Research paper

DuoBody-CD3xCD20 induces potent T-cell-mediated killing of malignant B cells in preclinical models and provides opportunities for subcutaneous dosing

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ABSTRACT

Background: DuoBody-CD3xCD20 (GEN3013) is a full-length human IgG1 bispecific antibody (bsAb) recognizing CD3 and CD20, generated by controlled Fab-arm exchange. Its Fc domain was silenced by introduction of mutations L234F L235E D265A.

Methods: T-cell activation and T-cell-mediated cytotoxicity were measured by flow cytometry following coculture with tumour cells. Anti-tumour activity of DuoBody-CD3xCD20 was assessed in humanized mouse models in vivo. Non-clinical safety studies were performed in cynomolgus monkeys.

Findings: DuoBody-CD3xCD20 induced highly potent T-cell activation and T-cell-mediated cytotoxicity towards malignant B cells in vitro. Comparison of DuoBody-CD3xCD20 to CD3 bsAb targeting alternative B-cell antigens, or to CD3xCD20 bsAb generated using alternative CD20 Ab, emphasized its exceptional potency. In vitro comparison with other CD3xCD20 bsAbs in clinical development showed that DuoBody-CD3xCD20 was significantly more potent than three other bsAbs with single CD3 and CD20 binding regions and equally potent as a bsAb with a single CD3 and two CD20 binding regions. DuoBody-CD3xCD20 showed promising anti-tumour activity in vivo, also in the presence of excess levels of a CD20 Ab that competes for binding. In cynomolgus monkeys, DuoBody-CD3xCD20 demonstrated profound and long-lasting B-cell depletion from peripheral blood and lymphoid organs, which was comparable after subcutaneous and intravenous administration. Peak plasma levels of DuoBody-CD3xCD20 were lower and delayed after subcutaneous administration, which was associated with a reduction in plasma cytokine levels compared to intravenous administration, while bioavailability was comparable.

Interpretation: Based on these preclinical studies, a clinical trial was initiated to assess the clinical safety of subcutaneous DuoBody-CD3xCD20 in patients with B-cell malignancies.

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1. Introduction

Targeting of CD20-expressing cells with CD20-specific monoclonal antibodies (mAb) has shown to be highly successful for the treatment of B-cell malignancies and specific autoimmune diseases.

Rituximab was the first CD20-specific mAb to obtain regulatory approval and has rapidly become a key part of the standard of care for both non-Hodgkin’s lymphoma and B-cell chronic lymphocytic leukaemia patients. Rituximab is also used in CD20-positive B-lineage acute lymphoblastic leukaemia (B-ALL) [1,2]. Nevertheless, disease relapse or recurrence occurs in many patients [3]. Since resistance of B-cell malignancies to CD20-targeting mAb is rarely caused by loss of CD20 expression or CD20 mutations [4,5], CD20 remains an attractive tumour target, due to its restricted normal tissue expression and prevalent expression on malignant B cells. However, alternative and more powerful targeting strategies are warranted.
Bispecific antibodies (bsAb) specifically recognizing the T-cell antigen CD3 in addition to a tumour-associated antigen (TAA), CD3 bsAb, have begun to deliver their promise in treatment of cancer. CD3 bsAb are able to efficiently induce cytotoxic synapse formation and target cell kill independent of the T-cell receptor (TCR) specificity. BSAb that recognize CD3 and EpCAM (withdrawn in Jun-2017) and blinatumomab, recognizing CD3 and CD19, have been approved and marketed; catumaxomab, recognizing CD3 and EpCAM, was approved in both formats, for example by prolonging plasma half-life and improving manufacturability, as well as in expanding the arsenal of TAA that can be targeted with bsAb.

We previously developed a method to generate whole IgG1 bsAb using controlled Fab-arm exchange (cFAE); termed DuoBody® technology. Here, two parental IgG1 mAb, each with a matched single point mutation in the IgG Fc region, reorganize into full-length bispecific IgG1 molecules upon exposure to controlled manufacturing conditions [7,8]. The cFAE process was shown to be an easy and robust method to generate highly stable bsAb with superior yield compared to other bsAb platforms, both at research and manufacturing scale [9]. Importantly, the method provides the opportunity to generate and screen large and diverse panels of bsAb, to identify bsAb with optimal functional characteristics.

