Evaluation of Quantitative Relationship Between Target Expression and Antibody-Drug Conjugate Exposure Inside Cancer Cells

Sharad Sharma, Zhe Li, David Bussing, and ©Dhaval K. Shah

Department of Pharmaceutical Sciences, School of Pharmacy and Pharmaceutical Sciences, The State University of New York at Buffalo, Buffalo, New York

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ABSTRACT

Antibody-drug conjugates (ADCs) employ overexpressed cell surface antigens to deliver cytotoxic payloads inside cancer cells. However, the relationship between target expression and ADC efficacy remains ambiguous. In this manuscript, we have addressed a part of this ambiguity by quantitatively investigating the effect of antigen expression levels on ADC exposure within cancer cells. Trastuzumab-valine-citrulline-monomethyl auristatin E was used as a model ADC, and four different cell lines with diverse levels of human epidermal growth factor receptor 2 (HER2) expression were used as model cells. The pharmacokinetics (PK) of total trastuzumab, released monomethyl auristatin E (MMAE), and total MMAE were measured inside the cells and in the cell culture media following incubation with two different concentrations of ADC. In addition, target expression levels, target internalization rate, and cathepsin B and multidrug resistance protein 1 (MDR1) protein concentrations were determined for each cell line. All the PK data were mathematically characterized using a cell-level systems PK model developed for ADCs. Our results demonstrate a strong quantitative relationship between antigen expression level and intracellular exposure of ADCs in cancer cells.

SIGNIFICANCE STATEMENT

In this manuscript, we have demonstrated a strong linear relationship between target expression level and antibody-drug conjugate (ADC) exposure inside cancer cells. We have also shown that this relationship can be accurately captured using the cell-level systems pharmacokinetics model developed for ADCs. Our results indirectly suggest that the lack of relationship between target expression and efficacy of ADC may stem from differences in the pharmacodynamic properties of cancer cells.

Introduction

Antibody-drug conjugates (ADCs) are novel anticancer agents that are designed to achieve wide therapeutic index by selectively delivering potent cytotoxic agents to the cancer cells. At present, there are 5 US Food and Drug Administration–approved ADCs in the clinic (Adcetris, Kadcyla, Mylotarg, Besponsa, and Polivy), and more than 80 ADCs are in clinical trials (Beck et al., 2017; Moek et al., 2017). ADCs work by binding to cancer cells via the antibody backbone and internalizing inside the cells to release the cytotoxic agent that ultimately kills the cancer cells (Kalim et al., 2017). Most ADCs target cell surface antigens that are overexpressed on cancer cells and have minimal or no expression on normal tissue cells (Bornstein, 2015). Targeting specific, overexpressed antigens is key for achieving the wide therapeutic index of ADCs (Hinrichs and Dixit, 2015).

It is widely believed that the efficacy of an ADC depends on the expression level of the target on the cancer cells, as well as the inherent potency of the cytotoxic agent. In fact, the mechanism-of-action for ADCs suggests that the higher the target expression, the better the efficacy of an ADC (Tolcher, 2016; Lambert and Morris, 2017). However, there has been no preclinical or clinical investigations that comprehensively examines this belief. There is a lack of quantitative understanding regarding the relationship between target expression and ADC efficacy. Some in vitro investigations even show that the efficacy of an ADC is not always correlated with the expression levels of the target (O’Brien et al., 2008; Barok et al., 2011; Li et al., 2013; Bornstein, 2015). Consequently, the importance of antigen expression level in selecting the patient population for ADC treatment, and implementation of a precision medicine strategy for ADCs, remains ambiguous. In this publication, we have addressed a part of this ambiguity by quantitatively investigating the relationship between antigen expression levels and the exposure of ADC inside the cancer cells.

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ABBREVIATIONS: ABC, antibody-binding capacity; ADC, antibody-drug conjugate; d8, deuterated; DAR, drug-antibody ratio; HER2, human epidermal growth factor receptor 2; IS, internal standard; LC-MS/MS, liquid chromatography–tandem mass spectrometry; MDR1, multidrug resistance protein 1; MFI, mean fluorescence intensity; MMAE, monomethyl auristatin E; QC, quality control; QSC, Quantum Simply Cellular; T-vc-MMAE, trastuzumab-vc-monomethyl auristatin E.
We have used T-vc-MMAE (trastuzumab-valine-citrulline-monomethyl auristatin E) as a tool ADC and four different cell lines with different levels of HER2 expression as in vitro models. The pharmacokinetics (PK) of total trastuzumab, released (or “free”) MMAE, and total MMAE inside the cells and in the cell culture media was analyzed following incubation of ADC with the four cell lines. We also measured ADC/antibody internalization rate and cellular cathepsin B and MDR1 protein level in each cell line. Cathepsin B is a lysosomal enzyme responsible for intracellular cleavage of the valine citrulline peptide linker of ADCs, causing release of free MMAE (Dorywalska et al., 2016), and MDR1 is a drug transporter known to efflux MMAE out of cells, leading to drug resistance against MMAE based ADCs (Chen et al., 2015). Finally, in vitro PK data were mathematically characterized using our previously published cell-level PK model for ADCs (Singh and Shah, 2017b) to establish a quantitative relationship between antigen expression levels and intracellular exposure of ADC.

