CD25-targeted antibody–drug conjugate depletes regulatory T cells and eliminates established syngeneic tumors via antitumor immunity

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ABSTRACT

Background Regulatory T cells (Tregs) contribute to an immunosuppressive tumor microenvironment. They play an important role in the establishment and progression of tumors with high Treg infiltration and present a major obstacle to tumor eradication by immunotherapies. Numerous strategies have been attempted to deplete or block Tregs, although their success has been limited.

Methods A CD25-targeted, pyrolobenzodiazepine (PBD) dimer-based antibody–drug conjugate (ADC) was investigated for its ability to deplete Tregs and induce antitumor immunity. Antitumor activity of CD25-ADC either alone or in combination with an anti-programmed cell death protein 1 (PD-1) antibody was evaluated in CD25-negative syngeneic models that exhibit tumor infiltration of CD25-expressing Tregs, and its pharmacodynamics and pharmacokinetics were assessed.

Results Single low doses of CD25-ADC resulted in potent and durable antitumor activity in established syngeneic solid tumor models and the combination of a suboptimal dose was synergistic with PD-1 blockade. Tumor eradication by the CD25-targeted ADC was CD8+ T cell-dependent and CD25-ADC induced protective immunity. Importantly, while CD25-ADC mediated a significant and sustained intratumoral Treg depletion, accompanied by a concomitant increase in the number of activated and proliferating tumor-infiltrating CD8+ T effector cells, systemic Treg depletion was transient, alleviating concerns of potential autoimmune side effects.

Conclusions This study shows that a PBD dimer-based, CD25-targeted ADC is able to deplete Tregs and eradicate established tumors via antitumor immunity. This represents a novel approach to efficiently deplete Tregs via a very potent DNA damaging toxin known to induce immunogenic cell death. Moreover, this study provides proof of concept for a completely new application of ADCs as immunotherapeutic agents, as the main mode of action relies on the ADC directly targeting immune cells, rather than tumor cells. These strong preclinical data warrant the clinical evaluation of camidanlumab tesirine (ADCT-301), a PBD-based ADC targeting human CD25, either alone or in combination with checkpoint inhibitors in solid tumors with known Treg infiltration. A phase I trial (NCT03621982) of camidanlumab tesirine in patients with selected advanced solid tumors is ongoing.

INTRODUCTION

Antibody–drug conjugates (ADCs), which consist of a monoclonal antibody (mAb) conjugated to a cytotoxic drug via a chemical linker, have emerged as a novel class of anti-cancer therapeutics. The antibody component of the ADC binds to tumor-specific or tumor-associated antigens and delivers a potent cytotoxic agent at the target site. Several ADCs have received approval for cancer therapy, with many more under evaluation in various phases of clinical development.

Camidanlumab tesirine (ADCT-301) is an ADC comprising HuMax-TAC, a human IgG1 mAb directed against human CD25, stochastically conjugated via a cathespin-cleavable valine–alanine peptide linker to the potent pyrolobenzodiazepine (PBD) dimer warhead SG3199, with a drug-to-antibody ratio (DAR) of 2.3. PBD dimers have distinct advantages over other warheads as they form non-distortive interstrand cross-links in the minor groove of DNA, which are refractory to DNA repair allowing persistence of the DNA interstrand cross-links. SG3199 has been shown to be highly cytotoxic against multiple human solid and hematological cancer cell lines, with mean 50% growth inhibitory concentrations in the pM range. Further, PBD-based ADCs are able to target low copy number antigens and typically have a low DAR (~2) compared with other ADCs based on more conventional auristatin or maytansine warheads (DAR: ~ 4).

CD25 (interleukin (IL)-2Rα) is part of the heterotrimetric IL-2 receptor that regulates normal immune function and is widely expressed on the surface of leukemias and lymphomas. Clinical trials of CD25-targeted radioimmunoconjugates and immunotoxins in patients with CD25-expressing lymphomas have demonstrated clinical proof of concept.
for CD25 as a potential therapeutic target. Based on promising data in preclinical models, camidanlumab tesirine is currently being evaluated in multiple clinical trials in Hodgkin and non-Hodgkin’s lymphoma (NCT02432235 and NCT04052997), with promising interim results.

Regulatory T cells (Tregs) play an important role in the establishment and progression of tumors and are considered a major obstacle to tumor eradication by immunotherapies. Infiltration of Tregs contributes to the immunosuppressive tumor microenvironment (TME) in a variety of cancers including, but not limited to, ovarian, lung, pancreatic, colorectal, and melanoma. Moreover, the intratumoral balance between Tregs and effector T cells (Teffs) appears to influence the outcome of immunotherapies, and poor prognosis in solid tumors is often associated with high tumor infiltration by Tregs and a low ratio of Teffs to Tregs.