Evidence before this study

CD3 targeting with monoclonal antibodies (mAb) has been shown to be highly successful for the treatment of B-cell malignancies. Since resistance to CD3-targeting mAb is rarely caused by loss of CD20 expression or CD20 mutations, CD20 remains an attractive tumour target, due to its restricted normal tissue expression and prevalent expression in B-cell malignancies. T-cell redirecting CD3 bispecific antibodies (bsAb) have begun to deliver their promise in the treatment of cancer. CD3 bsAb trigger T cell kill of tumour cells expressing the target antigen, irrespective of T-cell receptor specificity. Generation of bispecific antibodies using controlled Fab-arm exchange (cFAE) has been shown to be very efficient (yield >95%) and results in stable, fully functional IgG1 bsAb.

Added value of this study

The preclinical studies described here show that DuoBody-CD3xCD20, a CD3 bsAb generated by cFAE, induced T-cell activation and T-cell-mediated cytotoxicity towards CD20-expressing malignant B cells with high potency, which could not be explained by CD20 expression levels or binding characteristics of its CD20-binding Fab arm. In comparison with other CD3xCD20 bsAb in clinical development, DuoBody-CD3xCD20 showed equal or superior potency in functional assays in vitro. Furthermore, subcutaneous administration of DuoBody-CD3xCD20 induced profound, long-lasting and reversible B-cell depletion from peripheral blood and lymph nodes in cynomolgus monkeys and resulted in reduced plasma cytokine levels compared to intravenous administration, while bioavailability was comparable.

Implications of all the available evidence

The present evidence suggests that DuoBody-CD3xCD20 is a promising agent for treatment of B-cell malignancies. The subcutaneous administration route may provide a method to reduce peak cytokine levels in patients, in addition to a reduced treatment burden for patients and improved resource utilization at the treatment facility. Convenience is increasingly important to patients and health care professionals and systems. Based on results of the preclinical studies described here, a clinical trial to assess the safety and preliminary efficacy of subcutaneous DuoBody-CD3xCD20 (GEN3013) in patients with relapsed, progressive or refractory B-cell lymphoma is currently enrolling patients (GCT3013-01, NCT03625037).

2. Materials & methods

2.1. Antibodies and cell lines

Commercially available mAb and cell lines used in the experiments are listed in Suppl. Tables 1 and 2, respectively. IgG1 mAb that are listed in Suppl. Table 3 were recombinantly produced [12], with an F405L mutation in all CD3 mAb, a K409R mutation in all TAA-specific mAb [7] and FEA (L234F, L235E and D265A) mutations in both. BsAb were generated by cFAE [8], in some cases using the HIV-1 gp120-specific mAb IgG1-b12 [13] to generate bsAb with one non-binding arm. Binding of the bsAb to their antigens was determined by flow cytometry as described (Suppl. data and methods). Four other CD3xCD20 bsAb were produced based on variable and constant region sequences available from published patent applications and literature (bsAb1: WO2014047231, WO2009018411 [Regeneron Pharmaceuticals]; bsAb2: US20170349657 A1, US20140370013 [Xencor Inc.]; bsAb3: US20170349657 A1, US20140370013 [Xencor Inc.]; bsAb4: US20160075785 A1 [Hoffmann-La Roche]). Binding of these bsAb to their targets, CD3 on healthy donor T cells and CD20 on Daudi cells, was confirmed (data not shown).

2.2. Antibody binding assay

Binding of bsAb to cell surface-expressed antigens was determined by flow cytometry as described [15], using an R-phycocerythrin (R-PE)-labelled detection Ab (Suppl. Table 1) to detect primary Ab binding. Binding was detected using an iQue screener (InteliCyt Corporation, USA), a BD LSRFortessa or a BD Canto II flow cytometer (BD Biosciences, Europe).