Materials and Methods

Cell Culture and Reagents. HER2 expressing breast cancer cell lines SKBR-3, MDA-MB-453, MCF-7, and MDA-MB-468 were purchased from American Type Culture Collection (Manassas, VA). The cells were cultured at 37°C in a humidified incubator with 5% CO2 in the recommended media with 10% FBS. Trastuzumab (Herceptin; Genentech, San Francisco, CA) was purchased from the local pharmacy and valine-citrulline-p-aminobenzylcarbamate monomethyl auristatin E (VC-PABC-MMAE) linker-payload was purchased from DC Chemicals (Shanghai, China). Goat anti-human IgG (Fc-specific) and goat anti-human IgG-Fab antibody (Fab-specific) were purchased from Sigma-Aldrich (St. Louis, MO). Deuterated (d8) MMAE was purchased from MedChem Express (Missouri Junction, NJ).

Synthesis and Characterization of Trastuzumab-vc-MMAE. T-vc-MMAE was synthesized by random conjugation of MMAE to interchain disulfide bonds of trastuzumab via the Valine-Citrulline dipeptide linker. This method results in a heterogeneous formulation of ADC with a range of drug-antibody ratios (DARs). The detailed description of T-vc-MMAE synthesis has been previously described in Singh et al. (2016b) and Singh and Shah (2017b). The purified T-vc-MMAE ADC was analyzed for potential aggregates using size-exclusion chromatography. The abundance of different DAR species in the ADC formulation and average DAR was quantitatively determined using the hydrophobic interaction chromatography. An average DAR value of the ADC was also confirmed by UV spectroscopic analysis (Singh et al., 2016b).

Receptor Count. HER2 receptor count on cells was determined by Quantum Simply Cellular kit (Bang Laboratories, Fishers, IN) as per the manufacturer’s recommended protocol. Briefly, Quantum Simply Cellular (QSC) beads and 0.2 million cells were incubated with 100 nM Alexa Fluor-488 conjugated trastuzumab (fluorochrome to antibody ratio ~ 5, developed using Alexa Fluor antibody labeling kit, molecular probes; Thermo Fisher Scientific, Waltham, MA) in PBS on ice. After 1 hour of incubation, beads were combined, washed, and resuspended in 0.5 ml PBS. Similarly, all four cells were washed, resuspended in 0.5 ml PBS, and stored on ice. Beads and cells were analyzed using BD Accuri C6 flow cytometer (BD Biosciences, San Jose, CA). Mean fluorescence intensity (MFI) of each bead type was plotted against manufacturer provided antibody-binding capacity (ABC) values to create a standard curve. MFI of individual cell line was then used to calculate the HER2 receptor count using QuickCal v.2.3 Data Analysis program provided by the manufacturer.

Rate of Internalization. T-vc-MMAE internalization rate in each HER2 positive cancer cell line was measured using Alexa Fluor-488 labeled trastuzumab as a surrogate molecule. Internalization was measured in the form of internalization score using Imagestream flow cytometer (Amnis; MilliporeSigma, Burlington, MA). Briefly, 2 × 10⁶ cells were incubated with 100 nM Alexa Fluor-488 conjugated trastuzumab in sterile PBS on ice for 1 hour. Cells were washed with PBS and resuspended in respective cell culture media with 10% FBS.

TABLE 1

<table>
<thead>
<tr>
<th>Variable</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>MMAEMediaFree</td>
<td>Amount of MMAE in the media space</td>
</tr>
<tr>
<td>MMAEMediaBound</td>
<td>Number of molecules of unbound (free) MMAE in a single tumor cell</td>
</tr>
<tr>
<td>MMAEMCellBound</td>
<td>Number of tubulin-bound MMAE molecules in a single tumor cell</td>
</tr>
<tr>
<td>ADCMediaFree</td>
<td>Concentration of T-vc-MMAE in the media space</td>
</tr>
<tr>
<td>ADCMediaBound</td>
<td>Number of T-vc-MMAE molecules bound to HER2 receptors on a single cell</td>
</tr>
<tr>
<td>ADCendo/lyso</td>
<td>Number of T-vc-MMAE molecules internalized in endosomal/lysosomal space</td>
</tr>
<tr>
<td>DAR</td>
<td>Average number of MMAE molecules conjugated to Trastuzumab</td>
</tr>
<tr>
<td>NCell</td>
<td>Number of cells in culture flask</td>
</tr>
</tbody>
</table>
followed by incubation at 37°C for 0, 1, 4, 16, 24, or 48 hours. Cells were collected and analyzed for membrane bound and cytosolic fraction of Alexa Fluor-488 labeled trastuzumab using the erode masks that were defined from the bright field image of each cell. The extent of internalization at each time point was determined using IDEAS software internalization wizard, which calculates an internalization score based on the ratio of cytosolic intensity to total cell intensity determined using bright field image of each cell. The internalization score versus % media uptake was plotted and the half-life (using the upper quartile of pixel intensities). The half-life was calculated as the first order rate of proteases induced intracellular degradation and MMAE release, which was assumed for cathepsin B and MDR1 (Hodges et al., 2011), respectively.

