DCDT2980S, an anti-CD22-Monomethyl Auristatin E antibody-drug conjugate, is a potential treatment for non-Hodgkin's Lymphoma

DCDT2980S for the treatment of non-Hodgkin’s lymphoma

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Abstract

Antibody-drug conjugates (ADCs), potent cytotoxic drugs linked to antibodies via chemical linkers, allow specific targeting of drugs to neoplastic cells. We have used this technology to develop the ADC DCDT2980S that targets CD22, an antigen with expression limited to B-cells and the vast majority of non-Hodgkin’s lymphomas (NHL). DCDT2980S consists of a humanized anti-CD22 monoclonal IgG1 antibody with a potent microtubule disrupting agent, monomethyl auristatin E (MMAE), linked to the reduced cysteines of the antibody via a protease cleavable linker, maleimidocaproyl-valine-citrulline-p-aminobenzoyloxycarbonyl (MC-vc-PAB). We describe the efficacy, safety and pharmacokinetics of DCDT2980S in animal models to assess its potential as a therapeutic for the treatment of B-cell malignancies. We did not find a strong correlation between in vitro or in vivo efficacy and CD22 surface expression, nor a correlation of sensitivity to free drug and in vitro potency. We show that DCDT2980S was capable of inducing complete tumor regression in xenograft mouse models of NHL and can be more effective than rituximab plus combination chemotherapy at drug exposures that were well tolerated in cynomolgus monkeys. These results suggest that DCDT2980S has an efficacy, safety, and pharmacokinetics profile that support potential treatment of NHL.
Introduction

Malignancies of B-cell origin are a heterogeneous group of neoplasms, which vary in genetic drivers of transformation, B-cell subtype of origin and clinical outcome (1). Major unmet medical needs for B-cell malignancies include non-Hodgkins lymphoma (NHLs) and chronic lymphocytic leukemia (CLL). The subclasses of NHL range in clinical outcome from slow-growing indolent and incurable diseases with a median survival of 8-10 years, such as follicular non-Hodgkins lymphoma (NHL) (2), to more aggressive intermediate- to high-grade lymphomas (such as diffuse large-cell lymphoma), which can have a median survival of 6 months if left untreated or long-term remission in more than 50% of patients with appropriate treatment (3). Mantle cell lymphoma (MCL) shares characteristics of both types, being incurable with a short remission following treatment. Diffuse large B-cell lymphoma (DLBCL) is the most common type of NHL accounting for approximately 30-40% of all new patients, whereas follicular lymphoma (FL) and MCL account for approximately 20 to 25% and 6 to 10% of new lymphomas, respectively. CLL is the most common chronic leukemia in adults with approximately 15,000 new cases per year in the United States (4) and like follicular lymphoma is indolent but incurable. Despite advances in the clinical outcomes of patients with NHL and CLL using treatments such as the CD20-specific monoclonal antibody rituximab (Rituxan, MabThera), indolent B-cell malignancies remain incurable as are approximately half of aggressive NHL patients. Thus, there is an unmet medical need for new therapies that are preferably better tolerated.
than the current standards of care.

Previously we identified CD22 and CD79 as superior targets for antibody-drug conjugate (ADC), potent cytotoxic drugs linked to antibodies via chemical linkers, therapy for the treatment of NHL (5-7). CD22 is a 130 kDa type 1 transmembrane glycoprotein, normally expressed on B-lineage cells from the pre-B-cell stage, and to being fully expressed on all subtypes of mature B-cell and down regulated from the surface of plasma cell. CD22 is also expressed on most malignant mature B cells, including follicular NHL, marginal zone lymphoma (MZL), MCL, DLBCL, small lymphocytic lymphoma (SLL), and CLL ((6, 8), this manuscript). CD22 is not expressed on non-B lymphoid cells including, myeloid cells, hematopoietic stem cells, or any other non-hematopoietic lineage thus making it an excellent target for antibody and ADC therapy for a wide range of B-cell malignancies (9). Antibodies and antibody-drug conjugates targeted to CD22 are being tested in clinical trials for efficacy in NHL. An unconjugated humanized anti-CD22 antibody, Epratuzumab, is currently in clinical trials for the treatment of B-cell malignancies (10) and lupus (11). The ADC inotuzumab ozogamicin (IO, CMC-544) a human IgG4 monoclonal antibody to CD22 conjugated via an acid-labile hydrazone linker to a cytotoxic DNA damaging agent calicheamicin (12, 13) is being tested in clinical trials for treatment of NHL. Both of these anti-CD22 therapeutics have shown efficacy and have not displayed major toxicities relating to the targeting of CD22.

In this paper we describe the efficacy, safety and pharmacokinetics of the anti-CD22 ADC DCDT2980S in animal models. DCDT2980S consists of a humanized anti-CD22 monoclonal IgG1 antibody, MCDT2219A, with a potent antimitotic
chemotherapeutic agent, monomethyl auristatin E (MMAE) chemically linked to the reduced cysteines of the antibody via a protease labile linker, maleimidocaproyl-valine-citrulline-p-aminobenzoyloxycarbonyl (MC-vc-PAB). We have investigated efficacy and safety of DCDT2980S in animal models to assess its potential as a therapeutic for the treatment of B-cell malignancies.