Simultaneous binding of bsAb to B and T cells was measured as follows: Heparinized whole blood from a healthy donor was incubated with Ab at 37 °C for 2 h. Cells were washed twice and incubated with mAb specific for CD4 or CD8 and CD19 (Suppl. Table 1) at 4 °C for 30 min. Erythrocytes were lysed by addition of erythrocyte lysis buffer (10 mM KHCO₃/0.01 mM EDTA/155 mM NH₄Cl dissolved in dH₂O). Samples were analysed by flow cytometry, using a BD Canto II (BD Biosciences, Europe). The number of CD4⁺CD19⁺ or CD8⁺CD19⁺ double-positive events, indicative of simultaneous binding of DuoBody-CD3xCD20 to human T and B cells, was quantified by CD4/CD19 and CD8/CD19 quadrant analysis, after measuring a fixed sample volume.
2.3. Determination of target expression levels (QIFi)

Target expression, in terms of specific antibody-binding capacity (sABC), was measured using the QIFi kit (DAKO) according to manufacturer’s instructions. Ab used in these experiments are listed in Suppl. Table 1.

2.4. In vitro T-cell assays

Buffy coats from healthy donors (Sanquin, Amsterdam) were used to isolate either peripheral blood mononuclear cells (PBMC) using Lymphocyte separation medium (Lonzas, Basel, Switzerland) or pan-T cells, CD4+ T cells or CD8+ T cells by negative selection using RosetteSep™ Enrichment cocktail kits (Stem Cell Technologies, Vancouver, Canada). CD3 bsAb-induced T-cell-mediated cytotoxicity was determined with a chromium release, alamarBlue or flow cytometric assay. Chromium-release assays with isolated T cells and target cells were performed as described [16]. E:T ratios tested are indicated in the Figure legends. Specific lysis was calculated as: % specific lysis = ((CPM sample – CPM background lysis)/(CPM maximal lysis – CPM background lysis)) x 100, where CPM refers to counts per minute. A release of 0.52 x [length] x [width]^2. Blood leukocyte cell counts, and activation status were determined by flow cytometry (Ab listed in Suppl. Table 1). Sample sizes for in vivo experiments in mice were chosen based on power analysis, the growth characteristics of the tumour model (homogenous vs. heterogeneous outgrowth in untreated animals) and the expected efficacy of the drug. Using historical data, growth of five different human xenograft models was compared after SC implantation in mice. From this comparison, the average coefficient of variance (CV) was calculated to be 31% at a mean tumour size of 883 mm^2, at the last day that all groups in the study were intact. Analysis using GraphPad StatMate indicated that group sizes of n = 10, n = 8 and n = 6 indicated a power of 80% for a reduction in tumour growth of 41.4% (n = 10), 47% (n = 8) and 55.9% (n = 6) to be detected with a significance of 0.05. For co-engraftment of Raji cells and PBMC from one donor, sample sizes of n = 4 (Fig. 4A) and n = 5 (Fig. 4B) were selected. Since two PBMC donors were used in these experiments, a total of n = 8 (A) and n = 10 (B) mice was used per group. For Fig. 4C-G, a sample size of n = 8 was selected.

For PK and clearance analysis of Ab, an analysis was performed using five tests with a total of 80 mice, in which five different Ab were evaluated in different mouse strains. Power analysis indicated that with a group size of n = 3, differences of 0.2 in log-clearance could be demonstrated with 80% power. This means that differences can be determined if plasma clearance rate of one Ab is 66% faster or slower than that of another antibody. Practice has shown that changes in Ab structure easily lead to large differences in clearance rate. Therefore, group sizes of n = 3 were considered sufficient (Suppl. Fig. 1). CB17/lcr-Pkrdcsdci/lcrlocCrl, female mice were used, 10–11 weeks at the time of Ab injection. At the time of Ab injection for PK studies, animals were picked randomly from the cages and treatment groups were randomized over all cages.

All animal experiments were performed in compliance with European directives (2010/63/EU) and approved by the local Ethical committees. Animals were housed and handled in accordance with good animal practice as defined by FELASA, in an AAALAC and ISO 9001:2000 accredited animal facility (GDL). For experiments performed at Axenis, animal experimental procedures have received an approval form the local animal Ethical Committee (CETEA 89), Institut Pasteur de Paris.