Cathepsin B and MDR1 ELISA. Cellular cathepsin B and MDR1 protein expression was determined using human cathepsin B ELISA Kit (ab19584; Abcam, Cambridge, MA) and human ABCB1/MDR1/P Glycoprotein ELISA Kit (LS-F25310; LSBio, Seattle, WA), respectively. For both the kits, manufacturer’s recommended protocol was followed. Briefly, 1 million cells were lysed in 200 µl of PBS by sonication. Cell lysate was used to quantify the concentration of cathepsin B and MDR1 based on standard curve created using known standards provided with the kit. Concentrations measured in picograms per milliliter were converted into picomolar based on cellular volume of 3 pl for MDA-MB-453 and MDA-MB-468 cells, 4.18 pl for SKBR-3 cells (Kenny et al., 2007), and 8.14 pl for MCF-7 cells (Singh and Shah, 2017b). Molecular weight of 37 and 170 kDa was assumed for cathepsin B and MDR1 (Hodges et al., 2011), respectively.

**ELISA to Quantify Total Trastuzumab in Cells and Media.** Five million SKBR-3, MDA-MB-453, MCF-7, and MDA-MB-468 cells were treated with 1 and 10 nM concentration of T-vc-MMAE in a 100 mm tissue culture dish with 8 ml of cell culture media. At each time point (4, 8, and 24 hours), cell culture media was collected from the treatment dishes and stored in tubes, and the cells were trypsinized, washed, and collected separately. Cells were divided into five different fractions of 1 million each. Cells and culture media collected at each time point were individually analyzed by ELISA for intracellular and extracellular total trastuzumab concentrations, respectively. The trastuzumab ELISA is covered in detail in Singh and Shah (2017b). Briefly, an assay plate (MaxiSorp 96-well clear plate, Nunc; Thermo Fisher Scientific) was coated with Fc-specific goat anti-human IgG as a capture antibody and blocked with 1% bovine serum albumin (Pierce, Thermo Fisher Scientific) buffer. The plate was incubated with T-vc-MMAE standards, quality controls (QC), and diluted test samples, and detected by the sequential addition of goat anti-human IgG-alkaline phosphatase. The bound alkaline phosphatase activity was detected by colorimetric conversion of p-nitro phenyl phosphate solution (1 mg/ml in diethanolamine buffer) (Sigma-Aldrich) and measurement of absorbance at 405 nm. The optical density of each well was recorded using Filter Max F-5 microplate analyzer (Molecular Devices, Sunnyvale, CA), and the standard curve with four-parameter logistic model was created using SoftMax Pro software (Molecular Devices). Standards and QC samples were prepared by performing serial dilutions of T-vc-MMAE in media or 10% of cell lysate in RIPA buffer. Cell samples were lysed in RIPA buffer (Pierce, Thermo Fisher Scientific) with protease inhibitor cocktail at a concentration of 1 million cells/100 µl. Cell lysate samples were diluted 10-fold with PBS before subjecting to ELISA.