**Materials and methods**

**Antibodies and antibody-drug conjugates**

The antibody for DCDT2980S (MCDT2219A, Hu10F4) was generated and humanized as previously described (6). The antibody was conjugated as previously described (14). Briefly, ADCs were prepared by incubating the maleimido drug derivative with the partially reduced antibodies for 1 hr at 4°C. After quenching the reaction with excess N-acetyl-cysteine, to react with any free linker-drug, the conjugated antibody was purified. Conjugation conditions were chosen to achieve an average drug-to-antibody ratio (DAR) of approximately 3.5. ADC protein concentrations were calculated using absorbance at 280 nm (320 nm reference) and the molar extinction coefficient of the antibody. The average DARs were calculated from the integrated areas of the DAR species resolved by hydrophobic interaction chromatography (HIC) on an analytical column (TSK butyl-NPR 4.6mm x10cm 2.5mm, Tosoh Bioscience, King of Prussia, PA, US).
Cell lines

The NHL cell lines DOHH2, Granta-519, Karpas-1106P, MHH-PREB-1, NU-DUL-1, NU-DHL-1, Ramos, RC-K8, REC-1, SC-1, SU-DHL-1, SU-DHL-4, SU-DHL-5, SU-DHL-6, SU-DHL-8, SU-DHL-10, SU-DHL-16, U-698-M, WSU-DLCL2 and WSU-NHL were obtained from DSMZ (Braunschweig, Germany). The DB, Farage, HT, MC116, Pfeiffer and Toledo cell lines were obtained from the American Type Culture Collection (Manassas, VA). The A4/Fukada, SCC-3, TK were obtained from JHSF (Osaka, Japan). All cell lines were maintained in RPMI 1640 supplemented with 10% fetal bovine serum (Sigma, St. Louis, MO) and 2 mM L-glutamine. Each cell line authenticated by Short Tandem Repeat (STR) Profiling using the Promega PowerPlex 16 System and compared to external STR profiles of cell lines to determine cell line ancestry. In addition an SNP fingerprint is generated from the original thaw to serve as our internal master fingerprint. The SNP fingerprinting is performed each time a new batch is frozen down. Cell lines were typically used for several months before thawing a new passage.

CD22 expression levels in xenograft tumors

To measure the CD22 expression on xenograft tumors, recovered tumors were minced and put through a 30 μm cell strainer (BD Biosciences) to achieve a single-cell suspension. The tumor cells were subsequently prepared by the
standard density centrifugation over lymphocyte separation medium (MP Biomedical). The resulting single-cell suspension was stained with anti-human CD22-APC antibody (BD Biosciences, clone S-HCL-1), anti-human CD20-Pacific Blue (Beckman Coulter) and 7-amino-actinomycin D (BD Biosciences). Data were analyzed using FlowJo (Tree Star), and the mean fluorescent intensity (MFI) was calculated from the CD20+ and 7-amino-actinomycin D negative population. CD22 expression on cell lines was measured with the same CD22-APC antibody.

**CD22 expression levels in patient tumors**

Tissue biopsies were dissociated using a needle and syringe. FACSFlow (10 ml) (BD Biosciences, Oxford, UK, Cat. No. 342003) was injected into the biopsy to flush leucocytes from the mass. Leucocytes from tissue, bone marrow aspirates and peripheral blood were all then prepared using ammonium chloride lysis to remove red cells. A volume of sample was incubated with a 10-fold excess of ammonium chloride (8.6 g/L in distilled H₂O, Vickers Laboratories, Pudsey, UK, Cat. No.0055-CO) for 5 min at 37 °C. Cells were then washed twice in buffer (FACSFlow containing 0.3% bovine serum albumin (Sigma, UK, Cat. No. A3059)). Leucocytes (1x10⁶) were stained with 30 µL of a pre-prepared cocktail of antibodies. Samples were assayed using the panel of antibodies described. Following incubation in the dark at 4 °C for 20 minutes cells were washed twice in buffer and re-suspended in FACSFlow. Samples were acquired on a FACS Canto II analyser using FACSDiva software (BD Biosciences, Oxford, UK). A minimum of 50,000 events were acquired for each tube of the panel.
Data analysis was carried out using FacsDiva software (Biosciences, Oxford, UK). B-cells were gated using CD19 PE-Cy5.5 (SJ25-C1), FSC and SSC parameters. Appropriate non-B cell pan markers served as isotype controls for gating of positive and negative quadrants. CD8 APC (SK1) was used as an isotype control for CD22 APC (S-HCL-1). Mean Fluorescence intensities (MFIs) presented as normalized values (MFI CD22-MFI CD8).

**Cell viability assay**

Cells were plated in quadruplicate at 1-5x 10^3 per well in 384-well plates in RPMI containing 10% fetal bovine serum overnight before treated with DCDT2980S or the control anti-gD-drug conjugate. Each conjugate was added to experimental wells at a final concentration of 50, 16.7, 5.5, 1.8, 0.61, 0.2, 0.067, 0.022 or 0.007 µg/mL. For MMAE a final concentration of 200, 40, 8, 1.6, 0.32, 0.064, 0.0128, 0.0026, 0.00051 nmol/L was tested with control wells receiving medium alone. After 72 h incubation at 37 °C, cell viability was measured using the Celltiter-Glo Luminescent Cell Viability Assay (Promega Corp., Madison, WI). The concentration of DCDT2980S or MMAE resulting in the 50% inhibition of cell...
viability was calculated from a four-variable curve analysis and was determined from a minimum of three independent biological replicate experiments.