2.7. Non-clinical safety studies in cynomolgus monkeys

Non-clinical safety studies were conducted at Charles River Laboratories (Tranent, UK) in accordance with the European Convention for the Protection of Vertebrate Animals Used for Experimental and Other Scientific Purposes (Council of Europe), under control of the UK Home Office. This facility is a GLP compliant facility and the practices and procedures adopted during the conduct were consistent with the OECD Principles of Good Laboratory Practice as incorporated into the United Kingdom Statutory Instrument for GLP.
Fig. 1. Functional characterization of DuoBody-CD3xCD20 in vitro. A: Isolated T cells were incubated with DuoBody-CD3xCD20 and Daudi cells (E:T ratio 2:1). Percentages of CD69-positive CD4+ (left) and CD8+ T cells (middle) were determined by flow cytometry at the indicated time points. Data for one representative donor out of four donors is shown. Graphs show the mean ± standard error of the mean (SEM) of duplicate wells. The right panel shows percentages of CD69-positive CD4+ T cells. Data shown are pooled data from 6–8 donors for T cells treated with 3.3 ng/mL DuoBody-CD3xCD20. Boxes extend from 25th to 75th percentiles. Whiskers are drawn down to minimum and up to maximum percentages. Median percentages are indicated. A significant linear trend was observed in time (one-way ANOVA with post-test for linear trend). B: Isolated T cells were co-cultured with Daudi cells (E:T ratio 1:1) in the presence of 1 μg/mL DuoBody-CD3xCD20 for 24 h. Images were taken at a 60x magnification (plus extra optical magnification of the left pictures) and are representative of three experiments. C: Isolated CD4+ (left) or CD8+ T cells (right) were incubated with DuoBody-CD3xCD20 and chromium-labelled Daudi cells (E:T ratio 20:1; this relatively high E:T ratio was chosen to make sure that a signal could be measured for CD4+ T cells and at early time points) for the indicated time periods. Data shown are mean percentages of Daudi cell lysis ± SEM of triplicates. Data from one representative donor out of six donors tested is shown. D: Isolated CD4+ or CD8+ T cells were incubated with DuoBody-CD3xCD20 or bsAb-CD3xctrl and chromium-labelled Daudi cells (E:T ratio 8:1) for 24 h. Data shown are mean percentages of Daudi cell lysis ± SEM of triplicates. Data from one representative experiment out of 3 performed is shown. E: PBMC were incubated with indicated Ab for 72 h. Percentages of CD69-positive CD4+ (left) and CD8+ T cells (middle) and the remaining number of intact endogenous B cells (right) were determined by flow cytometry. Data shown are from one representative experiment out of four performed and are mean percentages ± SEM of duplicate wells. F: Isolated T cells were incubated with DuoBody-CD3xCD20, bsAb-CD3xctrl or bsAb-ctrlxCD20 and indicated B-cell lymphoma cell lines (E:T ratio 2:1) for 48 h. The remaining number of intact lymphoma cells (right) were determined by flow cytometry. Data shown for each cell line are from one representative donor (for number of donors tested see Table 1) and are mean percentages ± SEM of duplicate wells.
Cynomolgus monkeys (purpose-bred cynomolgus monkeys [Macaca fascicularis, of Mauritian origin, Fig. 5A-D: 26–29 months old at allocation to the study; 2.65–3.03 kg; 2.71–3.06 kg at dosing; Fig. 5 E-F: 4–6 years at the time of dosing, 3.2–7.9 kg]) received DuoBody-CD3xCD20 either via an IV infusion or a SC injection. Blood samples and lymph node biopsies were taken to determine B-cell counts and concentrations of Ab and cytokines. Single-cell suspensions were prepared from lymph node biopsies using the Media- machine System (Becton Dickinson). The number of B cells in whole blood samples or lymph node biopsies was determined by flow cytometry using a two-laser five-color Beckman Coulter FC500 or a BD LSRFortesa, gating for CD19+ cells (antibodies listed in Suppl. Table 1).

Plasma was isolated from whole blood by centrifugation (1500g, 4 °C, 10 min). DuoBody-CD3xCD20 concentrations were determined with a Single Molecule Counting ImmunoAssay method (PRA, Assen, The Netherlands). Cytokine concentrations were measured using the Milliplex MAP NHP Cytokine Magnetic Bead Panel (Millipore) on a Bio-Plex 200 reader (BioRad).

