Clinical pharmacology of vc-MMAE antibody–drug conjugates in cancer patients: learning from eight first-in-human Phase 1 studies

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ABSTRACT

vc-MMAE antibody–drug conjugates (ADCs) consist of a monoclonal antibody (mAb) covalently bound with a potent anti-mitotic toxin (MMAE) through a protease-labile valine-citrulline (vc) linker. The objective of this study was to characterize the pharmacokinetics (PK) and explore exposure–response relationships of eight vc-MMAE ADCs, against different targets and for diverse tumor indications, using data from eight first-in-human Phase 1 studies. PK parameters of the three analytes, namely antibody-conjugated MMAE (acMMAE), total antibody, and unconjugated MMAE, were estimated using non-compartmental approaches and compared across the eight vc-MMAE ADCs. Relationships between analytes were assessed by linear regression. Exposure–response relationships were explored with key efficacy (objective response rate) and safety (Grade 2+ peripheral neuropathy) endpoints. PK profiles of acMMAE, total antibody and unconjugated MMAE following the first dose of 2.4 mg/kg were comparable across the eight ADCs; the exposure differences between molecules were small relative to the inter-subject variability. acMMAE exposure was strongly correlated with total antibody exposure for all the eight ADCs, but such correlation was less evident between acMMAE and unconjugated MMAE exposure. For multiple ADCs evaluated, efficacy and safety endpoints appeared to correlate well with acMMAE exposure, but not with unconjugated MMAE over the doses tested. PK of vc-MMAE ADCs was well characterized and demonstrated remarkable similarity at 2.4 mg/kg across the eight ADCs. Results from analyte correlation and exposure–response relationship analyses suggest that measurement of acMMAE analyte alone might be adequate for vc-MMAE ADCs to support the clinical pharmacology strategy used during late-stage clinical development.

Introduction

Antibody–drug conjugates (ADCs) are a novel class of therapeutic agents consisting of a monoclonal antibody (mAb) covalently bound with a cytotoxic drug through a chemical linker. The mAb preferentially targets cell surface antigens overexpressed in tumor cells. Upon binding, the ADC is internalized by the tumor cell, where it undergoes lysosomal degradation, leading to the release of the cytotoxic drug. The use of targeted delivery of highly potent cytotoxic drugs is designed to enhance the antitumor activity while minimizing the toxicity in normal tissues. Additionally, this therapeutic approach takes advantage of the favorable pharmacokinetic (PK) properties of the antibody to provide a sustained delivery of the cytotoxic drug.

Many ADCs in clinical development use a protease-labile dipeptide linker (valine-citrulline [vc]), conjugated to monomethyl auristatin E (MMAE) via solvent-accessible thiolis present in mAb cysteines (vc-MMAE ADO).1 A representative structure of a vc-MMAE ADC is shown in Figure 1.2 Conjugation through reduced inter-chain disulfide cysteine residues results in a heterogeneous mixture of conjugated antibodies, with even-numbered drug-to-antibody ratio (DAR) species of 0, 2, 4, 6 or 8. The average DARs for most of vc-MMAE ADCs in clinical development are approximately 3–4. In 2011, the United States Food and Drug Administration approved a vc-MMAE ADC, brentuximab vedotin (ADCETRIS™) for the treatment of relapsed anaplastic large cell lymphoma and Hodgkin’s lymphoma.3

Given the complex structure of ADCs, which contain both large and small molecule components, and the mixture of various DAR species in the dosing solution, the PK of ADCs are expected to be complex. ADCs are expected to be cleared through two major pathways: proteolytic degradation and deconjugation.4,5 Similar to mAbs, ADC clearance through proteolytic degradation is driven primarily by catabolism mediated by target-specific or nonspecific cellular uptake followed by lysosomal degradation. In contrast, deconjugation clearance is usually mediated by enzymatic or chemical cleavage (e.g., maleimide exchange) of the linker leading to the release of the cytotoxic drug from the ADC.6 It is expected that ADC catabolism and deconjugation in vivo change the concentration and relative fractions of individual DAR species with time, by converting high DAR species to low DAR
species, resulting in a gradual decrease in average DAR over time. This phenomenon was observed in a preclinical study as well.\textsuperscript{7} Considering the heterogeneity and complex changes in ADC concentration and composition after ADC administration, multiple analytes have been measured in order to characterize the PK properties of an ADC.\textsuperscript{8,9} These measurements include: conjugate (evaluated as either conjugated-antibody or antibody-conjugated drug); total antibody (sum of conjugated, partially deconjugated, and fully deconjugated antibody); naked antibody (antibody without the conjugated drug); and the unconjugated drug.\textsuperscript{8–10} For the vc-MMAE ADCs clinically developed at Genentech, three analytes are routinely measured for PK assessment: antibody-conjugated MMAE (acMMAE), total antibody, and unconjugated MMAE.\textsuperscript{9}

Over the past decade, Genentech has developed eight vc-MMAE ADCs, against different targets and for diverse tumor indications, which provides a unique opportunity to characterize the clinical pharmacology for this class of ADCs. Here, we report and compare the PK characteristics of the eight vc-MMAE ADCs using data from eight first-in-human (FIH) Phase 1 studies in cancer patients. The correlation between analytes was assessed across the eight ADCs. Additionally, exposure–response relationship analysis was conducted to explore the key analyte that correlates with efficacy and safety endpoints.