**Xenograft experiments**

Xenograft experiments were performed as previously described (15). Briefly, all animal studies were performed in compliance with National Institutes of Health guidelines for the care and use of laboratory animals and were approved by the Institutional Animal Care and Use Committee (IACUC) at Genentech, Inc. Cells were inoculated subcutaneously into the flanks of female CB17 ICR SCID mice. When mean tumor size reached desired volume, the mice were divided into groups of 7-9 mice with the same mean tumor size and dosed intravenously (IV) via the tail vein with ADCs or antibodies. Rituximab was dosed at 30 mg/kg intraperitoneally (IP), which is above the maximum efficacious dose. CHOP (single IV injection of 30 mg/kg cyclophosphamide, 2.475 mg/kg doxorubicin, 0.375 mg/kg vincristine and oral dosing of 0.15 mg/kg prednisone once a day for 5 days) was dosed to the maximum tolerated dose defined as the dose, where a 5% weight loss in the animals was observed.

**Pharmacokinetic study in SCID mice**

The PK study in SCID mice was approved by the Institutional Animal Care and Use Committee at Genentech, Inc. Female SCID mice received a single intravenous (IV) dose of 0.5 or 5 mg/kg DCDT2980S via the tail vein.
(n=20/group). Blood samples were collected via retro-orbital bleeds performed on alternating eyes and the terminal blood sample was collected via cardiac stick from each animal in each dosing group at the following time points: predose; 5 minutes; 1, 6, and 24 hours; and 2, 3, 4, 7, 10, 14, 21, and 28 days post-dose, and processed to collect serum. Three blood samples were taken from each mouse and there were four mice per time point. Serum concentration–time data were used to estimate relevant PK parameters.

Multiple dose toxicity and toxicokinetics study in cynomolgus monkeys

The toxicity and toxicokinetics study in cynomolgus monkeys was approved by the Institutional Animal Care and Use Committee and conducted at Covance Laboratories, Inc. (Madison, WI). Animals were dosed intravenously every three weeks for a total of five doses on study days 1, 22, 43, 64, and 85 with vehicle or DCDT2980S at 1, 3, or 5 mg/kg and were necropsied on day 92 (3 animals/sex/group), day 127 (2 animals/sex/group) or after confirmation of B-cell recovery on day 232 (2 animals/sex/group). Blood was collected pre-study and at selected time points throughout each study for analyses of hematology, serum chemistry, coagulation, toxicokinetics, anti-therapeutic antibodies, and measurement of circulating lymphocyte populations by flow cytometry. Additionally, ophthalmologic and physical examinations were conducted during the pre-dose phase, and at the end of the first and last dose cycles as applicable. Cardiovascular safety pharmacology end points were also evaluated over the
course of the study using either electrocardiogram external leads. At necropsy, organ weights were measured on select organs and tissues were thoroughly examined by gross and microscopic examination.

**Anti-tumor efficacy study in mice bearing BJAB and WSU-DLCL2 human non-Hodgkin’s lymphoma xenografts**

The efficacy studies in mice were approved by the Institutional Animal Care and Use Committee at Genentech, Inc. Female C.B.-17 SCID mice (8-10 week old) were injected with 20 million human non-Hodgkin’s lymphoma BJAB or WSU-DLCL2 cells, suspended in 0.2 mL of HBSS, subcutaneously in the right flank. When tumors reached the desired volume, mice were randomized into nine groups (n=9-10/ per group). Mice bearing BJAB xenografts received a single IV dose of vehicle (control group), CNJ1135 (a non-binding control MC-vc-MMAE conjugate) at 4 mg/kg, MCDT2219A (unconjugated anti-CD22 antibody) at 4 mg/kg, or DCDT2980S (treatment groups) at doses of 0.1, 0.5, 1, 1.5, 2, and 4 mg/kg. Mice bearing WSU-DLCL2 xenografts received a single IV dose of vehicle (control group), CNJ1135 (a non-binding control MC-vc-MMAE conjugate) at 16 mg/kg, MCDT2219 (unconjugated anti-CD22 antibody) at 16 mg/kg, or DCDT2980S (treatment groups) at doses of 1, 2, 4, 8, 12, and 16 mg/kg. Tumors were measured twice each week for the duration of the study using UltraCal-IV calipers and tumor volume was calculated using the following formula: Tumor Volume (mm$^3$) = (length × width$^2$) × 0.5. The results were plotted as mean tumor volume ± SEM of each group over time. Tumor growth inhibition (TGI) as a
percentage of vehicle was calculated using the following formula: \( \% \text{TGI} = 100 \times \left[1 - \frac{\text{AUC}_{\text{treatment/day}}}{\text{AUC}_{\text{vehicle/day}}} \right] \). Partial response (PR) was defined as a tumor regression of > 50% but < 100% of the starting tumor volume, and complete response (CR) was defined as 100% tumor regression (i.e., no measurable tumor) on any day during the study. Time to tumor doubling (TTD) was defined as the time in days for a tumor to double in volume from its initial volume on Day 0. Student’s t-test was used to evaluate the difference in TTD distributions between groups with a p-value of \( \leq 0.05 \) to be significant.