The numbers of cynomolgus monkeys evaluated in toxicity studies were the minimum required to properly characterise the effects of the test article and to avoid an unnecessary number of animals to accomplish the study objectives. Fig. 5A-D: n = 2 female monkeys per group; Fig. 5E-F: n = 6 monkeys (3 males and 3 females) per group. For allocation to dose groups, animals were randomly allocated to groups and subsequently assigned individual animal numbers. Individual body weights were reviewed to ensure a balanced distribution across groups. For quantitative numerical data where $n \leq 3$, no statistical analyses were applied. These data were evaluated by comparing the individual data in a descriptive manner.

2.8. Data analysis and statistics

Dose-response curves were generated using four-parameter nonlinear regression analysis (GraphPad Prism 7.02 software [La Jolla, CA, USA]; equation log[agonist] vs. response – variable slope). Half maximal effective concentration (EC$_{50}$) values were derived from these curves. In in vivo experiments, differences in tumour growth or frequency of cellular subsets were analysed in treated versus control mice by Mann-Whitney two-tailed test or by a Kruskal-Wallis analysis of variance [ANOVA] test followed by Dunn’s multiple comparison post-test, as indicated.

3. Results

3.1. Selection of mutations to obtain CD3 bsAb with a silenced Fc region

A fully functional IgG1 Fc region that is capable of inducing Fc-mediated effector functions is not desirable for CD3 bsAb, because Ab-dependent Fc gamma receptor (FcyR)-mediated cross-linking of CD3 on T cells may induce TAA-independent T-cell activation and cytotoxicity. In addition, Ab-dependent cell-mediated cytotoxicity (ADCC) and complement-dependent cytotoxicity (CDC), may lead to cytotoxicity towards T cells. Furthermore, we observed ineffective T-cell redirection by an Fc-effector function-competent (active) CD3 bsAb in a syngeneic mouse tumour model [20]. We set out to identify mutations to silence Fc-mediated effector functions, that were compatible with generation of (CD3) bsAb using FcAE. Point mutations N297Q, L234F, L235E, D265A, P331S [21–27], alone or in combination, were introduced in CD3-specific IgG1 mAb or bsAb, and tested for their capacity to prevent FcyR and complement binding, while retaining neonatal FcR (FcRn) binding.

The combination of three mutations L234F, L235E and D265A (FEA) provided the most favourable characteristics in vitro and in vivo: CD3-specific mAb carrying the FEA mutations showed normal antigen binding (Suppl. Fig. 1A), did not induce Ab-dependent FcR-mediated T-cell activation or proliferation (Suppl. Fig.1B-C) and did not induce untargeted cytotoxicity in tumour cells (Suppl. Fig. 1D). Similarly, CD3 bsAb carrying the FEA mutations induced cytotoxicity in HER2-expressing tumour cells when the TAA-specific Fab-arm recognized HER2 (Suppl. Fig. 1E-F). CD3 mAb bearing the FEA mutations were unable to bind membrane expressed FcYRII (Fig. 1G) and immobilized CD3 mAb with the FEA mutations had lost the capacity to bind FcYRII, FcYRII (data not shown) and complement component C1q (Suppl. Fig. 1H). CD3 mAb with the FEA mutations showed normal binding to human and mouse FcRn in vitro (data not shown) and demonstrated a normal plasma half-life in vivo (Suppl. Fig.1I-J). Finally, introduction of the FEA mutations did not have impact on the efficiency of FcAE (data not shown).

3.2. Functional characterization of DuoBody-CD3xCD20 in vitro

DuoBody-CD3xCD20 was generated by FcAE of a humanized variant of the mouse anti-human CD3 mAb SP34 [28] and the human anti-CD20 mAb IgG1-7D8 [10,11], carrying the matched DuoBody mutations F405L and K409R (respectively) and the FEA mutations.