**Results**

**vc-MMAE ADCs and PK data**

The drug targets, tumor indications, number of patients, and dosing for the eight vc-MMAE ADCs are summarized in Table 1. In addition to sharing the same linker-drug and conjugation chemistry, the average DARs for the eight vc-MMAE ADCs were approximately the same (i.e., ~3.5). Additionally, the mAb components of the eight ADCs all used humanized IgG1 isotype. Of the eight vc-MMAE ADCs, three molecules, namely DMOT4039A, DMUC5754A, and DFRF4539A, were able to bind circulating soluble antigens besides the corresponding member-bound antigens. Three of the eight molecules were developed for hematological tumors, while the remaining five were for solid tumors. The PK analyses were based on data from FIH Phase 1 studies in cancer patients that included both a dose-escalation component and at least one expansion cohort at a dose identified for further clinical study. The dose-escalation schemes were slightly different across the eight ADCs, but the dose levels assessed were all within the range of 0.1 to 3.2 mg/kg when given every 3 weeks (q3w) (Table 1). Nevertheless, the same dose regimen of 2.4 mg/kg q3w for the expansion cohort(s) was selected for all eight ADCs based on the overall benefit and risk profiles for each molecule. All data available in the clinical database at the time of data extraction were included in the analysis. A total of 564 patients (ranging from 33 to 95 patients for each study) across eight FIH Phase 1 studies were included in the analysis. Around half of the patients (~53%) were given the 2.4 mg/kg q3w dose. For some ADCs, single-agent weekly dosing was also pursued, but the results are not included in this analysis because the data set is small.

**PK characteristics of vc-MMAE ADCs**

As shown in Figure 2 and Figure S1, the concentration–time profiles of acMMAE, total antibody and unconjugated MMAE following the first 2.4 mg/kg intravenous dose of vc-MMAE ADCs (the dose used in the expansion cohort) were largely comparable across the eight molecules. Both acMMAE and
Figure 2. (Table 2) shows that plasma concentrations of unconjugated MMAE were low after Cycle 1 across the eight ADCs ranged from 103 L/kg, which was approximately the same as that of total antibody across eight ADCs, respectively. The differences in $C_{\text{max}}$ for acMMAE and total antibody, which was numerically shorter than that of total antibody (4.62 vs 3.92 d), although the differences between the two analytes were within the assay variability of 20% (Table 2). The mean steady-state volume of distribution ($V_{\text{ss}}$) of acMMAE for the eight ADCs ranged from 60.1 to 103 L/kg, which was approximately the same as that of total antibody (61.5–128 L/kg) (Table 2). The inter-subject variability of the exposure for the two analytes was similar across the eight ADCs with %CV for area under the concentration–time curve from time zero to infinity ($AUC_{\text{inf}}$) at Cycle 1 ranging from 20–36% and 25–46% for acMMAE and total antibody, respectively (Table 2). Consistent with concentration–time profiles, following the first dose of 2.4 mg/kg, the key PK parameters of acMMAE and total antibody (e.g., $C_{\text{max}}$, $AUC_{\text{inf}}$, Clearance, $V_{\text{ss}}$, and $t_{1/2}$) were overlapping and largely comparable across the eight ADCs (Table 2 and Figure 3). The inter-molecule variability of mean exposure (e.g., $C_{\text{max}}$ and $AUC_{\text{inf}}$ at Cycle 1) across the eight ADCs ranged from 7–19% and 16–34% for acMMAE and total antibody, which was smaller than or at the lower range of inter-subject variabilities of the two analytes (Figure 3). Modest accumulation of acMMAE and total antibody analytes were observed for the eight vc-MMAE ADCs upon repeated dosing of 2.4 mg/kg q3w, with trough concentration ratios of 1.1–1.7 and 1.0–2.1 between Cycle 3 and Cycle 1 for acMMAE and total antibody analytes, respectively.

Plasma concentrations of unconjugated MMAE were low relative to those of acMMAE and total antibody following single dose of 2.4 mg/kg (Figure 2 and Figure S1). Mean $C_{\text{max}}$ of unconjugated MMAE was 3.15–7.01 ng/mL across the eight vc-MMAE ADCs, which were more than 100-fold lower than that of acMMAE (Table 2). There was an apparent delay for unconjugated MMAE to reach its $C_{\text{max}}$ with median time to $C_{\text{max}}$ ($T_{\text{max}}$) of approximately 2–3 d post-infusion. The mean terminal half-lives of unconjugated MMAE were 2.4 mg/kg, the mean clearance of acMMAE was numerically higher than that of total antibody analyte for all the eight molecules, with mean value of 16.6–30.2 mL/dy/kg and 8.24–23.6 mL/d/kg across the eight ADCs for acMMAE and total antibody, respectively (Table 2). The mean terminal half-lives ($t_{1/2}$) for the eight ADCs ranged from 3.8 to 6.2 d for acMMAE, which was numerically shorter than that of total antibody (3.9–10.7 d) for the majority of the vc-MMAE ADCs except DEDN6526A. The mean terminal half-life of acMMAE for DEDN6526A was numerically longer than that of total antibody (4.62 vs 3.92 d), although the differences between the two analytes were within the assay variability of 20% (Table 2).
Table 2. PK parameters (mean ± SD [%CV]) of acMMAE, total antibody and unconjugated MMAE post the first dose of vc-MMAE ADCs at 2.4 mg/kg (90 min intravenous infusion).