Numerous attempts are underway to exploit the therapeutic potential of depleting Tregs. One of them relies on targeting CD25, which is highly expressed on tumor-infiltrating Tregs, but absent on naïve Teffs. Depleting or suppressing Tregs via anti-CD25-based therapies, alone or in combination with checkpoint inhibitors, could, therefore, be an effective strategy for tumor eradication, particularly in solid tumors that harbor a large number of tumor-infiltrating Tregs.

We, therefore, proposed that camidanlumab tesirine, in addition to its direct cytotoxic activity in CD25-expressing tumor cells, may have antitumor activity by depleting CD25-expressing Tregs and enabling tumor eradication through Teffs. Preclinical studies evaluating the efficacy of immune modulatory agents such as checkpoint inhibitors are generally performed in immunocompetent mice grafted with mouse tumor cell lines (syngeneic models). As the antibody component of camidanlumab tesirine is directed against human CD25 and does not cross react with mouse CD25, we generated CD25-ADC, a CD25-targeted ADC composed of PC61, a mAb directed against mouse CD25, stochastically conjugated to tesirine, the same PBD drug-linker payload employed in camidanlumab tesirine.

This study evaluated the antitumor activity of CD25-ADC either alone or in combination with an antiprogrammed cell death protein 1 (PD-1) antibody in CD25-negative syngeneic colon cancer models that exhibit tumor infiltration of CD25-expressing Tregs, and assessed its pharmacodynamics and pharmacokinetics (PK).

MATERIALS AND METHODS

Generation of ADCs

PC61, a rat IgG1k mAb purified from hybridoma PC61 5.3, was conjugated to tesirine (SG3249) for generating CD25-ADC, essentially as previously described for camidanlumab tesirine. In brief, antibody was buffer exchanged into a proprietary histidine buffer at pH 6 using a TRIS/EDTA pH 8.5 buffer, and the solution was reduced with Tris (2-carboxyethyl) phosphine reductant. Dimethylacetamide and SG3249 (threefold excess relative to antibody) were added to the solution. The conjugation reaction was incubated, then quenched with threefold molar excess of N-acetyl cysteine and incubated again. The pH was then decreased to 6.0 using histidine hydrochloride solution and the generated CD25-ADC was purified by tangential flow filtration, filtered, and stored at −70°C. Final yield was estimated by ultraviolet–visible spectrophotometry based on starting antibody.

The rat isotype control ADC (C0010-5-SG3249) and the non-binding control ADC (B12-SG3249) were generated by stochastic conjugation of C0010-5 (rat isotype antibody, Crown Bioscience, California, USA) and B12 (human IgG1 non-binding antibody, produced in house), respectively, to SG3249.

Characterization of CD25-ADC

Characterization of synthesized CD25-ADC was performed by size exclusion chromatography (SEC), hydrophobic interaction chromatography (HIC), and reduced reverse-phase (RP) liquid chromatography, using standard techniques as previously described.

In vitro binding assay

Binding of PC61 to mouse recombinant CD25 (R&D Systems) was determined by ELISA, using a mouse CD25/human Fc chimeric antigen (R&D Systems) and a secondary goat antirat HRP (Jackson Immunoresearch Laboratories). Optical density was measured using an Envision plate reader at 450 nm.

CD25 expression

Analysis of CD25 expression on murine cell lines was performed by flow cytometry. Data were analyzed using ForeCyt software, and the data were reanalyzed using FlowJo software (V.10) to determine the half-maximal effective concentration (EC₅₀) values.

Murine cell lines

The source of cell lines used in this study along with cell growth media is shown in online supplementary table S6.

In vitro cell killing

Cytotoxicity of CD25-ADC, the free warhead SG3199, and isotype control ADC C0010-5-SG3249 was determined. In brief, cell lines were incubated with serial dilutions of CD25-ADC, the isotype control ADC, or the free warhead SG3199 for 5 days at 37°C in a 5% carbon dioxide-gassed, humidified incubator. Cell viability was measured with CellTiterGlo luminescent assay (Promega Corporation) according to manufacturer’s instructions. The data were normalized to vehicle-treated cells. EC₅₀ values were determined by using GraphPad software (GraphPad). The mean and standard errors of three independent EC₅₀ values were determined.
Animal care compliance
The Institutional Animal Care and Use Program at Charles River Discovery Services is accredited by the Association for Assessment and Accreditation of Laboratory Animal Care International, which assures compliance with accepted standards for the care and use of laboratory animals.