DuoBody-CD3xCD20 induced dose-dependent activation of CD4+ and CD8+ T cells upon incubation in the presence of CD20-expressing Daudi cells, as assessed by upregulation of CD69 (Fig. 1A) and CD25 (Suppl. Fig. 2A), with comparable kinetics for CD4+ and CD8+ T cells. The capacity of DuoBody-CD3xCD20 to activate T cells was confirmed by confocal microscopy. In addition to the presence of CD69 on CD4+ and CD8+ T cells, perforin, a component of cytotoxic granules, was detected in T cells that were near Daudi cells (Fig. 1B). DuoBody-CD3xCD20 induced potent dose-dependent T-cell-mediated cytotoxicity towards Daudi cells. In contrast to the kinetics of T-cell activation, the kinetics of DuoBody-CD3xCD20-induced tumour cell kill were different for CD4+ and CD8+ T cells. While CD8+ T-cell-mediated cytotoxicity was already measurable after 3 h, CD4+ T-cell-mediated cytotoxicity could not be detected at that time point (Fig. 1C). Furthermore, at the same effector to target cell (E:T) ratio, CD8+ T cells induced more efficient target cell lysis than CD4+ T cells (Fig. 1D).

Selective targeting of CD20-positive cells by DuoBody-CD3xCD20 was confirmed in healthy donor PBMC, that contain both CD3-expressing effecter cells and CD20-expressing target cells. Flow cytometry experiments demonstrated formation of CD4/CD19 and CD8/CD19 double-positive events, indicative of T/B-cell doublets, in the presence of DuoBody-CD3xCD20 (Suppl. Fig. 2B). This demonstrates that DuoBody-CD3xCD20 can simultaneously bind B and T cells. Moreover, DuoBody-CD3xCD20 induced activation of both CD4+ and CD8+ T cells and subsequent lysis of normal B cells in PBMC, as assessed by flow cytometry after 72 h of incubation (Table 1, Fig. 1E). Both T-cell activation and cytotoxicity strictly required simultaneous binding of CD3 and CD20, since bsAb controls bsAb-CD3xctrl and bsAb-ctrlxCD20 did not induce T-cell activation or B-cell depletion (Fig. 1E) in PBMC.

The capacity of DuoBody-CD3xCD20 to induce cytotoxicity in malignant B cells was further analysed in a panel of cell lines derived from various types of B-cell lymphoma, with CD20 cell surface expression ranging from 33,000–350,000 specific antibody-binding capacity (sABC). DuoBody-CD3xCD20 induced potent cytotoxicity in all cell lines tested, with EC$_{50}$ values in the low picomolar range. There was no relation between antigen expression and cytotoxicity, suggesting that antigen expression of 33,000 CD20 molecules per cell is sufficient to achieve maximal T-cell-mediated cytotoxicity in the presence of DuoBody-CD3xCD20 (Table 1, Fig. 1F).

In summary, DuoBody-CD3xCD20 induced highly potent kill of CD20-positive tumour cells, which was mediated by both CD4+ and CD8+ T cells. T-cell activation and cytotoxic activity were strictly dependent on binding of both CD3 and CD20.
Average EC50 values of DuoBody-CD3xCD20-induced cytotoxicity. Averages of CD20 expression levels and EC50 values of DuoBody-CD3xCD20-induced T-cell-mediated cytotoxicity and T-cell activation (% CD69 positive cells) measurements for each B-cell lymphoma cell line tested, calculated from multiple experiments. CD20 expression levels were determined by quantitative flow cytometry assays at E:T ratio 2:1 (% kill; % CD69 positive after 48-h incubation). (GCB = germinal centre B-cell diffuse large B-cell lymphoma, ABC = activated B-cell diffuse large B-cell lymphoma, BL = Burkitt's lymphoma, n = number of donors).