**Bioanalysis of serum samples from pharmacokinetic and toxicokinetic studies**

After administration of DCDT2980S, total antibody (conjugated and unconjugated anti-CD22 antibody) concentrations were analyzed in serum by two ELISAs. For serum samples from the pharmacokinetic and toxicokinetic studies in cynomolgus monkeys, microtiter plates were coated with an anti–complementarity determining region antibody against anti-CD22 and incubated overnight, followed by blocking. DCDT2980S standards, controls, and samples were added to the coated plate and incubated for 2 hours to allow for binding. After washing, sheep anti–human IgG conjugated to horseradish peroxidase was added, incubated for 1 hour, and then washed again. TMB substrate (Kirkegaard & Perry Laboratories,
Inc.) was used for detection and absorbance was measured at 450 nm against a reference wavelength of 620 or 630 nm. The minimum quantifiable concentration of the assay was determined to be 80 ng/mL. For serum samples from the pharmacokinetic study in SCID mice, anti-CD22 standards and samples diluted in sample diluent were incubated for 2 hours with biotinylated sheep anti-human IgG antibody (Genentech, Inc.) and goat anti-human IgG antibody conjugated to horseradish peroxidase in either micronic tubes (National Scientific) or polypropylene round-bottom plates (Corning Costar) to form a bridging complex. Complexed samples were transferred to NeutrAvidin coated plates (Thermo Fisher Scientific Inc.) and incubated for 1 hour. After washing, the detection step was carried out using TMB substrate (Kirkegaard & Perry Laboratories, Inc.). Absorbance was measured at 450 nm against a reference wavelength of 620 nm. The minimum quantifiable concentration of the assay was determined to be 63 ng/mL.

**Pharmacokinetic data analysis**

Serum concentration-time profiles were used to estimate the following PK parameters in mouse and monkey, using non-compartmental analysis (WinNonlin, version 5.2.1; Pharsight Corporation, Mountain View, CA): total drug exposure defined as area under the serum concentration–time curve extrapolated to infinity (AUC_{inf}), area under the concentration-time curve from time 0 to TK day 21 (AUC_{0-21}), area under the concentration-time curve from TK day 84 to TK day 105 (AUC_{105}), and observed maximum serum concentration.
(Cmax). A naïve pooled approach was used in mouse to provide one estimate for each dose group, while in monkey, each animal was analyzed separately and results for each dose group were summarized as mean ± standard deviation (SD).

Results

Generation and Characterization of DCDT2980S

The antibody anti-CD22 10F4 has been previously identified as an antibody to CD22 that is particularly effective as an ADC compared to other anti-CD22 antibodies when tested in xenograft models (6). The humanized form of 10F4 (MCDT2219A; Hu10F4) (6) was conjugated to MC-vc-PAB-MMAE though the interchain disulfide cysteines (Figure 1A) as previously described (14). Hydrophobic interaction chromatography (HIC) analysis of DCDT2980S to detect conjugated antibody showed that the average drug to antibody ratio (DAR) for DCDT2980S was 3.6 and consisted of 5% 0-drug, 28% 2-drug, 49% 4-drug, 14% 6-drug, and 3% 8-drug species (Figure 1B). The antibody MCDT2219A binds cynomolgus monkey and human CD22 but not rat and mouse CD22 ((6), Figure S1). The ADC DCDT2980S demonstrated a high binding affinity to human CD22 (Kd = 1.7 +/- 0.2 nM) by equilibrium binding similar to that of MCDT2219A. Binding affinity of DCDT2980S was similar between cynomolgus monkey (Kd = 1.8 +/- 0.1 nM) and human.
Surface CD22 and sensitivity to MMAE are poor predictors of sensitivity to DCDT2980S in vitro

To test the efficacy and reveal parameters that could determine sensitivity to DCDT2980S we assessed cell viability in a large panel of 35 NHL cell lines (Table S1). The degree of sensitivity to DCDT2980S and an isotype control anti-gD-vcMMAE was determined from a dose-response curve of 0 – 50 µg/mL and IC$_{50}$ values were calculated (Table S1). To ensure that the IC$_{50}$ values calculated were specific to DCDT2980S, we compared IC$_{50}$ values to that of the isotype control anti-gD-vcMMAE. In most of the cell lines that expressed surface target the IC50 of anti-gD-vcMMAE was much greater than the IC50 of DCDT2980S, however, five cell lines had similar IC50 for targeted and control ADC (Table S1). These data demonstrate that DCDT2980S has very potent and broad activity across a large panel of NHL cell lines. Since surface expression levels of CD22 might be an important driver of response to DCDT2980S, we performed flow cytometry across the panel of NHL cell lines. We then assessed whether there was a correlation with the IC$_{50}$ values obtained with DCDT2980S (Figure 2A). The overall correlation was low but statistically significant ($R^2 =0.23$, $P =0.003$). However the data suggest that while the higher expressing cell lines (MFI >1000) predict a sensitive cell line (IC50 < 5 ug/mL); cell lines that express lower levels of CD22 are not predictive of response. That is, cell lines with low levels of CD22 could either be quite sensitive to DCDT2980S or comparatively resistant. We also examined how sensitivity of the cell line to MMAE and DCDT2980S correlated (Figure 2B)
and found a moderate and statistically significant correlation ($R^2 = 0.39$, $p < 0.0001$). These data suggest that sensitivity to the free drug and the amount of surface target effect how a cell line responds to DCDT2980S in vitro but neither is a predictor of response.