| ADC Notation | ADC Names | N  | C_{max} (ng/mL) | AUC_{inf} (d*ng/mL) | CL (mL/d/kg) | V_{dss} (mL/kg) | t_{1/2} (d) |
|--------------|-----------|----|----------------|---------------------|--------------|----------------|-------------|
| (a) acMMAE analyte | | | | | | | |
| ADC1 | Pinatuzumab vedotin | 39 | 878 ± 240 [27.4] | 2340 ± 805 [34.4] | 21.8 ± 13.3 [61.1] | 103 ± 33.1 [32.3] | 6.18 ± 2.69 [43.5] |
| ADC2 | Polatuzumab vedotin | 45 | 887 ± 230 [25.9] | 2610 ± 860 [33.0] | 19.4 ± 10.9 [56.2] | 91.7 ± 30.3 [33.0] | 5.85 ± 2.46 [42.1] |
| ADC3 | DEDN6526A | 31 | 904 ± 191 [21.2] | 2080 ± 743 [35.8] | 22.6 ± 7.24 [32.0] | 60.1 ± 19.9 [33.0] | 4.62 ± 1.78 [38.5] |
| ADC4 | DMOT4039A | 36 | 725 ± 178 [24.5] | 1440 ± 332 [23.1] | 30.2 ± 7.36 [24.4] | 96.0 ± 29.3 [30.5] | 3.80 ± 1.15 [30.4] |
| ADC5 | DMUC5754A | 39 | 828 ± 201 [24.3] | 1800 ± 501 [27.8] | 25.6 ± 7.69 [30.0] | 85.3 ± 24.7 [28.9] | 5.12 ± 1.56 [30.5] |
| ADC6 | DNIB0600A | 51 | 923 ± 187 [20.3] | 2300 ± 541 [23.5] | 19.4 ± 4.71 [24.3] | 92.3 ± 30.7 [33.2] | 5.45 ± 1.50 [27.4] |
| ADC7 | DSTP3086S | 37 | 884 ± 172 [19.5] | 2430 ± 489 [20.1] | 17.4 ± 3.95 [22.7] | 75.4 ± 21.5 [28.5] | 5.84 ± 2.24 [38.4] |
| ADC8 | DFRF4539A | 16 | 885 ± 213 [24.1] | 2710 ± 872 [32.2] | 16.6 ± 7.69 [46.5] | 77.4 ± 41.2 [53.2] | 4.88 ± 2.48 [50.9] |
| (b) total antibody | | | | | | | |
| ADC1 | Pinatuzumab vedotin | 38 | 42.3 ± 13.1 [30.9] | 255 ± 105 [41.1] | 13.7 ± 4.5 [30.9] | 128 ± 40.5 [31.5] | 10.7 ± 6.00 [56.1] |
| ADC2 | Polatuzumab vedotin | 45 | 37.6 ± 11.0 [29.2] | 208 ± 44.7 [35.0] | 17.1 ± 6.5 [34.8] | 110 ± 34.5 [31.4] | 7.26 ± 3.75 [51.6] |
| ADC3 | DEDN6526A | 31 | 45.4 ± 10.9 [24.1] | 128 ± 44.7 [35.0] | 20.9 ± 7.18 [34.3] | 70.6 ± 25.2 [35.7] | 3.92 ± 1.68 [42.8] |
| ADC4 | DMOT4039A | 36 | 37.3 ± 9.21 [24.7] | 120 ± 40.6 [34.0] | 23.6 ± 13.8 [58.3] | 118 ± 34.5 [31.4] | 4.26 ± 1.45 [34.0] |
| ADC5 | DMUC5754A | 39 | 41.3 ± 8.15 [19.7] | 175 ± 61.1 [35.0] | 15.6 ± 6.47 [41.4] | 93.1 ± 28.5 [30.6] | 6.33 ± 2.85 [45.0] |
| ADC6 | DNIB0600A | 52 | 49.3 ± 10.4 [21.1] | 200 ± 50.9 [25.4] | 12.7 ± 3.24 [25.5] | 98.1 ± 42.0 [42.8] | 6.55 ± 2.76 [42.1] |
| ADC7 | DSTP3086S | 37 | 56.1 ± 13.6 [24.2] | 315 ± 80.7 [25.6] | 8.24 ± 2.66 [32.3] | 63.2 ± 19.7 [31.2] | 7.52 ± 3.18 [42.3] |
| ADC8 | DFRF4539A | 16 | 55.9 ± 17.3 [30.9] | 301 ± 107 [35.6] | 9.02 ± 4.73 [52.4] | 61.5 ± 23.8 [53.3] | 6.00 ± 2.98 [49.9] |
| (c) Unconjugated MMAE | | | | | | | |
| ADC1 | Pinatuzumab vedotin | 39 | 6.47 ± 4.81 [74.3] | 3.09 (0.0660, 6.96) | 51.2 ± 32.4 [63.3] | 3.64 ± 0.879 [24.1] |
| ADC2 | Polatuzumab vedotin | 44 | 6.66 ± 4.63 [69.6] | 2.95 (0.198, 9.75) | 53.6 ± 28.0 [52.3] | 3.75 ± 0.973 [26.0] |
| ADC3 | DEDN6526A | 32 | 6.72 ± 4.26 [63.4] | 2.07 (1.04, 7.01) | 50.3 ± 26.1 [51.7] | 3.99 ± 0.648 [21.7] |
| ADC4 | DMOT4039A | 36 | 4.43 ± 2.52 [57.0] | 2.03 (1.04, 7.01) | 33.7 ± 17.8 [52.7] | 3.35 ± 0.673 [20.1] |
| ADC5 | DMUC5754A | 39 | 7.01 ± 4.41 [63.0] | 2.96 (1.07, 8.20) | 57.6 ± 44.6 [77.4] | 3.76 ± 1.57 [41.8] |
| ADC6 | DNIB0600A | 52 | 5.14 ± 3.26 [63.5] | 2.06 (0.899, 7.22) | 47.9 ± 32.4 [67.7] | 4.15 ± 1.83 [44.1] |
| ADC7 | DSTP3086S | 36 | 5.71 ± 3.24 [56.7] | 2.99 (1.98, 4.19) | 43.3 ± 21.7 [50.0] | 3.91 ± 0.654 [16.7] |
| ADC8 | DFRF4539A | 16 | 3.15 ± 1.51 [47.7] | 3.10 (2.79, 7.27) | 35.3 ± 18.4 [52.1] | 5.00 ± 1.98 [39.6] |