Assessment of efficacy using in vivo models
MC38 tumors were established in 9-week-old, female C57BL/6 mice (Charles River, Massachusetts, USA) by implanting 5x10^5 MC38 cells subcutaneously (s.c.) into their flanks. CT26 tumors were established in 9-week-old, female BALB/c mice (Charles River) by implanting 3x10^5 CT26 cells s.c. into their flanks. When group mean tumor volumes reached specified target ranges, animals were randomly sorted into groups and treatments were initiated. CD25-ADC, the non-binding control ADC (B12-SG3249) or isotype control ADC (C0010-5-SG3249) were administered on day 1, intraperitoneally (i.p.) as a single dose at 5 mg/kg on days 2, 5, and 8). A vehicle-initiated CD25-ADC (0.5 mg/kg) was started at specified group mean tumor volumes. Mice were sorted into groups to receive treatment with CD25-ADC (0.5 mg/kg on day 1), and/or anti-PD-1 antibody (5 mg/kg on days 2, 5, and 8), or vehicle control (phosphate buffer saline (PBS)). For the efficacy analysis, 10 mice/group were monitored for 36 days. For the T-cell immunophenotype analysis, untreated mice (n=8) were collected on day 1 (pre-dose). Samples of tumor, blood and thymus were collected from all groups including vehicle (n=8 per group) at specified time points. Cell suspensions were processed for flow cytometry.

Flow cytometry
Blood samples were prepared by centrifugation and lysis of the red blood cells. Immediately following blood collection, tumor and/or selected organs were harvested from each animal and processed for flow cytometry. Mouse tumor samples were dissociated according to the manufacturer’s instruction using the gentle MACS protocol “Tumor Dissociation Kit.” Samples were filtered through a 70 µm cell strainer and rinsed twice in PBS/2.5% fetal bovine serum buffer to remove enzymatic buffer. Spleen, lymph nodes and thymus samples were dissociated by gently grinding the tissue across a 70 µm cell strainer with RPMI-1640 media, followed by lysis by the red blood cells. The final single-cell suspensions were prepared in Staining Buffer (2.5% fetal bovine serum, 0.09% sodium azide, in PBS, pH 7.4) at 2x10^7 cells/mL. Briefly, 100 µL of single-cell suspensions were pelleted, resuspended in Live/Dead Aqua (Life Technologies) as per manufacturer’s instructions, and stained for 30 min at 4°C. After two washes with Staining Buffer, Fc receptors were blocked using Mu TruStain fcX (Biolegend) for 10 min at 4°C, washed twice with Staining Buffer and probed with antibody panels for 30 min at 4°C protected from light. Details on cell surface markers used for flow cytometry are shown in online supplementary table S7 along with representative staining for cell surface markers used for flow cytometry are shown.

T-cell immunophenotype studies
Seven-week-old, female C57BL/6 non-tumor-bearing mice were sorted into groups (n=24/group) to receive treatment with a single dose of CD25-ADC (0.5 mg/kg), isotype control ADC C0010-5-SG3249 (0.5 mg/kg), or no treatment (control). Spleen, lymph nodes, blood, and thymus were collected from six mice/timepoint and cell suspensions were processed for flow cytometry.

MC38 tumors were established as previously described in 9-week-old, female C57BL/6 mice and treatments were started at specified group mean tumor volumes. Mice were sorted into groups to receive treatment with CD25-ADC (0.5 mg/kg on day 1), and/or anti-PD-1 antibody (5 mg/kg on days 2, 5, and 8), or vehicle control (phosphate buffer saline (PBS)). For the efficacy analysis, 10 mice/group were monitored for tumor volume over 36 days. For the T-cell immunophenotype analysis, untreated mice (n=8) were collected on day 1 (pre-dose). Samples of tumor, blood and thymus were collected from all groups including vehicle (n=8 per group) at specified time points. Cell suspensions were processed for flow cytometry.

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Treatment tolerability was assessed by body weight measurements and by frequent observation for clinical signs of treatment-related side effects.
with phorbol 12-myristate 13-acetate (PMA, 20 ng/mL), ionomycin (500 ng/mL) and GolgiPlug in complete Roswell Park Memorial Institute (RPMI) medium for 5 hours at 37°C. For quantification of absolute number of cells, a defined number of fluorescent beads (CountBright Absolute Counting Beads, ThermoFisher) was added to each sample before acquisition and used as a counting reference.

**PK of CD25-ADC in mice**

PK analysis of CD25-ADC was performed in 7-week-old, female C57BL/6 non-tumor-bearing mice. Serum samples were collected for each timepoint after a single-dose administration of CD25-ADC (0.1, 0.5, or 1 mg/kg). Quantitation of total (unconjugated and conjugated) antibody was determined by electrochemiluminescence immunoassay using recombinant mouse CD25 as capture and a biotin-labeled polyclonal goat anti-rat IgG (mouse adsorbed; Bio-Rad) in combination with sulfoTAG streptavidin as detector. For each dose group, six mice were intravenously injected with CD25-ADC; serum was collected from three animals/group at 1, 6, 48, 96, 180, and 504 hours following dosing, and from the other 3 animals at 3, 24, 72, 120, and 336 hours following dosing. PK parameters were determined by non-compartmental analysis (Phoenix WinNonLin).