Table 1

| Name       | Lymphoma Type | Average CD20 expression level (ng/mL) | n tests | Range (x1000) | EC50 (pm) | Average DuoBody-CD3xCD20-induced cytotoxicity (pm) | n donors | Average DuoBody-CD3xCD20-induced CD4+ T cell activation (pm) | n donors | Average DuoBody-CD3xCD20-induced CD8+ T cell activation (pm) | n donors |
|------------|---------------|--------------------------------------|---------|---------------|----------|-------------------------------------------------|----------|-------------------------------------------------|----------|-------------------------------------------------|----------|
| OCI-Ly18  | GCB           | 166 (84)                             | 2       | 36            | 453      | 0.531 [0.813]                                   | 6        | 0.121 [0.300]                                    | 6        | 0.807 [1.300]                                    | 6        |
| RC-K8      | ABC           | 139 (78)                             | 4       | 67.604        | 450.693  | 450.693 [766.193]                               | 4        | 450.693 [766.193]                               | 4        | 450.693 [766.193]                               | 4        |
| OCCM184   | BL            | 33 (0)                               | 8       | 88            | 88       | 0.120 [0.105]                                   | 2        | 0.120 [0.105]                                   | 2        | 0.120 [0.105]                                   | 2        |
| SU-DHL-8  | GCB           | 33 (0)                               | 8       | 88            | 88       | 0.120 [0.105]                                   | 2        | 0.120 [0.105]                                   | 2        | 0.120 [0.105]                                   | 2        |

3.3. Assessment of the contribution of the B-cell-targeting Fab arm to CD3 bsAb efficacy in vitro

The CD20-specific Fab arm of DuoBody-CD3xCD20 was derived from the previously described CD20-specific human IgG1-7D8 [10,11]. To study the contribution of this specific Fab arm to the observed potency of DuoBody-CD3xCD20, we compared the capacity to induce T-cell-mediated cytotoxicity of a CD3 bsAb based on 7D8, with CD3 bsAbs using B-cell-targeting arms derived from mAbs against seven other well-known B-cell membrane molecules: CD22, CD24, CD37, CD70, CD79b, CD138 and human leukocyte antigen (HLA)-DR. CD3 bsAbs targeting the different tumour antigens showed strong variation in cytotoxic capacity, as demonstrated in a 24-h chromium release assay, and cytotoxicity did not correlate with antigen expression levels (Fig. 2A). The CD20-specific CD3 bsAb (generated using a 7D8-derived Fab arm) was among the most potent CD3 bsAbs. Interestingly, none of CD3 bsAbs targeting HLA-DR or CD138, both of which show antigen expression levels comparable to or higher than CD20, were able to induce cytotoxicity, demonstrating that high levels of antigen expression do not guarantee efficacy of CD3 bsAbs.

In a next set of experiments, CD20, in addition to CD37 and CD70, the three antigens that were most efficiently targeted with CD3 bsAbs in the first experiment, were used to assess the contribution of binding characteristics of the TAA-specific Fab arm to CD3 bsAb-induced cytotoxicity. DuoBody-CD3xCD20 was compared with three other CD20-specific CD3 bsAbs and, similarly, four different CD3 bsAbs targeting CD37 and CD70 were generated and compared. The tumour-targeting Fab arms were selected based on diversity in binding epitopes [29,30] and Suppl. Table 4) and target antigen-binding characteristics (EC50 and maximal binding), as assessed by flow cytometry (Fig. 2B).

Binding characteristics of the TAA-binding Fab arm (Fig. 2B) did not predict the capacity of CD3 bsAbs to induce T-cell-mediated cytotoxicity (Fig. 2C) in experiments using Daudi cells as target cells and purified healthy donor T cells as effector cells. Despite the difference in EC50 and maximal binding of CD3xCD202C6 compared to CD3xCD202C6 and CD3xCD202C6, all three bsAbs displayed comparable cytotoxic activity in Daudi cells. DuoBody-CD3xCD20, while demonstrating lower apparent affinity than CD3xCD202C6, demonstrated higher maximum levels of binding and considerably more potent cytotoxicity. The CD37-targeting CD3 bsAbs, CD3xCD37Ab3, CD3xCD37Ab5 and CD3xCD37Ab6, showed considerable differences in maximal antigen binding but induced comparable cytotoxicity. CD3xCD37Ab showed reduced binding, and reduced potency compared to the other CD3xCD37 bsAbs. For the CD70 bsAbs, the potency observed for Daudi cells was considerably lower than for RPMI-8226 cells that were used in the first experiment (Fig. 2A), despite higher expression of CD70 on Daudi cells (RPMI-8226: 3,900 sABC; Daudi: 12,900 sABC). Nonetheless, also for CD70, the differences in potency of CD3 bsAbs could not be predicted based on the binding characteristics.