**Efficacy of DCDT2980S in xenograft models of B-cell lymphoma**

In addition to testing the activity of DCDT2980S in vitro, we tested the activity of DCDT2980S in vivo using xenograft tumor models with a wide range of expression of CD22. We evaluated the expression level of CD22 in several xenograft models and found the expression level of CD22 changed when grown in vivo compared to the expression levels when grown in culture (Figure 3). We selected two models for efficacy studies, BJAB for its relatively low expression, (91% of primary tumors have greater surface expression) and WSU-DLCL2 for its relatively high in vivo expression of CD22 (8% of primary tumors have greater surface expression) (Figure 3). In vitro, these cell lines are equally sensitive to free drug (IC50 = 0.07 nm/L, Figure 2 and Table S1). Mice bearing BJAB tumors (volume range of 160–285 mm$^3$) were given a single dose of vehicle; DCDT2980S at 0.1, 0.5, 1, 1.5, 2, or 4 mg/kg; MCDT2219A (unconjugated DCDT2980S) at 4 mg/kg; or a isotype control (non-binding) conjugate (CNJ1135) at 4 mg/kg (Figure 4A). Based on the comparisons of tumor growth inhibition (TGI) and time to tumor doubling (TTD), DCDT2980S showed clear dose-dependent inhibitory activity versus the vehicle
group at dose levels > 0.5 mg/kg (p<0.0001). Of the 10 mice receiving 1.5 mg/kg DCDT2980S, 4 had partial responses (PRs) and 6 had CRs. DCDT2980S at a dose of ≥ 2 mg/kg reached the maximal efficacy, resulting in (complete responses) CRs in all treated mice. The inhibitory effects of a single dose of DCDT2980S were durable, such that 6 of 10 mice receiving 1.5 mg/kg DCDT2980S and all the mice given ≥ 2 mg/kg DCDT2980S maintained CRs until the end of study.

MCDT2219A (the unconjugated anti-CD22 antibody) at 4 mg/kg did not have an effect on tumor growth. The non-binding conjugate CNJ1135 at 4 mg/kg had a minimal effect on tumor growth. Low level of activity of the control conjugate might be due to a low level of CD22-independent delivery of MMAE to tumor cells, however activity from the group given DCDT2980S at the matching dose level was substantially superior to that from the group given CNJ1135.

Pilot experiments with the tumor xenograft model of human diffuse large B-cell lymphoma WSU-DLCL2 indicated that this model that was more resistant to DCDT2980S; therefore tumor bearing mice (volume range of 200−325 mm3) were given a single dose of vehicle; DCDT2980S at 1, 2, 4, 8, 12, or 16 mg/kg; MCDT2219A at 16 mg/kg; or a control (non-binding) conjugate at 16 mg/kg (Figure 4B). Based on comparisons of TGI and TTD, DCDT2980S showed clear dose-dependent inhibitory activity versus the vehicle group at dose levels greater than 2 mg/kg (p < 0.0001). Four of 9 mice given 8 mg/kg DCDT2980S and 7 of 9 mice given 12 mg/kg DCDT2980S had PRs. Finally, DCDT2980S at 16 mg/kg exhibited the greatest efficacy, with all treated mice experiencing tumor regression (one PR and eight CRs). MCDT2219A (unconjugated DCDT2980S) at 16 mg/kg
did not have an effect on tumor growth. The non-binding conjugate CNJ1135 at 16 mg/kg had a modest effect on tumor growth, possibly due to a low level of CD22-independent delivery of MMAE to tumor cells or non-specific uptake of the ADC, and was not unexpected at this high level. TGI from the group given DCDT2980S at the matching dose level was substantially superior to that from the group given CNJ1135. As in the in vitro study a lower level CD22 of expression did not predict lower activity of DCDT2980S in the in vivo study.

**Safety profile of DCDT2980S in cynomolgus monkey**

Administration of DCDT2980S in mouse xenograft models did not appear to negatively affect the mice since they appeared healthy and gained weight at all does levels. However, since mice are particularly tolerant of microtubule inhibiting drugs (16) and DCDT2980S does not bind mouse or rat CD22, therefore rodents are not an appropriate model to test the safety of DCDT2980S. DCDT2980S binds cynomolgus monkey and human CD22 with almost equal affinity (Figure S1) so cynomolgus monkey was chosen as the relevant experimental animal for evaluating the tolerability of DCDT2980S.