**Statistical analyses**

Statistical analyses were done using GraphPad Prism version 8.1 (Log-rank test and analysis of variance with a Tukey post-hoc test, followed by Kruskal-Wallis test with Dunn’s post-hoc test) or JMP software (Dunn method for joint ranking). Results were considered significant when p<0.05, **p≤0.05; ***p≤0.01; ****p≤0.001.

**Patient and public involvement**

No patients were involved in this study.

**RESULTS**

**Generation and in vitro characterization of CD25-ADC**

CD25-ADC is an ADC composed of PC61, a rat IgG1k anti-CD25 antibody directed against mouse CD25, stochastically conjugated to SG3249 (tesirine; figure 1A). The resulting ADC was determined to be 96.5% monodisperse by SEC, with a DAR of 2.1 by RP liquid chromatography and 2.0 by HIC. As a control, an isotype ADC composed of a rat isotype IgG1 antibody stochastically conjugated to tesirine (C0010-5-SG3249) was prepared (98% Monomer, DAR-RP 2.01, DAR-HIC 1.87). The PC61 antibody demonstrated strong binding affinity to both recombinant mouse CD25 (EC₅₀ 1.182 nM; figure 1Bi) and to CD25 expressed on the surface of a mouse lymphoma (Yac-1) cell line (EC₅₀ 0.9 nM); however, the isotype control antibody did not bind to CD25 (figure 1Bii).

The in vitro cytotoxicity of CD25-ADC was determined in a panel of murine tumor-derived cell lines including CD25-positive Yac-1 and CD25-negative MC38, CT26, 4T1, LL/2, and B16-F10. CD25-ADC showed potent and specific cytotoxicity in the CD25-expressing Yac-1 cell line (CD25-ADC EC₅₀ 11.5 pM (±1.56); isotype-ADC EC₅₀ 8575 pM (±1072); figure 1Bv), but it did not bind or have any specific activity in the CD25-negative cell lines (figure 1Biii,iv,vi,vi and online supplementary figure S1i–vi). In contrast, SG3199, the warhead component of CD25-ADC, showed potent cytotoxicity in all the CD25-negative cell lines tested, in line with its non-targeted mode of action (figure 1Bvii and online supplementary figure S1ii,vii).

**In vivo antitumor activity of CD25-ADC**

CD25-ADC had strong, dose-dependent, and durable antitumor activity in established s.c. syngeneic tumor models. In the CD25-negative MC38 colon cancer model, a single dose of CD25-ADC at 0.5 and 1 mg/kg resulted in rapid antitumor responses, sustained for the 60-day study period, with 8/10 TFS and 2/10 PR at both 0.5 and 1 mg/kg (figure 2vi,vii and online supplementary table S1) and significant increased survival compared with the vehicle group (p<0.001, figure 2xi and online supplementary figure S2). Strong synergy (CDI: 0.3) was observed when a suboptimal single dose of 0.1 mg/kg of CD25-ADC was combined with an anti-PD-1 antibody (figure 2viii) resulting in significant increased survival compared with each monotherapy group (p<0.05, figure 2xi and online supplementary figure S2). No additional benefit was obtained from adding an anti-PD-1 antibody to CD25-ADC at 0.5 mg/kg and 1 mg/kg doses owing to the strong antitumor activity of CD25-ADC monotherapy at these doses (figure 2ix–x). Vehicle alone had no effect on disease activity, with no responses observed (figure 2i). Anti-PD-1 antibody administered alone at 5 mg/kg gave 0/10 PR and 3/10 TFS (figure 2ii). A non-binding control ADC (B12-SG3249; 97.8% Monomer, DAR-RP 2.08, DAR-HIC 1.79) dosed at 1 mg/kg had some activity alone (2/10 PR and 2/10 TFS) and in combination with anti-PD-1 antibody (2/10 PR and 5/10 TFS; online supplementary table S1); however, this combination was not synergistic (CDI: 2.1) and the activity was inferior to that observed with the same dose of CD25-ADC (figure 2ii,iv and online supplementary figure S2).