AUC_{inf}: area under the concentration–time curve from time zero to infinity; CL: clearance; C_{max}: maximum concentration; V_{dss}: volume of distribution at steady state; t_{1/2}: terminal half-life; T_{max}: time to C_{max}; CV: coefficient of variation; SD: standard deviation.
antibody, although numerically shorter for some ADCs, with mean values ranging from 3.0 to 5.0 d across the eight ADCs. As shown in Figures 2 and 3, the PK profiles and exposure of unconjugated MMAE were largely comparable across the eight ADCs following the first dose of 2.4 mg/kg. The differences in exposure of unconjugated MMAE (i.e., $C_{\text{max}}$ and $AUC_{\text{inf}}$ at Cycle 1) at 2.4 mg/kg between molecules were relatively small as compared to the inter-subject variability (Figure 3). The inter-subject variability of unconjugated MMAE was relatively large, with %CV for the eight ADCs ranging from 48–74% and 50%-77% for Cycle 1 $C_{\text{max}}$ and $AUC_{\text{inf}}$, respectively (Table 2(c)), while the inter-molecule variability of unconjugated MMAE across the eight ADCs was 24% and 18% for mean $C_{\text{max}}$ and $AUC_{\text{inf}}$ at Cycle 1, respectively. There was no apparent accumulation of unconjugated MMAE upon repeated dosing of vc-MMAE ADCs at 2.4 mg/kg q3w with trough concentration ratios of 1.0–1.3 between Cycle 3 and Cycle 1.

The dose levels evaluated in Phase 1 dose–escalation studies for the q3w regimen ranged from 0.1 to 3.2 mg/kg across vc-MMAE ADCs. As shown in Figure 4, PK linearity appeared to vary with different vc-MMAE ADCs. A trend of nonlinearity was observed for pinatuzumab vedotin (ADC1), DEDN6526A (ADC3) and DMUC5754A (ADC5). The clearance of acMMAE and total antibody for these three ADCs decreased with increasing doses; consistently, the corresponding inter-subject variability also decreased with the doses tested (Figure 4(a) and Figure S2). The PK for these three ADCs appeared to approach linearity at the expansion dose of 2.4 mg/kg. The remaining five ADCs exhibited apparent dose-independent clearance, indicating that PK was linear over the doses tested (Figure 4(a)). Unconjugated MMAE appeared to exhibit dose-proportional PK for all the eight ADCs, as dose-normalized exposure was overlapping and comparable over the dose tested, although inter-subject variability was relatively large (Figure 4(b)). The assessment of PK linearity is limited by small sample sizes per dose level in the q3w dose cohorts.

### Analyte correlation analyses

As shown in Figure 5(a), acMMAE exposure (i.e., $AUC_{\text{inf}}$ at Cycle 1) was strongly correlated with that of total antibody for all eight vc-MMAE ADCs. Values of $R^2$ ranged from 0.96 to 0.99 for the eight vc-MMAE ADCs. In contrast, the correlation between acMMAE and unconjugated MMAE exposure was less evident for the eight ADCs, with $R^2$ value ranging from 0.63 to 0.78 (Figure 5(b)).