Cynomolgus monkeys were intravenously administered DCDT2980S at 1, 3, or 5 mg/kg every 3 weeks for 5 doses. Doses up to 5 mg/kg were well tolerated with no adverse effects on body weight or clinical evidence of toxicity. Notable findings associated with DCDT2980S exposure included reversible bone marrow toxicity
and associated hematopoietic changes. Specifically, monkeys were observed with dose dependent reversible neutropenia (Figure 5A) and reticulocytopenia (Figure 5B) at 3 and 5 mg/kg. We also detected minor decreases in erythrocytes, hemoglobin, and hematocrit at 3 and 5 mg/kg, and corresponding bone marrow hypocellularity at all doses (Data not shown). No cardiovascular, respiratory, renal, gastrointestinal, neurologic, or ophthalmic abnormalities were observed at any dose.

Pharmacokinetics of DCDT2980S in mice and monkeys.

In mice, total antibody exposure in the area under concentration–time curve in serum extrapolated to infinity (AUC_{inf}) following single IV administration of 0.5 mg/kg and 5 mg/kg of DCDT2980S was 72.1 day•μg/mL and 821 day•μg/mL, respectively; suggesting a dose-proportional PK behavior.

In cynomolgus monkeys, exposure of total antibody increased slightly more than dose proportionally from 1 to 3 mg/kg, and dose proportionally from 3 mg/kg to 5 mg/kg. The exposure of last dose interval (AUC_{84-105}) at 3 and 5 mg/kg was higher than that of the first dose interval (AUC_{0-21}), due to depletion of the target B-cells up DCDT2980S upon treatment (F.K. Fuh and M. Williams). Thus, concentration- and time- dependent pharmacokinetics of DCDT2980S were observed in cyno, as expected.

To gain a better understanding of the therapeutic potential of DCDT2980S, we employed a strategy that assessed the efficacy of DCDT2980S in mouse...
xenograft models at an exposure (AUC) consistent with the highest non-severely toxic dose (HNSTD) in cynomolgus monkeys. From the HNSTD of 5 mg/kg in the multiple dose cyno toxicity study we calculated an AUC$_{84-105}$ (Table 1), representing the exposure to DCDT2980S after a single IV dose. Using this exposure value and mouse PK data, it was determined that a mouse dose of 4 mg/kg would provide similar DCDT2980S exposure in mice as the dose achieved at the cyno HNSTD ($628 \pm 71.6$ day$\cdot$µg/mL). This dose level, 4 mg/kg, has been shown to be efficacious in multiple xenograft models (Figures 4 and 6) suggesting that efficacious DCDT2980S exposures can be achieved at doses that are well-tolerated.

Comparison of the activity of DCDT2980S standard of care in vivo

Since we had determined doses in mice that resulted in exposures that might be achievable in humans we invested how the activity of DCDT2980S compared to current standards of care in xenograft models. The current front-line standard of care for most NHLs is a combination of rituximab (anti-CD20 antibody) cyclophosphamide, doxorubicin, vincristine and prednisone (R-CHOP). To compare the activity DCDT2980S to R-CHOP and extend our understanding of the vivo efficacy of DCDT2980S we assessed the activity of DCDT2980S in RAMOS-RA1 and Granta-519 lymphoma models. In these studies, we compared the activity of a single dose of DCDT2980S with a single cycle of CHOP,
rituximab, or R-CHOP. Rituximab was given above the maximum efficacious
dose and CHOP was given close to the maximum tolerated dose for mice (15).
Since the exposure equivalent of the maximum tolerated dose of DCDT2980S
was approximately 4 mg/kg for a single dose exposure we dosed DCDT2980S
once at 5 mg/kg as top end of our exposure and 2 mg/kg once to provide a very
conservative dose. In the Granta519 model DCDT2980S was substantially more
effective than R-CHOP in inhibiting the tumor growth (Figures 6A; p< 0.0001).
Based on the comparisons of TGI and TTD, DCDT2980S showed clear inhibitory
activity versus the vehicle group for the two dose levels tested (p < 0.0001).
DCDT2980S at 2 and 5 mg/kg produced a TGI of 120% and 126%, and
ultimately led to sustained tumor regression in all treated mice. Seven of the 9
mice given 2 mg/kg DCDT2980S and all of the mice given 5 mg/kg DCDT2980S
maintained CRs until the end of study, Day 28 (see Figure 6). In contrast, the
dose of CHOP or R-CHOP delayed tumor growth and did not achieve any tumor
regression, with a TGI of 79% and 90%, respectively, and a TTD of 12 and 13.5
days. Rituximab alone at 30 mg/kg had a modest effect on delaying tumor
growth. The non-binding conjugate CNJ1135, given at 5 mg/kg, had a noticeable
effect on tumor growth, yielding a TGI of 68% and a TTD of 10 days. Activity due
to a low level of CD22-independent delivery of MMAE to tumor cells is not
unexpected at this dose level. Even so, the TGI from the group given
DCDT2980S at the matching dose level was still superior to that from the group
given CNJ1135.
In the Ramos RA1 model a single dose of DCDT2980S resulted in dose-dependent inhibition of tumor growth (Figure 6). Based on the comparisons of TGI and TTD, DCDT2980S showed clear inhibitory activity versus the vehicle group for the two dose levels tested ($p \leq 0.0005$). DCDT2980S at 2 mg/kg and R-CHOP had comparable activity, with a TGI of 76% and 73%, respectively, and a TTD of and 6.5 days, compared with a TTD of 2.5 days in the vehicle group. The effects of 5 mg/kg DCDT2980S were more pronounced, with a TGI of 112%, and ultimately led to sustained tumor regression in all treated mice. Of the 7 mice given 5 mg/kg DCDT2980S, 5 maintained CRs until the end of the study. As expected, the non-binding conjugate CNJ1135, given at 5 mg/kg, did not have an effect on tumor growth. These data suggest that DCDT2980S could be very effective compared to standard of care. At a minimum, they suggest that the excellent tumor responses we observe are not due to an extreme sensitivity of the models to antibody therapy or chemotherapy.