In the CD25-negative CT26 colorectal cancer model, a single dose of CD25-ADC (0.1, 0.5, or 1 mg/kg) showed dose-dependent antitumor activity (figure 3vi–vii) and significant increased survival compared with the vehicle group (p<0.001, figure 3xi and online supplementary figure S3). Synergistic activity was observed when an anti-PD-1 antibody was combined with a low dose (0.1 mg/kg) of CD25-ADC (CDI: 0.3; figure 3viii) resulting in significant increased survival compared with each monotherapy group (p<0.001, figure 3xi and online supplementary figure S3). The antitumor activity of CD25-ADC at 0.5 mg/kg and 1 mg/kg doses was further increased on combining with anti-PD-1 antibody (7/10 TFS in the group treated with 0.5 mg/kg of CD25-ADC + anti-PD-1...
antibody and 8/10 TFS in the group treated with 1 mg/kg of CD25-ADC + anti-PD-1 antibody; figure 3ix,x and online supplementary table S2). There were no responses in the group treated with vehicle (figure 3i), anti-PD-1 alone (figure 3ii), or isotype control ADC, although minor antitumor activity was observed when the isotype control ADC was combined with the anti-PD-1 antibody (1/10 TFS; figure 3iii,iv and online supplementary table S2).

In both studies, all regimens were well tolerated, with minimal body weight loss (online supplementary figure S4) and no clinical observations reported.

**In vivo mode of action studies**

We first investigated whether treatment with CD25-ADC induces immunological memory. Following treatment with CD25-ADC alone or in combination with anti-PD-1 antibody, TFS mice were rechallenged on the opposite flank with the same tumor cells as originally implanted (MC38 or CT26). Tumor-naïve mice were also implanted and used as controls. None of the TFS mice developed new tumors as opposed to all treatment-naïve controls (online supplementary figure S5), indicating that CD25-ADC induced antitumor immunity. Interestingly, following treatment with the control ADCs, either monotherapy or in combination with anti-PD-1 antibody, the few TFS animals from both the MC38 and CT26 studies also did not develop new tumors on rechallenge (online supplementary figure S5).

Next, we evaluated whether CD8+ T effs contribute to the mode of action of CD25-ADC. A CD8-depleting antibody was administered together with CD25-ADC to mice bearing established MC38 tumors. Depletion of CD8+ T cells abolished the antitumor activity of CD25-ADC both alone and together with anti-PD-1 (figure 4 and online supplementary table S3), indicating that CD25-ADC antitumor activity is mediated by CD8+ T effs and that CD8+ T effs are not negatively impacted by CD25-ADC.

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**Figure 1** Structure and in vitro characterization of CD25-ADC. (A) Structure and (B) in vitro characterization of CD25-ADC. (i) ELISA showing binding of anti-CD25 antibody PC61 to mouse recombinant CD25. (ii–iv) Flow cytometry measurement of PC61 and isotype-control antibody binding to Yac-1, MC38 and CT26 cells. (v–vii) Yac-1, MC38 and CT26 cells’ viability after exposure to CD25-ADC and isotype-ADC (and the naked pyrrolobenzodiazepine-dimer SG3199 in MC38 and CT26 cell lines). MFI, median fluorescence intensity; PABA, para amino benzoic acid.
Finally, we performed a longitudinal T-cell immunophenotype study following a single dose of CD25-ADC (0.5 mg/kg), either alone or in combination with an anti-PD-1 antibody, in mice bearing established MC38 tumors. CD25-ADC monotherapy had greater antitumor activity than vehicle or anti-PD-1 treatment (online supplementary figure S6), and synergistic activity was observed when CD25-ADC treatment was combined with anti-PD-1 (CDI: 0.133), with the combination resulting in 4/10 TFS at the end of the study (online supplementary table S4). The

Figure 2  In vivo antitumor activity of CD25-ADC in the s.c. MC38 syngeneic model. Treatment with (i) vehicle, (ii) anti-PD-1 antibody (5 mg/kg, on days 2, 5, and 8), (iii) non-binding ADC (1 mg/kg, single dose on day 1) alone or (iv) in combination with anti-PD-1 antibody, (v–vii) CD25-ADC (0.1, 0.5, and 1 mg/kg single dose on day 1) alone or (viii–x) in combination with anti-PD-1 antibody, started at a group mean tumor volume of 103 mm³. Data are shown as tumor volumes (mm³) over time for each individual mouse (10 mice/group). (xi) Survival of mice shown in i–x. Lines for G4, G5, G8 and G9 are overlapping.
immunophenotype analysis was carried out at 2, 8, and 11 days post dose of CD25-ADC, which was 1 day after the first dose of anti-PD-1 and 1 and 4 days after the last dose of anti-PD-1, respectively. A single dose of CD25-ADC, either alone or in combination with anti-PD-1, induced robust and durable intratumoral Tregs depletion, which was still maintained at 11 days post dose (Figure 5A, top panel). CD8+ T-eff levels were not negatively impacted by