### Exposure–response relationship

The exposure–efficacy relationship was explored for the following four ADCs: pinatuzumab vedotin (ADC1), polatuzumab vedotin (ADC2), DEDN6526A (ADC3) and DNIB0600A (ADC6). These four ADCs were selected because they demonstrated efficacy in the Phase 1 studies with reasonable response events to enable the analysis. Given different tumor indications may have different exposure–efficacy relationships, only one tumor indication with adequate efficacy across tested doses was selected for each ADC if multiple tumor types were assessed in the study. Total antibody was not included in the analysis, given the strong correlation on the PK exposure between acMMAE and total antibody analytes. As shown in Figure 6, for all four ADCs, patients with higher acMMAE exposure tended to have higher objective response rate (ORR). A significant exposure–response relationship was observed between acMMAE Cycle 1 $AUC_{\text{inf}}$ and ORR ($p < .05$, Figure 6).
Figure 4. Dose-dependency of acMMAE and unconjugated MMAE PK for the eight vc-MMAE ADCs.
A. acMMAE clearance vs dose
B. Dose-normalized AUC_{\text{inf}} at Cycle 1 of unconjugated MMAE vs dose
Open cycle: observed data, AUC_{\text{inf}}: area under the concentration–time curve from time zero to infinity, acMMAE: antibody-conjugated MMAE, ADC1 = Pinatuzumab vedotin (CD22); ADC2 = Polatuzumab vedotin (CD79b); ADC3 = DEDN6526A (ETBR); ADC4 = DMOT4309A (MsLN); ADC5 = DMUC5754A (MUC-16); ADC6 = DNIB0600A (Napi2b); ADC7 = DSTP3086S (Steap1); ADC8 = DFRF4539A (FcRHS).
Figure 5. Correlation of AUC$_{\text{inf}}$ at Cycle 1 between analytes post the first dose of vc-MMAE ADCs over the doses tested (90 min IV infusion).
A. acMMAE vs total antibody
B. acMMAE vs unconjugated MMAE
Open cycle: observed data. The black line and shaded gray area represent linear regression model prediction and 90% confidence interval of predictions, acMMAE: antibody-conjugated MMAE, AUC$_{\text{inf}}$: area under the concentration–time curve from time zero to infinity, ADC1 = Pinatuzumab vedotin (CD22); ADC2 = Polatuzumab vedotin (CD79b); ADC3 = DEDN6526A (ETBR); ADC4 = DMOT4309A (MsLN); ADC5 = DMUC5754A (MUC-16); ADC6 = DNIB0600A (Napi2b); ADC7 = DSTP3086S (Steap1); ADC8 = DFRF4539A (FcRH5).
for three of the four ADCs, with DEDN6526A (ADC3) as the exception. For ADC3, a trend of positive exposure–response relationship was observed between acMMAE exposure and ORR, although it is not statistically significant ($p = .067$). In contrast, there was no significant positive relationship between unconjugated MMAE exposure and ORR ($p > .05$, Figure S3). The $p$ values of exposure–efficacy relationship for acMMAE exposure were consistently lower as compared to the corresponding unconjugated MMAE exposure for all four ADCs (Figure 6 and Figure S3), suggesting that the ORR correlation was stronger with acMMAE exposure compared with unconjugated MMAE exposure.

The exposure–safety relationship was also explored with the same four ADCs. Peripheral neuropathy (PN) was the adverse event of interest for vc-MMAE ADCs, as it is the most frequent adverse event resulting in dose reductions/discontinuations for vc-MMAE ADCs. As shown in Figure 7, patients with higher exposure of acMMAE appeared to have high probability to develop grade 2+ peripheral neuropathy. The relationship was statistically significant ($p \leq 0.05$) for three of four ADCs, with DEDN6526A (ADC3) the exception. In contrast, no significant relationship was observed between unconjugated MMAE exposure and grade 2+ peripheral neuropathy for all four ADCs ($p > .05$, Figure S4). For DEDN6526A (ADC3), a trend toward a positive exposure–response relationship was observed with acMMAE exposure, although it is not statistically significant ($p = .276$, Figure 7); a flat exposure–response was observed for unconjugated MMAE ($p = .855$, Figure S4). Compared with unconjugated MMAE, acMMAE exposure appeared to exhibit stronger correlation with probability to develop grade 2+ peripheral neuropathy with $p$ value of the exposure–safety relationship consistently lower for acMMAE than that for unconjugated MMAE across all four ADCs evaluated. It is worth noting that the exposure–response assessment for each Phase 1 study is limited by small patient and event numbers.