**Discussion**

Here we describe DCDT2980S, an anti-CD22-MC-vc-PAB-MMAE antibody-drug conjugate, and its preclinical safety and efficacy. We found that DCDT2980S was broadly active in vitro and that comparatively low amounts of target were not a barrier to efficacy. Further, the vitro activity translated to activity of DCDT2980S in in vivo models of NHL. Since the antibody is not cross-reactive to rodents and mice are uniquely insensitive to auristatin type drugs (16), cynomolgus monkeys
were used to assess the DCDT2980S safety profile. DCDT2980S was well tolerated in cynomolgus monkeys with neutropenia as a predominant safety finding. The pharmacokinetics of DCDT2980S in SCID mice where efficacy studies were conducted, appeared to be dose proportional. In cynomolgus monkeys, the pharmacokinetics of DCDT2980S appeared to be slightly non-linear in the dose range of 1–3 mg/kg and became dose proportional in the dose range of 3 to 5 mg/kg of DCDT2980S. Thus, the contribution of B-cell (CD22) mediated clearance to total clearance at the dose range of 1 to 3 mg/kg appeared to be moderate. In mouse and cynomolgus monkey PK studies, the concentrations of the conjugate (DCDT2980S) were also measured (data not shown). Comparison of the total antibody and DCDT2980S concentrations over time assessed the \textit{in vivo} stability of the conjugate. The DCDT2980S serum concentration decreased biexponentially, in a manner similar to total antibody, suggesting that MMAE is not rapidly released from DCDT2980S.

To understand the potential efficacy of DCDT2980S at drug exposures that were tolerated in cynomolgus monkeys, we translated the a single dose exposure found to be safe in a five dose cynomolgus monkey study to our mouse models and found that an exposure of a single dose would be equivalent to approximately 4 mg/kg in mice. This would be an efficacious dose in all four of the xenograft models we tested. Again, levels of surface CD22 that were low compared to the levels of CD22 expression found in primary NHL tumors were not a barrier to efficacy. The BJAB and WSU-DLCL2 models had very different sensitivities to the DCDT2980S, ED 90 of 1.1 vs 14.5 mg/kg respectively,
however, the BJAB model had lower surface expression of CD22 and the cell lines had equal sensitivity to MMAE in vitro. The drivers of in vivo sensitivity to DCDT2980S are unclear at this point, however our data suggest that DCDT2980S could be broadly active and not limited to patients with high CD22 expression. The observation that R-CHOP combination therapy dosed at near the maximum tolerated dose is not as efficacious as relevant doses of the ADC suggest that the excellent in vivo activity we observed is not due to a general sensitivity to chemotherapy in our xenograft models. Assuming that cynomolgus monkeys is a reasonable model for human PK properties and tolerability of DCDT2980S, DCDT2980S could be of therapeutic utility in the majority of NHL patients. This conclusion is also supported by our current clinical understanding of the target and linker-drug. CD22 is a clinically validated target for both monoclonal antibody therapy and ADC therapy (17, 18) and MC-vc-MMAE is the linker drug used in the anti-CD30 ADC brentuximab vedotin which is approved for the treatment of patients with Hodgkin lymphoma after failure of autologous stem cell transplant (ASCT) or after failure of at least two prior multi-agent chemotherapy regimens. The objective response rate in the brentuximab vedotin in Hodgkin lymphoma patients was 75% suggesting that the linker-drug technology used in DCDT2980S has the potential to be broadly effective in the treatment of NHL at tolerated dose (19). Based on the work and rationale described here DCDT2980S is being tested in Phase 1 trials for the treatment of NHL and CLL.
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References


Table 1. Group Mean (±SD) Pharmacokinetic Parameters of Total Antibody after a Single IV Administration of DCDT2980S to SCID Mice and Multiple (q3w x 5) IV Administrations of DCDT2980S to Cynomolgus Monkeys