**TABLE 2.** A list of parameters and their values used to drive the in vitro PK model for T-vc-MMAE

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Description</th>
<th>Units</th>
<th>Value (CV%)</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>V&lt;sub&gt;Media&lt;/sub&gt;</td>
<td>Volume of the media compartment</td>
<td>mL</td>
<td>8</td>
<td>Fixed</td>
</tr>
<tr>
<td>SF</td>
<td>Scaling factor to convert the number of molecules to nanoMoles</td>
<td>Unitless</td>
<td>10&lt;sup&gt;6&lt;/sup&gt;</td>
<td>Fixed</td>
</tr>
<tr>
<td>V&lt;sub&gt;Cell&lt;/sub&gt;</td>
<td>Volume of each cell</td>
<td>pl</td>
<td>SKBR-3 4.18, MDA-MB-453 8.14, MCF-7 3.0, MDA-MB-468 20</td>
<td>Kenny et al., 2007; Singh and Shah, 2017b</td>
</tr>
<tr>
<td>DT&lt;sub&gt;Cell&lt;/sub&gt;</td>
<td>Doubling time of each cell line</td>
<td>h</td>
<td>SKBR-3 65, MDA-MB-453 0.018, MCF-7 0.37, MDA-MB-468 0.014</td>
<td>Shah et al., 2012</td>
</tr>
<tr>
<td>k&lt;sub&gt;on&lt;/sub&gt;</td>
<td>First order net antibody-HER2 complex internalization rate</td>
<td>1/h</td>
<td>SKBR-3 0.028, MDA-MB-453 0.11, MCF-7 0.178, MDA-MB-468 0.178</td>
<td>Fixed</td>
</tr>
<tr>
<td>k&lt;sub&gt;off&lt;/sub&gt;</td>
<td>First order non-specific deconjugation rate of MMAE from ADC</td>
<td>1/h</td>
<td>0</td>
<td>Fixed</td>
</tr>
<tr>
<td>A&lt;sub&gt;HER2&lt;/sub&gt;</td>
<td>Average drug to antibody ratio of T-vc-MMAE</td>
<td>Unitless</td>
<td>SKBR-3 4.16, MDA-MB-453 0.8 X 10&lt;sup&gt;6&lt;/sup&gt;, MCF-7 0.052 X 10&lt;sup&gt;6&lt;/sup&gt;, MDA-MB-468 0.01 X 10&lt;sup&gt;6&lt;/sup&gt;</td>
<td>Fixed</td>
</tr>
<tr>
<td>k&lt;sub&gt;ADC&lt;/sub&gt;</td>
<td>First order rate of proteases induced intracellular ADC degradation and MMAE release</td>
<td>1/h</td>
<td>SKBR-3 0.196 (28), MDA-MB-453 0.168 (2), MCF-7 0.03, MDA-MB-468 0.03</td>
<td>Estimated</td>
</tr>
<tr>
<td>k&lt;sub&gt;off&lt;/sub&gt;</td>
<td>Average first order cellular influx and efflux rate constant for free MMAE</td>
<td>1/h</td>
<td>8.33, 0.199</td>
<td>Singh and Shah, 2017b</td>
</tr>
</tbody>
</table>

*Note: CV% indicates coefficient of variation.*
LC-MS/MS to Quantify MMAE in Cells and Media. As mentioned above, cells and culture media for each cell line treated with 1 and 10 nM T-vc-MMAE were collected separately at 4, 8, and 24 hours. Cells were divided into five different fractions of 1 million cells each. Cells and culture media were individually analyzed for intracellular and extracellular concentration of free and total MMAE (after forced deconjugation), respectively, by liquid chromatography–tandem mass spectrometry (LC-MS/MS). MMAE was quantified using Sciex API 3000 triple quadrupole (SCIEX, Framingham, MA) with high-pressure liquid chromatography.

Fig. 2. HER2 receptor number quantification for different breast cancer cells by flow cytometry. (A) Histogram overlay of control (blue line) and Alexa Fluor 488–conjugated trastuzumab labeled cells (red line). The histograms show one representative result from three separate experiments. (B) Histogram overlay of QSC beads labeled with Alexa Fluor 488–conjugated trastuzumab. The histograms show one representative result from three separate experiments. (C) Antigen binding capacity vs. MFI plot generated using QuickCal v. 2.3 Data Analysis program. Three separate experiments are represented by open circles, squares, or triangles, with individual R values. (D) HER2 receptor count (mean ± S.D., n = 3) on SKBR-3, MDA-MB-453, MCF-7, and MDA-MB-468 cells.

Fig. 3. Trastuzumab internalization rate determined for different breast cancer cells. (A) Bright green color in each image represents cell membrane bound and internalized fraction of Alexa Fluor 488–conjugated trastuzumab. Each image in the panel represents one cell out of 1500 gated cells that were used for calculation at each time point. (B) The plot of internalization score vs. time used to calculate internalization half-life. Solid square, upward triangle, and downward triangle represents SKBR-3, MDA-MB-453, and MCF-7 cells, respectively. Symbol represents mean and error bars represent S.D.
liquid chromatography by Shimadzu Prominence (Shimadzu Corporation, Kyoto, Japan). An XBridge BEH Amide column (Waters Corporation, Milford, MA) was used for LC, as described in Singh and Shah (2017b). d8 MMAE (MedChem Express) was used as an internal standard (IS). For each standard, the ratio of the peak areas for MMAE to IS was plotted against the standard’s concentration. The standard curve was fitted to data points using linear regression (using Analyst 1.4.2 software; Waters Corporation) and validated by low, mid, and high QC samples.