Figure 3  In vivo antitumor activity of CD25-ADC in the s.c. CT26 syngeneic model. Treatment with (i) vehicle, (ii) anti-PD-1 antibody (5 mg/kg, on days 2, 5, and 8), (iii) isotype-ADC (1 mg/kg, single dose on day 1) alone or (iv) in combination with anti-PD-1 antibody, (v–vii) CD25-ADC (0.1, 0.5, and 1 mg/kg single dose on day 1) alone or (viii–x) in combination with anti-PD-1 antibody, started at a group mean tumor volume of 110 mm³. Data are shown as tumor volumes (mm³) over time for each individual mouse (10 mice/group). (xi) Survival of mice shown in i–x.
CD25-ADC, and, in fact, there was a statistically significant increase in the absolute number of tumor-infiltrating CD8+ T cells on day 11, which was further increased when CD25-ADC was combined with anti-PD-1 antibody (figure 5A, mid panel). Following CD25-ADC treatment, the CD8+ T_{eff}/T_{reg} ratio continued to increase throughout the time course analyzed, and a further increase in the ratio was observed in the combination therapy group (figure 5A, lower panel), in line with the increased antitumor activity of CD25-ADC and anti-PD-1 combination treatment. Interestingly, tumor-infiltrating CD8+ T_{eff} from the CD25-ADC-treated groups (either alone or in combination with anti-PD-1) had significantly higher activation (p<0.01, figure 5B, top panels) and proliferation rates compared with CD8+ T_{eff} from the control groups on day 11 (p<0.01, figure 5B, middle panels) and their interferon-γ production peaked on day 8, although the increase did not reach statistical significance compared with the controls (figure 5B, bottom panels). CD25-ADC treatment also reduced the number of circulating T_{reg}, but the effect was not durable, with values returning to control levels by day 11 post dose (figure 6A, top panel). Circulating CD8+ T_{eff} levels were not affected by CD25-ADC treatment, either alone or in combination with anti-PD-1 antibody, and the CD8+ T_{eff}/T_{reg} ratio returned to control levels by day 11 post dose (figure 6A, middle and low panels). In the thymus, CD25-ADC induced a temporary small increase in T_{reg}, but the CD8+ T_{eff}/T_{reg} ratio did not change during the time course analyzed (figure 6B).

In vivo studies in non-tumor-bearing mice

The T_{reg} depletion activity of CD25-ADC was further assessed in non-tumor-bearing immunocompetent mice over a period of 21 days. A single dose of CD25-ADC (0.5 mg/kg) caused a substantial but transient depletion of T_{reg} in spleen (87% reduction; p≤0.001), lymph nodes (85% reduction; p≤0.001), and blood (96% reduction; p<0.001) compared with the vehicle and isotype control ADC at 6 days post dose, with no significant impact on T_{eff} levels (p>0.05; figure 7A,B and online supplementary figure S7). Conversely, CD25-ADC induced a significant increase in both T_{reg} (328%; p≤0.001) and T_{eff} (254%; p≤0.01) in the thymus at day 6 post dose (figure 7C). T_{reg} levels in spleen, lymph nodes, blood, and thymus were restored to similar levels as the control by day 13 post CD25-ADC dosing. No significant body weight loss (online supplementary figure S8) or clinical observations were recorded throughout the duration of the study in any treatment group.

PK analysis of CD25-ADC in non-tumor-bearing mice showed dose-dependent, target-mediated drug disposition with non-linear PK at low-dose levels and linear PK at higher dose levels (online supplementary figure S9). A dose-dependent increase in exposure (area under curve, AUC [\text{tau}]) was observed, ranging from 14,572 ng×hour/mL to 751,771 ng×hour/mL (online supplementary table S5).

DISCUSSION

ADCs are an emerging class of therapeutic agents for the treatment of various cancers, and they are generally employed to target specific antigens overexpressed on malignant cells.22 There is increasing interest in targeting tumor-infiltrating CD25-positive T_{reg} to disrupt the immunosuppressive TME, as these CD25-positive T_{reg} undermine the antitumor immune function of T_{eff}. Here, we evaluated the efficacy of CD25-ADC, a CD25-targeted ADC, to provide proof of concept for its immunotherapeutic application via depletion of CD25-positive tumor-infiltrating T_{reg} in CD25-negative established solid tumors. This represents a completely novel application of ADCs, as the main mode of action does not depend on the ADC directly targeting the tumor cells, but rather relies on the ADC targeting immune cells in the local TME to provide

Figure 4 Role of CD8+ T_{eff} cells in CD25-ADC antitumor activity in the MC38 syngeneic model. Depletion of CD8+ T_{eff} cells significantly reduces the antitumor activity of CD25-ADC. CD25-ADC was administered intraperitoneally (i.p.) at a group mean tumor volume of 89 mm³ as a single dose on day 1 at 0.5 mg/kg alone or in combination with anti-PD-1 antibody (5 mg/kg, on days 2, 5, and 8). Anti-CD8 T-cell depleting antibody (10 mg/kg) was injected i.p. on days 0, 5, 8, and 13. Data are shown as mean tumor volumes (mm³) ± SEM over time (n=10/group).
therapeutic activity regardless of whether the tumor cells express the ADC’s target antigen.