**Discussion**

vc-MMAE ADCs are one of the most commonly used drug-linker platforms in the clinical development of ADCs. Structurally, vc-MMAE ADCs share the same vc linker, cytotoxic drug (MMAE) and conjugation chemistry, but they incorporate different mAbs against different targets and are used for different tumor indications (Figure 1). For the eight vc-MMAE ADCs described here, the average DAR was approximately the same (3–4). Given that the patient numbers for each Phase 1 study were relatively small (ranging from 33 to 95) and the clinical data were rather limited (Table 1), leveraging the learning from other molecules with the same drug-linker can be valuable in better informing decision-making, such as identifying an optimal Phase 2 dose. Understanding the correlation between analytes and exploring the potential key analyte that correlates with efficacy and/or safety across ADC platform could also inform future clinical

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**Figure 6.** Logistic regression analysis of exposure-efficacy (ORR) response relationship by acMMAE AUC$_{inf}$ at Cycle 1. The black line and shaded gray area represent logistic regression model prediction and 90% confidence interval of predictions, open circle show exposure of individual patients with event ($p = 1$) and without events ($p = 0$), black solid circle and vertical lines show observed fraction of subjects with events in each exposure quartile and 90% confidence interval for these fractions. acMMAE: antibody-conjugated MMAE, AUC$_{inf}$: area under the concentration–time curve from time zero to infinity, ADC1 = Pinatuzumab vedotin (CD22); ADC2 = Polatuzumab vedotin (CD79b); ADC3 = DEDN6526A (ETBR); ADC6 = DNIB0600A (Napi2b); iNHL = indolent non-Hodgkin’s Lymphomas; DLBCL = diffuse large B-cell lymphoma.

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pharmacology strategy for ADCs in late-stage development. The objectives of this study were to conduct integrated analysis to characterize the PK and explore the exposure–response relationship of vc-MMAE ADCs to enable better decision-making and improved development strategy for vc-MMAE ADCs, and potentially for other ADCs, entering the clinical study.

Three analytes, namely acMMAE, total antibody, unconjugated MMAE were measured in eight FIH Phase 1 studies to characterize the PK behavior of vc-MMAE ADCs. As shown in Figure 2, each analyte exhibited a distinct PK behavior in vivo. Integrating the PK behavior of the three analytes is critical to understand the distribution, clearance pathway and elimination kinetics of a vc-MMAE ADC. As shown in Figure 2, acMMAE concentrations declined more rapidly than total antibody concentrations. This is likely because two clearance pathways drive the decrease in acMMAE concentrations, namely proteolytical degradation and deconjugation, while total antibody concentrations changes are driven solely by proteolytical degradation. As a result, the difference in the concentration decrease of the two analytes could infer the rate of deconjugation from the ADC.

Across the eight vc-MMAE ADCs, the PK of acMMAE at 2.4 mg/kg was characterized by mean clearance values ranging from 16.6 to 30.2 mL/d/kg, a mean volume of distribution at steady state limited to plasma and interstitial space volume and a mean terminal half-life of approximately of 3.8–6.2 d, which likely explains the observation that repeated dosing of vc-MMAE ADCs q3w only resulted in modest accumulation of acMMAE for the eight vc-MMAE ADCs. Similar PK characteristics were also reported for brentuximab vedotin, the clinically approved vc-MMAE ADC with average DAR of 4.0 (CL: 23–25 mL/d/kg and t1/2: 4.4–5.9 d at clinically approved dose of 1.8 mg/kg). Total antibody analyte was found to have a lower clearance (approximately 8.2–23.6 mL/d/kg) and longer t1/2 (approximately 3.9–10.7 d) compared to acMMAE. It was noted that total antibody clearance was faster than that of typical mAbs at linear range (approximately 3–6 mL/d/kg). The estimated elimination half-life of total antibody is also shorter than that for a typical IgG1 antibody (2–3 weeks). These results suggest that the mAb component of the ADCs undergoes additional clearance besides the target-mediated and nonspecific proteolytic degradation pathways through which a mAb is typically eliminated. In fact, Lyon et al. showed that ADCs with hydrophobic drug linkers (e.g., vc-MMAE) undergo selective uptake by the nonparenchymal cells (i.e., sinusoidal endothelium and Kupffer cells) of the liver, thus contributing to the accelerated clearance of total antibody analyte as compared to mAbs. Overall, the PK properties of acMMAE and total antibody are more similar to a mAb than to a small molecule drug, with relatively slow clearance and a small volume of distribution that is mostly confined to plasma and interstitial space.

There was an apparent delay for unconjugated MMAE in reaching its Cmax relative to acMMAE (Tmax approximately 2–3 d vs. post-infusion), but this was expected, as time is
needed for unconjugated MMAE to be formed following cellular uptake, lysosomal internalization and degradation of a vc-MMAE ADC, and then to be released back into the systemic circulation. The released unconjugated MMAE is expected to behave like a small molecule that could be metabolized by CYP3A4/5 or excreted unchanged into the urine/feces. The half-lives of unconjugated MMAE (3.0–5.11 d) were generally much longer than half-lives expected for typical small molecules and approximated the half-lives of acMMAE and total antibody, although the mean value was numerically shorter than those of acMMAE and total antibody. This suggests that PK of unconjugated MMAE follows formation of rate-limited kinetics. Additionally, no evidence of unconjugated MMAE accumulation in plasma was observed after repeated dosing of an vc-MMAE ADC at 2.4 mg/kg q3w.