For sample preparation, details can be found in Singh and Shah (2017b). Briefly, samples (unknowns, standards, and QCs) in either media or cell suspension (1 x 10^6 cells/200 μl) were spiked with d8 MMAE. To process cell samples, cells were permeabilized and pelleted, and the supernatant lysate was collected. All samples were evaporated under nitrogen and reconstituted in mobile phase B prior to analysis.

**Forced Deconjugation to Quantify Total MMAE.** Samples containing T-vc-MMAE were incubated with papain (Sigma-Aldrich), a cysteine protease, to enzymatically cleave the conjugated MMAE from the ADC construct such that total MMAE in the sample is present in the free, unconjugated form. Prior to papain treatment, cell suspensions were sonicated to release intracellular ADC and free MMAE. Samples of both media and cell lysate were incubated overnight at 40°C with 2 mg/ml papain (Li et al., 2016). Finally, the samples were prepared for LC-MS/MS analysis as described above.

**In-Vitro PK Model for ADC.** The cell-level PK model (Fig. 1) has been described in depth in previous work (Singh and Shah, 2017b). In short, the model accounts for binding of ADC to HER2, leading to its internalization and lysosomal degradation. This leads to free MMAE release inside the cells, which can bind with tubulin or efflux out of cells into the media space. The equations associated with this model are listed below, and detailed information about all symbols, state variables, and model parameters is provided in Tables 1 and 2. Modeling was conducted using ADAPT 5 (D’Argeo et al., 2009).

\[
\frac{d(A DA DG \bar{M} M E \text{ Media} \text{ Bound})}{dt} = \left( -K_{AC A}^{A DA DG \bar{M} M E \text{ Media} \text{ Bound} \rightarrow A DA DG \bar{M} M E \text{ Media} \text{ Free} + A DA DG \bar{M} M E \text{ Media} \text{ Bound} \text{ Bound} \right) + K_{AC A}^{A DA DG \bar{M} M E \text{ Media} \text{ Bound} + N(t)} \cdot \bar{C} \text{ MMAE} \text{ Media} \text{ Bound}
\]

\[
\begin{align*}
SF \frac{d(M M A E \text{ Media} \text{ Bound})}{dt} & = K_{AC A}^{M M A E \text{ Media} \text{ Bound} \rightarrow N(t)} \cdot N(t) \cdot \bar{C} \text{ MMAE} \text{ Media} \text{ Bound} + K_{AC A}^{M M A E \text{ Media} \text{ Bound} \rightarrow M M A E \text{ Media} \text{ Bound} \text{ Bound}} \\
N(t) \cdot \bar{C} \text{ MMAE} \text{ Media} \text{ Bound} & = K_{AC A}^{M M A E \text{ Media} \text{ Bound} \rightarrow M M A E \text{ Media} \text{ Bound} \text{ Bound}} \cdot V_{\text{ Media}} \\
\end{align*}
\]

**Results**

**Receptor Count.** HER2 receptor count on breast cancer cells was determined using QSC anti-Human IgG quantitation beads by flow cytometry. Figure 2 represents the flow cytometry histograms of SKBR-3, MDA-MB-453, MCF-7, and MDA-MB-468 cells labeled with Alexa Fluor 488-conjugated trastuzumab (Fig. 2A) and of QSC bead standards of known ABC value (Fig. 2B). This resulting relationship between ABC and MFI (Fig. 2C) allows for quantification of HER2 receptor count on the four cell lines examined (Fig. 2D). HER2 receptor count per cell (mean ± S.D.) quantified for SKBR-3 cells, MDA-MB-453 cells, MCF-7 cells, and MDA-MB-468 cells are 822,558 ± 163,770, 251,407 ± 18,763, 52,069 ± 4821, and 11,424 ± 1810, respectively.

**Internalization.** Cancer cell exposure of ADC depends on internalization of ADC-receptor complex inside the cells. Alexa Fluor 488-conjugated trastuzumab was used as a surrogate to analyze the rate of internalization of T-vc-MMAE in breast cancer cells. Figure 3A shows the increase in intracellular versus membrane bound fraction of Alexa Fluor 488-conjugated trastuzumab in breast cancer cells over 48 hours. Based on internalization score versus time plot shown in Fig. 2B, the internalization half-life of Alexa Fluor 488–conjugated trastuzumab in SKBR-3, MDA-MB-453, and MCF-7 cells was found to be 24.36 ± 6.18, 6.02 ± 1.60, and 3.89 ± 0.53 hours, respectively.
Cathepsin B and MDR1 Quantification. As shown in Fig. 4A, the cellular cathepsin B protein expression was comparable among three out of four breast cancer cell lines. MDA-MB-468 cells had ~2- to 3-fold higher concentration cathepsin B compared with the other three breast cancer cells. The mean cathepsin B concentrations in SKBR-3, MDA-MB-453, MCF-7, and MDA-MB-468 cells were 15.5 ± 1.5, 18.7 ± 3.5, 12.7 ± 1.2, and 36.8 ± 10.3 nM, respectively. As shown in Fig. 2B, the cellular MDR1 protein expression was also comparable among the four breast cancer cell lines. The mean MDR1 concentrations in SKBR-3, MDA-MB-453, MCF-7, and MDA-MB-468 cells were 3.6 ± 0.5, 4.1 ± 0.1, 4.3 ± 0.4, and 2.2 ± 0.4 nM, respectively.