Characterization of CD25-ADC showed that it is an appropriate surrogate for camidanlumab tesirine, a clinical-stage human CD25-targeting ADC, as evidenced by its specific binding to CD25-positive cells, its highly potent and specific cytotoxic activity against CD25-positive tumor cells, and its target-mediated clearance in vivo, with dose-dependent exposure and target-mediated drug disposition similar to camidanlumab tesirine. Moreover, similar to PC61, the antibody component of camidanlumab tesirine (HuMax-TAC) blocks the binding of IL-2 to the IL-2 receptor (data not shown).

A single dose of CD25-ADC exhibited potent, durable, and dose-dependent anticancer activity in two CD25-negative established solid tumor models with infiltrating Tregs. Notably, when CD25-ADC was tested at suboptimal doses, it showed strong synergy with an anti-PD-1 antibody.
Importantly, because synergistic antitumor activity was observed at suboptimal doses of the ADC, potent antitumor responses in patients could potentially be achieved at lower doses of the ADC, thereby minimizing ADC-induced toxicity.

CD25-ADC’s antitumor activity may be mediated through several distinct mechanisms. Given CD25-ADC’s potent and targeted activity against CD25-expressing cells and the selective CD25 expression by T_{reg} cells, we hypothesized that CD25-ADC could specifically deplete T_{reg}.

A single dose of CD25-ADC induced potent and durable depletion of intratumoral T_{reg}, and this was accompanied by an increase in the number of activated and proliferating tumor-infiltrating CD8+ T_{eff} cells. Interestingly, although T_{reg} depletion was similar when CD25-ADC was used alone or in combination with the anti-PD-1 antibody, the increase in tumor-infiltrating CD8+ T_{eff} was higher when CD25-ADC was combined with anti-PD-1 antibody, in line with the synergistic antitumor activity of the combination. CD25-ADC also depleted T_{reg} in blood, but contrary to the intratumoral milieu, the effect was transient and CD8+ T_{eff} cells were not affected. Similarly, in non-tumor-bearing mice, CD25-ADC transiently depleted T_{reg} in spleen, lymph nodes, and blood, with recovery of cell counts to control levels by day 13 post CD25-ADC. Conversely, neither CD8+ nor CD4+ T cells were affected by CD25-ADC. Importantly, the strong T_{reg} depletion observed in spleen, lymph nodes, and blood

Figure 6  Circulating and thymic T-cell immunophenotype analysis in MC38-bearing mice. (A) Absolute quantification of circulating T_{reg}, CD8+ T cells and CD8+/T_{reg} ratio following i.p. treatment with anti-PD-1 antibody or CD25-ADC or the combination of CD25-ADC and anti-PD-1. Blood was processed at the indicated times (days post CD25-ADC dose). (B) Absolute quantification of thymic T_{reg} cells and CD8+/T_{reg} ratio following i.p. treatment with anti-PD-1 antibody or CD25-ADC or the combination of CD25-ADC and anti-PD-1. Thymus was processed at the indicated times (days post CD25-ADC dose). Statistical differences between treatment groups were calculated using JMP 15 by the Dunn method for joint ranking. Results were considered significant when p<0.05. *, p≤0.05; **, p≤0.01.
was accompanied by a temporary but significant elevation of thymic Tregs, which may be a homeostatic mechanism in response to the transient but significant Tregs depletion.26

CD25 is also known to be expressed on activated CD8+ T eff cells,15 and this has historically raised skepticism over the use of CD25-targeted therapies to deplete Tregs as these therapies could also adversely affect CD25+ Teffs. Our data show that the CD8+ T eff population was not negatively affected by treatment with CD25-ADC, and, in fact, there was an increase in the number of tumor-infiltrating CD8+ T effs following depletion of intratumoral Tregs. We believe this is due to the significantly lower expression of CD25 on the CD8+ Teff population vs the Tregs population,27 coupled with the limited systemic exposure to CD25-ADC. The lack of CD25-ADC antitumor activity in the absence of CD8+ T cells indicates that CD25-ADC antitumor activity is CD8+ T cell dependent and reinforces the observations from the immunoprofiling studies that overall T eff responses were not negatively impacted by CD25-ADC. Moreover, in both MC38 and CT26 models, tumor-free mice that had previously received CD25-ADC alone or in combination with anti-PD-1 did not develop new tumors when rechallenged, indicating development of tumor-specific immunity.