Over the dose tested across eight ADCs (0.1–3.2 mg/kg), a trend toward faster clearance at lower doses was observed for three vc-MMAE ADCs, but not for the remaining five ADCs, suggesting that PK non-linearity depends on the target expression and/or disease burden. These results, though limited, suggest that there is target-mediated clearance for the three ADCs at low doses in addition to nonspecific proteolytical degradation and deconjugation clearance. The presence of circulating soluble antigen does not necessarily result in non-linear PK. DMOT4039A and DFRF4539A, which were able to bind soluble antigen, exhibited linear PK over the dose tested. Upon increasing dose levels, the clearance of the three ADCs reached a plateau when target-mediated clearance was saturated; all eight vc-MMAE ADCs appeared to approach linear PK at 2.4 mg/kg (Figure 4(a)). Given that the clearance pathways of vc-MMAE ADCs are expected to be similar when target-mediated clearance is saturated, it is not entirely unexpected that time-concentration profiles and key parameters of the three analytes for the eight vc-MMAE ADCs demonstrated remarkable similarity at 2.4 mg/kg regardless of target and tumor indication. These findings are consistent with results from platform population PK modeling of acMMAE analytes for the eight ADCs, which showed comparable acMMAE PK for the eight vc-MMAE ADCs, with much lower inter-molecule variability compared to the inter-individual variability. Therefore, for a future novel vc-MMAE ADC entering FIH clinical study, it is conceivable that the PK properties of vc-MMAE ADCs at 2.4 mg/kg reported here could be used to project human PK of a vc-MMAE ADC within a linear range and inform its FIH study design.

Given the complex structure of an ADC with both large and small molecule components, it is essential to measure multiple analytes in the FIH Phase 1 study to understand the clearance pathway and elimination kinetics of an ADC in humans, but whether there is a need to continue measuring the multiple analytes of an ADC during late-stage clinical development is not yet clear. In order to address this question, we assessed the correlation between analytes. As shown in Figure 5, there is a strong correlation between acMMAE and total antibody exposure (AUC_{inf} at Cycle 1) for all eight vc-MMAE ADCs, with R^2 values ranging from 0.96 to 0.99, supporting that measuring total antibody analyte in late-stage clinical development (e.g., Phase 2 or 3 studies) may not be warranted. In contrast, a weaker association was observed between acMMAE and unconjugated MMAE exposure (R^2: 0.63–0.78). Lack of strong apparent correlation between acMMAE and unconjugated MMAE exposure may be partially due to the complex multi-step formation and release processes of MMAE. Additionally, the released MMAE undergoes CYP3A-mediated metabolism and biliary/renal excretion.^{24} The inter-subject variability of these complex processes, especially large inherent inter-subject variability for CYP3A,^{25} may potentially contribute to the lack of strong exposure correlation between acMMAE and unconjugated MMAE.

Given the high correlation between acMMAE and total antibody exposure, exposure–response analysis was conducted with acMMAE and unconjugated MMAE analytes only. The results showed that higher acMMAE exposure was associated with better objective response; however, weaker or no apparent relationship was observed between unconjugated MMAE exposure and ORR (Figure 6 and Figure S3). Similar observations were also made for Grade 2+ peripheral neuropathy. The probability of patients having Grade 2+ peripheral neuropathy appeared to correlate better with acMMAE exposure, rather than unconjugated MMAE exposure, with p values of exposure–response relationship numerically lower for acMMAE compared with that of unconjugated MMAE (Figure 7 and Figure S4). These findings were observed consistently across the four ADCs evaluated. Similar exposure–response findings were also observed for brentuximab vedotin, the clinically approved vc-MMAE ADC.\(^3\) These results support that acMMAE, but not unconjugated MMAE, is the main analyte associated with key efficacy and safety endpoints for vc-MMAE ADCs, which is aligned with the design concept and mechanism of action of ADCs. Theoretically, the MMAE in the tissues can come from circulating ADC (measured as acMMAE) and/or unconjugated MMAE, but the exposure–response results suggest that circulating ADC appears to play a more important role in delivering MMAE to the tissues compared to circulating unconjugated MMAE, and thus its exposure correlated better with the efficacy and safety outcomes. It is worth noting that the exposure–response assessment for each Phase 1 study is rather limited due to small patient and event numbers, but the consistent trend of the exposure–response relationships observed across 4 vc-MMAE ADCs and brentuximab vedotin strengthens the robustness of the findings. Since there is weaker or no apparent relationship between unconjugated MMAE exposure and key efficacy and safety endpoints, monitoring unconjugated MMAE analytes may be of limited value to support late-stage clinical development (such as Phase 2 and/or 3 studies).