Intracellular Disposition of T-vc-MMAE. To analyze the quantitative relationship between target expression and ADC exposure in cancer cells, HER2 positive breast cancer cells were treated with 1 and 10 nM concentrations of T-vc-MMAE for 24 hours. Figure 5, A–H shows the time-dependent change in intracellular concentration of free MMAE, total MMAE, and total trastuzumab in SKBR-3, MDA-MB-453, MCF-7, and MDA-MB-468 cells. The intracellular concentrations of all three analytes increased with HER2 receptor count on cancer cells (Fig. 6A). There was no noticeable change observed for intracellular concentrations of total trastuzumab and total MMAE between 4 and 24 hours (Fig. 5, I–P).

Extracellular Disposition of T-vc-MMAE. Figure 5, I–P shows the time-dependent change in extracellular concentration of free MMAE, total MMAE, and total trastuzumab, after incubation of breast cancer cells with 1 or 10 nM concentration of T-vc-MMAE in the cell culture media. The extracellular free MMAE concentrations increased with HER2 expression level of the breast cancer cells, and the highest and lowest concentrations were observed for SKBR-3 and MDA-MB-468 cells, respectively (Fig. 5, I–P; Supplemental Fig. 2). The concentration of extracellular free MMAE also increased with time (Supplemental Fig. 2), and the exposure of extracellular free MMAE (AUC₀⁻²⁴) in the cell culture media was linearly correlated with HER2 receptor count on cancer cells (Fig. 6B). As shown in Fig. 5, K–O and...
Supplemental Fig. 3, medium and low level HER2 expressing cancer cell lines did not show any noticeable change in the total trastuzumab concentration in cell culture media over 24 hours. However, total trastuzumab concentration in cell culture media decreased by ~10-fold when 1 nM T-vc-MMAE was incubated with SKBR-3 cells (Fig. 5I). Total MMAE concentrations in the cell culture media followed the same trends as total trastuzumab concentrations.

Data Characterization Using Cellular PK Model. All 44 PK profiles representing the intracellular and extracellular PK of all three analytes at two doses in four cell lines, with the exception of intracellular PK of total trastuzumab for the 1 nM dose, were fitted simultaneously using the single-cell PK model for ADCs shown in Fig. 1. Figure 7 shows observed PK data superimposed over the model fittings. Most of the parameters of the PK model were fixed a priori based on experimentally measured or literature reported values. Only the rates of intracellular degradation of T-vc-MMAE for two cell lines (SKBR-3 and MDA-MB-453) were estimated using the data. As shown in Fig. 7, the model was able to capture the disposition of all three analytes in media and cellular space reasonably well. The goodness of fit was evaluated with respective $R^2$ values, after treatment with 1 (open square) and 10 nM (solid square) concentrations of T-vc-MMAE.

Fig. 6. Free MMAE exposure (AUC$_{0–24h}$) vs. HER2 receptor count plots. Intracellular (A) and extracellular (B) free MMAE exposure (mean ± S.E.M.) vs. HER2 receptor count (mean ± S.E.M.) plot, with respective $R^2$ values, after treatment with 1 (open square) and 10 nM (solid square) concentrations of T-vc-MMAE.

To facilitate the development of a quantitative relationship, HER2 receptor count was measured on each cell line. The HER2 receptor count was measured on each cell line. The HER2 receptor count was measured on each cell line.

Discussion

At present there are five Food and Drug Administration–approved ADCs (Beck et al., 2017; Lamb, 2017) and >80 ADCs under clinical development against 59 unique targets (Moek et al., 2017). The key to selecting a target antigen is overexpression of the target on the tumor cell surface with minimal or negligible expression in healthy tissue. Thus, it is imperative to understand the relationship between target expression level and ADC exposure inside cancer cells. In this study, we have evaluated a quantitative relationship between HER2 receptor expression and T-vc-MMAE exposure in breast cancer cells. In addition, for the first time we have reported a strong linear relationship between antigen expression level and intracellular (and extracellular) exposure of released drug inside the cancer cells.

