-T cell phenotypes (e.g., naïve/stem cell memory ($T_{cem}$)), central memory ($T_{cm}$), effector memory ($T_{em}$), and effector ($T_e$)) in the infused cell product or during the duration of the study, it was assumed that 100% of the infused CAR-T cells were of the effector phenotype, which gradually differentiate ($r_m$) into the memory phenotype at the site of action (bone marrow). However, with the availability of flow-based measurements pertaining to different subsets (CD4/CD8) and phenotypes of CAR-T cells in the peripheral blood and tumor site, the developed base model could be evolved to mathematically characterize distinct kinetics, efficacy, and expansion capabilities of different CAR-T subsets/phenotypes moving forward.

The impact of CAR-T dose and initial patient disease burden on CAR-T cell expansion and percentage decrease in serum sBCMA and M-protein levels was assessed by simulations shown in Figure 4. It was observed that the patient tumor burden could be more sensitive to apparent $C_{max}$ and $T_{max}$ after expansion in comparison to the administered doses of CAR-T cells. These results were not only consistent with our prior preclinical PK-PD modeling but also a similar observation has been observed in several clinical trials, including anti-CD19, anti-BCMA, and Tmune anti-prostate specific membrane antigen (PSMA)/transforming growth factor beta dominant negative (TGFβdn) CAR-T cell clinical studies. Within the Tmune anti-PSMA/TGFβdn clinical trial in patients with metastatic castration-resistant prostate cancer, it was observed that a ~10-fold increase in CAR-T doses does not result in a proportional increase to overall extent of exposure ($C_{max}$); however, the rate to maximum exposure is increased (decrease in $T_{max}$).
Similarly, in an anti-BCMA CAR-T cells clinical study in patients with multiple myeloma, a 10-fold increase in CAR-T dose levels among different cohorts (with or without lymphodepletion) led to no significant differences in $C_{\text{max}}$. Furthermore, in anti-CD19 CAR-T clinical trials in patients with B cell acute lymphoblastic leukemia, it was also observed that patients with higher bone marrow acute myeloid leukemia blast counts led to higher expansion (increased $C_{\text{max}}$) compared with patients with lower tumor burdens. These clinical observations provide further validity of adopting a systems approach while developing translational PK-PD relationships for CAR-T cells. Some other factors that could impact the cellular kinetics and efficacy of CAR-T cells include (i) product characteristics such as CAR-T cell subsets (CD4/CD8 ratios), phenotypes ($T_{\text{scm}}$, $T_{\text{em}}$, $T_{\text{cm}}$, and $T_{\text{e}}$), and exhaustion markers (PD-1, LAG-3, TIM-3) in patient apheresate and preinfusion products; (ii) presence of immunosuppressive immune cells and soluble factors; (iii) presence or absence of lymphodepletion, and (iv) resistance mechanisms leading to antigen escape. These factors are further discussed in the Supporting Information.

Later, the validated clinical PK-PD model was leveraged to describe the reported categorical response data for individual patients ($n = 33$; Figure 5 and Figure S2) based on the decrease in serum M-protein levels. Adopting this methodology not only bolstered our confidence in identifying mean parameter values but also enabled us to estimate intersubject variability around the mean estimates. PFS rates were calculated using this approach, which were later validated using the observed PFS rates, as shown in Figure 5b.

Once validated, the population PK-PD model was leveraged to do virtual population simulations ($n = 1000$) at different dose levels (50–800 $\times 10^6$ CAR-T cells; Figure 6a), and categorical dose-response curves were generated at 6 months (when CAR-T PK is active) and 21 months (CAR-T PK is inactive) post-CAR-T cell infusion based on the IMWG Uniform Response Criteria for Multiple Myeloma. Simulated dose-response curves suggested a very steep dose-exposure-response relationship (Figure 6). This observation is in concordance with the clinical behavior of two approved autologous CAR-T cell products (i.e., Kymriah and Yescarta), where a flat dose-response relationship was observed. Our model simulations for anti-BCMA CAR-T (bb2121) suggested that beyond a “threshold” CAR-T cell dose level, dose-escalation does not result in further improvement in expansion and patient responses. Recently reported clinical observations from the pivotal phase II karMMa trial on bb2121, evaluating three different dose levels (150, 300, and 450 million CAR + T cells) in 128 patients with r/r multiple myeloma also suggested very similar PK behavior across three dose levels and marginal improvement in response rates when escalating from 150 to 450 million dose levels. Not only are these clinical observations somewhat concordant with our model predictions but also when done a priori, they could significantly influence the clinical development of future CAR-T programs moving forward. Besides robust dose-efficacy relationships, when selecting recommended phase II dose level(s) for subsequent pivotal studies, dose-toxicity relationships and manufacturing capacity/costs of CAR-T cells should also be taken into consideration.

In summary, we have developed a multiscale translational PK-PD model to effectively characterize preclinical (in vitro and in vivo) and clinical PK-PD data sets for CAR-T cell therapy. This integrated PK-PD framework to describe the cellular kinetics of CAR-T cells could be leveraged in the future to characterize the two commonly observed toxicities with CAR-T cell therapies, i.e., cytokine release syndrome and immune effector cell–associated neurological syndrome. Currently the complexity of the model structure is limited by the availability of clinical data sets. Hence many important components, such as (i) CAR-T product-related attributes, (ii) integration of multiple bioanalytical measurements, and (iii) presence of a host-immune system (and its interactions with CAR-T cells), are currently missing. However, with exponential evolution of the cell-therapy platform, this modeling and simulation framework can be extended and evolved in the future to explore multiple infusion regimens and facilitate the discovery and development of other cell-therapy programs.

**ACKNOWLEDGMENTS**

The authors acknowledge Weirong Wang (Clinical Pharmacology and Pharmacometrics, Janssen R&D), Gopi Shankar (Biologics Development Sciences), Professor Yanguang (Carter) Cao (University of North Carolina Chapel Hill), and other team members from Discovery and Translational Research at Janssen Biotherapeutics for multiple discussions on the pharmacokinetic-pharmacodynamic modeling aspects of chimeric antigen receptor T cells. The authors are indebted to the teachings from Dr. Dhaval K. Shah Laboratory at University at Buffalo on “bench to bedside translational modeling,” which has significantly impacted the presented research in this article.

**CONFLICT OF INTEREST**

All authors were employees of Johnson and Johnson during the execution and completion of this study.

**AUTHOR CONTRIBUTIONS**

A.P.S. wrote the manuscript. A.P.S., W.C., X.Z., A.Z., and D.H. designed the research. W.C., X.Z., H.M., T.J.C., and A.P.S. performed the research. W.C., X.Z., T.J.C., A.P.S., and H.M. analyzed the data. All authors contributed new reagents/analytical tools.
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SUPPORTING INFORMATION
Additional supporting information may be found online in the Supporting Information section.

How to cite this article: Singh AP, Chen W, Zheng X, et al. Bench-to-bedside translation of chimeric antigen receptor (CAR) T cells using a multiscale systems pharmacokinetic-pharmacodynamic model: A case study with anti-BCMA CAR-T. CPT Pharmacometrics Syst Pharmacol. 2021;10:362–376. https://doi.org/10.1002/psp4.12598