In conclusion, the PK properties of acMMAE, total antibody and unconjugated MMAE of single-agent vc-MMAE ADCs are well characterized and demonstrated remarkable similarity at 2.4 mg/kg q3w across the eight vc-MMAE ADCs included in this analysis. Strong correlation between acMMAE and total antibody exposure, but not between acMMAE and unconjugated MMAE exposure were consistently observed for all the eight vc-MMAE ADCs. Additionally, exposure–response analysis for multiple vc-
MMAE ADCs showed that key efficacy and safety endpoints tested appeared to correlate well with acMMAE exposure, but not with that of unconjugated MMAE. Together these integrated platform analyses suggest that measurement of acMMAE analyte only might be adequate to support the clinical pharmacology strategy at the late stage of clinical development of vc-MMAE ADCs. Reducing the PK sample collection from three analytes to one analyte will not only increase the efficiency of clinical studies and reduce the development cost of an ADC, but also reduce unnecessary patient blood burden. However, it remains to be seen whether the recommended PK sampling strategy is applicable to other ADC platforms, such as ADCs with site-specific conjugation or different cytotoxic drugs.1

Materials and methods

Study designs and data

The PK analysis was based on data from FIH Phase 1 studies in cancer patients of eight ADCs that included both a dose-escalation component and at least one expansion cohort at a dose identified for further clinical study. Studied doses were in the range of 0.1–3.2 mg/kg given every 3 weeks (q3w) and in the range of 0.8–1.6 mg/kg given every week (qw). All studies included an expansion cohort at 2.4 mg/kg given every 3 weeks as a single agent. All data available in the clinical database at the time of data extraction were included in the analysis. PK sampling was rich with multiple samples per patient (n ~ 10) collected during cycle 1 (up to 21 d post first dose) and 4–5 samples/cycle up to cycle 4. In addition, samples pre-dose and post-infusion were collected every other cycle until at least cycle 8. PK concentrations of acMMAE, total antibody, and unconjugated MMAE were quantified in the clinical studies at prespecified timepoints. For two of the ADCs developed for non-Hodgkin lymphoma (pinatuzumab vedotin and polatuzumab vedotin), the FIH studies included single-agent treatment only as well as in combination with the standard of care drug, rituximab. The PK data from single-agent treatment given every 3 weeks were the main focus of the current analyses.

Bioanalytical methods for PK concentrations of the three analytes

The acMMAE concentrations in plasma were quantified using a validated method consisting of Protein A affinity chromatography to capture the ADC followed by the enzyme-mediated release of MMAE, and liquid chromatography-tandem mass spectrometry (LC-MS/MS) for detection of MMAE.9 The acMMAE assay measured the total concentration of MMAE conjugated to the antibody moiety. The total antibody assay quantified all forms of fully conjugated, partially deconjugated, and fully deconjugated antibody, which was measured in patient serum samples using a validated enzyme-linked immunosorbent assay (ELISA) method.9 Unconjugated MMAE, a measure of the total plasma MMAE that was unconjugated from the antibody, was quantified using a validated LC-MS/MS method.

PK assessment

PK analyses included all patients who received at least one vc-MMAE ADC dosing and adequate blood sampling for at least one PK parameter at cycle 1. PK parameters at cycles 1, including C\text{max}, AUC\text{inf}, clearance, V\text{dss}, and t\text{1/2} were estimated with non-compartmental analyses (NCA) using WinNonlin 5.2.1 when data permitted. PK parameters were summarized and compared by molecules and dose levels using descriptive statistics and graphically. Relationships between exposure of any two analytes were explored graphically and by simple linear regression.

Exposure–response analysis

The following efficacy and safety endpoints were examined in the exposure–response analyses: ORR and Grade 2+ peripheral neuropathy. AUC\text{inf} at cycle 1 for acMMAE and unconjugated MMAE analytes were used as the exposure metrics. Given different tumor indications may have different exposure–efficacy relationships, only one tumor indication with adequate efficacy across tested doses was selected for each ADC if multiple tumor types were assessed in the study. All the patients who received at least one dose of the ADC and had enough PK samples to estimate AUC\text{inf} at Cycle 1 across tested doses were included in exposure-safety analysis. Logistic regression analyses were conducted for ORR and Grade 2+ peripheral neuropathy by including AUC\text{inf} at Cycle 1 of acMMAE or unconjugated MMAE as a continuous variable. The analyses were performed using generalized linear model functions in Splus 8.2 (Solution Metrics, Sydney, Australia).

Compliance with ethical standards

All the eight clinical studies were approved by the relevant Institutional Review Boards and Medical Ethics Committees and were carried out in accordance with International Conference on Harmonization E6 Guidelines for Good Clinical Practice and with the Helsinki declaration of 1975 or comparable ethical standards. Written informed consent was obtained for all patients in accordance with federal and institutional guidelines.

Acknowledgments

The authors would like to thank all the patients and investigators for their participation in the clinical studies. The authors also thank clinical study teams for each project for their thoughtful study design and careful implementation of the clinical study. Support for third-party editing assistance for this manuscript was provided by F. Hoffmann-La Roche, Ltd.

Declaration of Interest Statement

CL, CZ, ZL, DS, DL, BW, SC, RZ, PA, BF and SG are paid employees of Genentech, Inc. and stockholders of Roche Holding Ltd.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.
**Funding**

This work was funded by F. Hoffmann-La Roche, Ltd.

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