Target abundance is crucial for the selection of an ADC, and it is important to identify the patients who can benefit the most from the treatment, to harness fully the therapeutic potential of an ADC (Lambert and Morris, 2017). The general hypothesis is that patients expressing a higher level of the target are more responsive than low or nonexpressers (Tolcher, 2016). But the literature reports also suggest that target expression may not always predict response to an ADC (Barok et al., 2011; Polson et al., 2011). A study with anti-CD22 ADC against non-Hodgkin’s lymphoma cells showed that surface expression of CD22 and sensitivity to the free drug may affect the ADC response in vitro. However, neither one was shown as a predictor of response, as CD22 expression and efficacy showed a poor correlation (Li et al., 2013). Certain studies also suggest that a threshold level of target antigen may be required for an ADC to be effective, and this threshold can vary among antigens based on their internalization rate and efficiency (Dornan et al., 2009; Polson et al., 2011; Sadekar et al., 2015). We hypothesize that the observed inconsistencies in the literature may arise from studies that explore the relationship between target expression and ADC efficacy directly but fail to consider the cellular exposure of the cytotoxic payload or the inherent sensitivity of cancer cells to the payload (i.e., the pharmacokinetics and pharmacodynamics of the cytotoxin).

So far, only few studies have explored the relationship between target expression and ADC exposure within tumor cells. For example, Okeley et al. (2010) have used brentuximab-vedotin and two different CD30 expressing cell lines to show that cells with higher CD30 expression achieved higher exposure of MMAE. Although such studies support qualitative target-dependent payload delivery in cancer cells, they lack quantitative evaluation of the target expression versus ADC exposure relationship. A previous study, Singh and Shah (2017b), began to explore this relationship but only examined the correlation in a high and a non–low expressing cell line under limited or continuous exposure to ADC. To expand on this earlier investigation, here we conducted a dedicated in vitro ADC disposition study using T-vc-MMAE as a tool ADC at high and low doses and four breast cancer cells with a broad range of HER2 expression: SKBR-3 (high), MDA-MB-453 (medium), MCF-7 (low), and MDA-MB-468 (very low) to explore a direct quantitative relationship between receptor number and ADC exposure in cancer cells. Furthermore, the present study demonstrates the validity of cell-level systems PK model for ADCs first proposed in Singh and Shah (2017b).

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ranged from ~800,000 (SKBR-3) to ~10,000 (MDA-MB-468) per cell (Fig. 2D), offering a wide range of target expression to evaluate ADC exposure. In addition, cell-specific internalization rates of HER2 were determined to understand how ADC internalization differs between cell lines. The internalization half-life of trastuzumab ranged from 4 to 24 hours (Fig. 3B), and there was an inverse relationship between HER2 expression level and internalization rate, which was similar to that observed by Ram et al. (2014). Of note, the rate of internalization could not be determined for MDA-MB-468 cells due to negligible expression of HER2. The measured internalization half-lives were used to calculate the rate of internalization of T-vc-MMAE, which was implemented as a parameter for the single-cell PK model (Table 2).

In vitro PK data (Fig. 5, A–H) revealed that intracellular concentrations of all three analytes increased with increasing expression of HER2. The high HER2 expressing SKBR-3 cells showed the highest concentrations of all three analytes followed by the medium and low HER2 expressing MDA-MB-453 and MCF-7 cells, respectively. As shown in Fig. 6A, a strong linear relationship was observed between HER2 receptor count and free MMAE exposure in the cells. Interestingly, the concentration of intracellular free MMAE generated was similar between the 1 and 10 nM T-vc-MMAE doses (Supplemental Fig. 4), and the overall intracellular free MMAE exposure (AUC0–24 h) increased less than twofold even for the highest HER2 expressing cells (Fig. 6A). This indicates that availability of HER2 is a rate limiting step

Fig. 7. Observed and model predicted concentration vs. time plots for intracellular and extracellular free MMAE, total MMAE, and total mAb concentrations. SKBR-3 (A), MDA-MB-453 (B), MCF-7 (C), and MDA-MB-468 (D) cells. All the data were simultaneously fitted using the single cell PK model for ADCs shown in Fig. 1.
for achieving intracellular exposure of ADC at both the concentrations tested. This observation is consistent with simulations conducted by Sadekar et al. (2015) and suggests that although higher target expression increases the exposure of an ADC, increasing the concentration of an ADC beyond target saturation does not increase intracellular payload exposure. Contrary to this observation, the concentrations of free MMAE in MDA-MB-468 cells increased by ~10-fold when media concentration of ADC increased from 1 to 10 nM (Supplemental Fig. 4). This observation suggests that for these very low targets expressing cells, the exposure of ADC was largely dependent on target independent uptake processes (e.g., pinocytosis) (Ait-Oudhia et al., 2017; Kalim et al., 2017).