<table>
<thead>
<tr>
<th>Dose (mg/kg)</th>
<th>Mouse (Single Dose)</th>
<th>Cynomolgus Monkey (q3w x 5)</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( C_{\text{max}} ) (µg/mL)</td>
<td>( AUC_{\text{inf}} ) (µg*day/mL)</td>
<td>( C_{\text{max}} ) (µg/mL)</td>
<td>( AUC_{0-21} ) (µg*day/mL)</td>
<td>( C_{\text{max}} ) (µg/mL)</td>
</tr>
<tr>
<td>1</td>
<td>NA</td>
<td>NA</td>
<td>24.9 ± 2.97</td>
<td>62.9 ± 6.06</td>
<td>26.9 ± 1.13</td>
</tr>
<tr>
<td>3</td>
<td>NA</td>
<td>NA</td>
<td>87.8 ± 11.0</td>
<td>259 ± 24.1</td>
<td>108 ± 13.4</td>
</tr>
<tr>
<td>5</td>
<td>134</td>
<td>821</td>
<td>134 ± 18.1</td>
<td>388 ± 39.6</td>
<td>159 ± 13.2</td>
</tr>
<tr>
<td>0.5</td>
<td>11.1</td>
<td>72.1</td>
<td>NA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4 (estimated)</td>
<td>NA</td>
<td>628 ± 71.6</td>
<td>NA</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Figure 1. Description of DCDT2980S. A) Structure of DCDT2980S. Only one MC-vc-MMAE attached to an interchain disulfide bond linker drug is shown for clarity. B) HIC chromatogram showing the drug distribution of DCDT2980. MCDT2219A, the unconjugated antibody is shown for comparison.

Figure 2. Efficacy of DCDT2980 in vitro. In vitro cell killing (IC50, µg/mL) of 35 non-Hodgkin’s lymphoma cell lines by DCDT2980 as compared to A) surface expression of CD22 (mean fluorescence intensity, MFI) and B) sensitivity to free MMAE.

Figure 3. Flow cytometry analysis of CD22 expression on NHL patient samples compared to xenograft tumors and cell lines. Surface expression of CD22 was assessed by FACS (mean fluorescence intensity, MFI) on primary tumor samples from patients with CLL (n=349), DLBCL (n=158), FL (n=236), HCL (N=16), MCL (n=66), and MZL (n=78) as well as on xenograft tumors and cell lines. Red lines show the median of each patient population.

Figure 4. Efficacy of DCDT2980 in tumor xenograft models. Tumor growth was plotted as mean (± SEM) tumor volume of each group receiving a single intravenous dose of vehicle, CNJ1135 (non-binding control conjugate), MCDT2219A (unconjugated anti-CD22) or DCDT2980S over the duration of the study. (A) BJAB xenografts (n=10 mice/group) with an average starting tumor
volume of 230 mm$^3$ were treated with the doses indicated (B) WSU-DLCL2 xenografts (n=9 mice/group) with an average starting tumor volume of 270 mm$^3$ were treated with the doses indicated.

**Figure 5.** Effect of DCDT2980S on selected hematology in cynomolgus monkeys. The effects of the DCDT2980S dosed every three weeks five times (arrows) with the indicated amounts on (A) Neutrophils and (B) Reticulocytes. Error bars represent standard deviation.

**Figure 6.** *In vivo* efficacy of DCDT2980S compared to standard of care. Mouse xenograft models of NHL with an average starting tumor volume of 200-300 mm$^3$ treated with a single dose of 2 or 5 mg/kg DCDT2980S compared to a single treatment of Rituximab, CHOP, Rituximab plus CHOP. Tumor growth was plotted as mean (± SEM) tumor volume of each group. (A) Granta-519 xenografts with 9 mice treated as indicated. (B) Ramos-RA1 xenografts with 7 mice per group were treated as indicated.
Figure 1

A

B

Elution Time (min)

Absorbance (280 nm)

0.0
200
100
0

MCOT2219A

0-drug
2-drug
4-drug
6-drug

UCOT2980S

0.0
5.0
10.0
15.0

0.0
100
200

2-drug
4-drug
6-drug
8-drug

Figure 1
Figure 2

A

DCDT2980S IC50 (μg/mL) vs. Surface expression (MFI)

- \( R^2 = 0.23 \)
- \( P = 0.003 \)

B

DCDT2980S IC50 (μg/mL) vs. free MMAE IC50 (nM)

- \( R^2 = 0.39 \)
- \( P < 0.0001 \)
CD22 expression in tumor models

8% of tumors have higher expression than WSU-DLCL2 model

91% of tumors have higher expression than BJAB model
**Figure 4**

**A**

- **BJAB**
  - Vehicle
  - Control ADC
  - Unconjugated Anti-CD22
  - 0.1 mg/kg
  - 0.5 mg/kg
  - 1 mg/kg
  - 1.5 mg/kg
  - 2 & 4 mg/kg

- **Tumor volume (mm³)**

- **Days**

- **Single dose**

  - **ED 50** = 0.4 mg/kg
  - **ED 90** = 1.1 mg/kg

**B**

- **WSU-DLCL2**
  - Unconjugated Anti-CD22
  - Control ADC, 16 mg/kg
  - 1 mg/kg
  - 2 mg/kg
  - 4 mg/kg
  - 8 mg/kg
  - 12 mg/kg
  - 16 mg/kg

- **Tumor volume (mm³)**

- **Days**

- **Single dose**

  - **ED 50** = 2.6 mg/kg
  - **ED 90** = 14.5 mg/kg

---

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Figure 5

**A**

Mean Absolute Neutrophils +/- SD

**B**

Mean Absolute Reticulocytes +/- SD

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Figure 6
Molecular Cancer Therapeutics

DCDT2980S, an anti-CD22-Monomethyl Auristatin E antibody-drug conjugate, is a potential treatment for non-Hodgkins Lymphoma

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