An additional mode of action that can contribute to CD25-ADC’s potent antitumor activity is the ability of PBD-based ADCs to induce immunogenic cell death (ICD) and to synergize with immuno-oncology therapies in vivo. PBD-induced apoptotic cells trigger ICD signals that stimulate the immune system and allow synergistic activity of PBD-based ADCs with immuno-oncology drugs.28 Moreover, bystander killing of neighboring CD25-negative tumor cells, following release of PBD warhead by dying Tregs, could also contribute to the antitumor activity of CD25-ADC, in line with previous data showing that camidanlumab tesirine and other tesirine-based ADCs induce bystander killing.4 5 However, whether bystander killing contributes to the overall antitumor response of CD25-ADC remains to be determined as intratumoral T eff levels were not affected by CD25-ADC, and local release of warhead could have also killed this cell population. Moreover, when CD8+ Teffs were depleted, CD25-ADC did not have any significant antitumor activity, indicating that bystander killing is unlikely to be the main mediator of CD25-ADC antitumor activity.

Figure 7  T-cell dynamic study in non-tumor-bearing mice. Effect of CD25-ADC on the percentage of Tregs and Teff levels in non-tumor-bearing mice. Female C57BL/6 mice were injected i.p. with vehicle, CD25-ADC (0.5 mg/kg), or isotype control ADC (0.5 mg/kg) on day 0. (A) Spleen, (B) lymph node, and (C) thymus were collected 4 hours post dose, and 6, 13, and 20 days post dose for T-cell immune profiling. Levels of Tregs, CD8+ T, and conventional CD4+ T cells in spleen, lymph nodes, and thymus are presented as % of CD45 cells±SEM over time.
The modest antitumor activity of a high dose of the non-binding control ADC observed in the MC38 model is not uncommon in preclinical studies with ADCs, and it does not invalidate the potent and CD25-mediated activity of CD25-ADC. Several explanations have been put forward to explain this phenomenon. For example, tumor-associated macrophages can process non-targeting human IgG1 ADCs via FcγR, allowing drug release. Moreover, warhead may have been released from the non-binding control ADC by extracellular enzymes such as cathepsin B released from tumor-associated macrophages. Notably, antitumor activity of the non-binding control ADC was considerably inferior to CD25-ADC at same dose and it was model dependent, as the isotype control ADC had no significant antitumor activity in the CT26 model. The observation that TFS mice previously treated with the control ADCs developed antitumor immunity reinforces the hypothesis that PBD-driven ICD may be an important comediatory of CD25-ADC antitumor activity. Because the control ADCs do not directly target and kill CD25-positive Treg (online supplementary figure S10), it is plausible that the released warhead acts by bystander killing of tumor cells, followed by triggering of ICD with subsequent recruitment of Treg cells. An interesting follow-up study to help dissect the contribution of each individual mechanism will be to generate non-cleavable versions of the ADCs, as the use of non-cleavable linkers will eliminate bystander cell killing caused by the released warhead. It has previously been shown that using CD25-specific immunotoxins and/or using Fc-mutated CD25-specific mAbs to deplete CD25-positive Treg is an effective way to achieve antitumor activity in established tumor models. Our molecule differs from previously published approaches as it employs a highly potent DNA damaging toxin known to induce ICD. Consequently, the dose levels required to elicit activity are modest, and exposure to the ADC is, therefore, only a few days after systemic administration. This ensures that Treg depletion is only transient and alleviates concerns of prolonged systemic Treg depletion, potentially leading to serious autoimmune side effects. Moreover, because of its unique mode of action, low doses of CD25-ADC could be tested as preconditioning for other immunotherapies such as additional checkpoint inhibitors, immune agonist antibodies targeting costimulatory receptors or cancer vaccines.

CONCLUSIONS

ADCs are currently being predominantly used to directly target antigen-positive tumor cells. Here, we provide proof of concept for a novel application of ADCs as immuno-therapeutic agents. In particular, this study demonstrates a novel immuno-oncology application of a surrogate of the camidanlumab tesirine ADC either as monotherapy or in combination with PD-1 inhibitors, when used at a suboptimal dose, to treat established CD25-negative solid tumors that exhibit tumor infiltration of CD25-positive Treg. Further work is needed to better understand the interplay of Treg and T eff and to investigate if other immune cells such as macrophages and conventional dendritic cells contribute to CD25-ADC antitumor activity, as in the case of previously reported antitumor activity of an anti-CD25 immunotoxin (2E4-PE38). Translation of these preclinical data in the clinic is now being investigated in a phase I trial (NCT03621982) evaluating the efficacy of camidanlumab tesirine in patients with selected advanced solid tumors.

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Contributors FZ designed the study, analyzed and interpreted data, and was the primary writer of the paper. KH and SC designed part of the studies, and analyzed and interpreted data. FB and BV performed experiments and analyzed data. PnHB designed the study, interpreted data, and edited the paper. All authors contributed to development of the final manuscript.

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