Because the MMAE molecules generated inside the cells can also diffuse out of the cells, a similar trend was observed in media, where extracellular free MMAE concentrations increased with increasing expression of HER2 (Supplemental Fig. 2). A linear relationship was found between HER2 receptor count and extracellular free MMAE exposure (Fig. 6B). However, the extracellular free MMAE exposure was ~150–650-fold lower than intracellular exposure. The extracellular concentration of total trastuzumab and total MMAE showed no apparent change for very low to medium HER2 expressing cells (Fig. 5, K–O). However, for SKBR-3 cells, total trastuzumab concentration in media decreased by ~10-fold after treatment with 1 nM ADC dose, representing target mediated elimination of ADC from media (Supplemental Fig. 3A).

To further evaluate the effect of other mechanistic components on cellular disposition of ADC, we measured Cathepsin-B and MDR1 protein concentrations in the cancer cells. Cathepsin B is considered to be a critical lysosomal protease responsible for the cleavage of VC peptide linker of ADCs (Dorywal ska et al., 2016). However, Caculitan et al. (2017) have shown that cathepsin B is not the only protease causing cleavage of VC linker to release free MMAE. Similarly, MDR1 is a drug transporter known to efflux MMAE out of cells and it can affect the cellular PK of free MMAE (Chen et al., 2015). Cathepsin B concentrations in tested cell lines ranged between ~13 and 37 nM (Fig. 4A), and MDR1 expression ranged between ~2 and 4 nM (Fig. 3B). Concentrations of both proteins were within 2–3-fold, and there was no relationship observed between cellular PK of ADC and concentrations of these proteins. Therefore, our observations suggest that either the range of expression for both proteins was not large enough among all the cell lines, or neither protein significantly contributed to the cellular PK of the ADC.

Finally, all the PK data were simultaneously characterized using the model shown in Fig. 1 to quantitatively validate the observed relationship between receptor expression and ADC exposure. This process also helps in validation of experimentally determined values of biomarkers and helps evaluate the ability of the cell-level systems PK model to predict cellular PK of ADC in diverse cell lines (Singh and Shah, 2017b). The values of most system parameters, such as DT Cell, Tubulinint, Kint, Tuboff, Koff, KADC, KADCC, KADC, KADC, and KADC, were taken directly from literature reports or our previous studies (Table 2). Several other parameters, such as DTCell, KADCC, KADCC, DARC, and AgCell, HER2, were directly measured in the current study. As shown in Fig. 7, the model was able to simultaneously capture all 44 PK profiles reasonably well while fitting only two parameters (ADC degradation rates in MDA-MB-453 and SKBR-3 cell lines). The estimated ADC degradation rates in the high/intermediate HER2 expression cell lines were significantly faster than low HER2 expressing cell lines, suggesting they may have altered endo-lysosomal processing. Although the model did a reasonable job, in certain instances there was a systemic bias between observed and model predicted profiles, such as antibody PK in SKBR-3 cells and free MMAE PK in MDA-MB-468 media. These biases may stem from experimental errors, model mis specification, errors in parameter values, or unverified model assumptions. For example, contamination by a fraction of cell membrane bound ADC/antibody in the lysate of high HER2 expressing SKBR-3 cell can result in higher value of observed antibody concentrations compared with model predictions, which only considers intracellular molecules. In addition, higher levels of cathepsin B in MDA-MB-468 cells may result in higher release of MMAE from the ADC into this media, which is not accounted for by the model. As such, more dedicated investigations may be able to reveal the reasons behind each model deviation. Nonetheless, considering the mechanistic nature of the cell-level PK model and its performance in such a low degree of freedom (only two estimated parameters), this model holds promise for inclusion in an in vivo systems PK/PD model (Shah et al., 2012; Singh et al., 2016a; Singh and Shah, 2017a). Toward this end, we are in the process of conducting in vivo disposition studies with xenografts expressing different levels of tumor antigen. Results from these studies will help us better understand tumor disposition of ADCs in a more natural setting where tumor physiology may impact the “antigen expression versus tumor exposure” relationship observed in vitro. Together, the in vitro/in vivo correlation will aid in the discovery, development, and preclinical-to-clinical translation of novel ADCs.

In sum, we have demonstrated a strong quantitative relationship between antigen expression level and cellular PK of ADC, and our data indirectly suggest that differences in cancer cell PD may be the reason for the ambiguous relationship between target expression and ADC efficacy.

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Contributed in developing analytical techniques: Sharma.
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Wrote or contributed to the writing of the manuscript: Sharma, Li, Bussing, Shah.

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Address correspondence to: Dhaval K. Shah, Department of Pharmaceutical Sciences, 455 Pharmacy Bldg., School of Pharmacy and Pharmaceutical Sciences, University at Buffalo, The State University of New York, Buffalo, NY 14214-8033. E-mail: dshah4@buffalo.edu