Taken together, while the CD20-specific Fab arm derived from IgG1-7D8 determines the unique potency of DuoBody-CD3xCD20, neither TAA expression levels nor the TAA binding characteristics could predict the potency of B-cell-specific CD3 bsAbs in vitro.

Finally, the potency of DuoBody-CD3xCD20 in vitro was compared to four other, previously described, CD3xCD20 bsAbs that are currently being evaluated in the clinic (sequences and format specifics were obtained from patent applications). The capacity of these bsAbs to induce T-cell activation (Fig. 3A-B) and T-cell-mediated cytotoxicity (Fig. 3C) was analysed by incubation with isolated T cells and Daudi cells for 48 h. DuoBody-CD3xCD20 was found to be significantly more potent than three IgG1-like bsAbs with a single CD3 and CD20 binding region, and equally potent as a bsAb with a single CD3 and two CD20 binding regions (Fig. 3D).
3.4. DuoBody-CD3xCD20 induced potent anti-tumour activity in humanized mouse models

The anti-tumour activity of DuoBody-CD3xCD20 was assessed in vivo using CD20-positive lymphoma xenograft models in mice engrafted with human immune cells. In a co-engraftment model where NOD-SCID mice were injected SC with a 1:1 mixture of PBMC and Raji B-cell lymphoma cells, followed by IV administration of Ab 2 h later, DuoBody-CD3xCD20 induced a dose-dependent decrease in tumour outgrowth compared to phosphate-buffered saline (PBS) (Fig. 4A). In the same model, we assessed the capacity of DuoBody-CD3xCD20 to exert anti-tumour activity in the presence of circulating...
rituximab (IgG1-RTX), which competes with 7D8 for CD20 binding [31]. To ensure that anti-tumour activity observed in this study could be attributed to DuoBody-CD3xCD20 and not rituximab, Fc-mediated effector functions of IgG1-RTX were silenced by introducing the FEA mutations (IgG1-RTX-FEA). The presence of up to 10 mg/kg IgG1-RTX-FEA did not reduce the potency of 0.05 mg/kg DuoBody-CD3xCD20 in vivo (Fig. 4B). This provides a preclinical indication that patients that still have rituximab in the circulation after failing to respond to a rituximab-containing treatment regimen could still benefit from treatment with DuoBody-CD3xCD20.

The therapeutic potential of DuoBody-CD3xCD20 was assessed in HIS mice, using both an IV and a SC xenograft model [19]. In the IV tumour model, representing the leukemic phase of B-cell malignancies, mice were injected IV with Daudi cells and Ab treatment was initiated 3 days later. Treatment with DuoBody-CD3xCD20, but not the control bsAb-CD3xctrl, resulted in inhibition of tumour outgrowth (Fig. 4C). The more solid manifestation of Non-Hodgkin's lymphoma was mimicked by injecting HIS mice SC with Raji lymphoma cells and tumours were allowed to establish before Ab treatment was initiated. DuoBody-CD3xCD20 induced a profound inhibition of tumour growth at doses of 0.1 mg/kg and 1 mg/kg, whereas bsAb-CD3xctrl did not show anti-tumour activity (Fig. 4D). Interestingly, bsAb-ctrlxCD20 (but not bsAb-CD3xctrl) also induced some anti-tumour activity in this model. In addition to evaluating the anti-tumour activity of DuoBody-CD3xCD20, this HIS-model allowed the opportunity to study depletion of normal human B cells from the circulation. On day 4, 9 and 16, a significant decrease in circulating B cells was observed after treatment with DuoBody-CD3xCD20 (0.1 and 1 mg/kg) and bsAb-ctrlxCD20 (p<0.05) compared to the PBS-treated mice (Fig. 4E). In mice treated with 1 mg/kg DuoBody-CD3xCD20, but not bsAb-ctrlxCD20, a transient upregulation of the T-cell activation marker CD69 was observed in peripheral blood CD4+ and CD8+ T cells at day 4 (Fig. 4F, G).

In summary, DuoBody-CD3xCD20 induced anti-tumour activity in vivo towards malignant B cells at SC sites as well as in the blood circulation. This was accompanied by a transient T-cell activation and was not hampered by the presence of circulating rituximab.

3.5. DuoBody-CD3xCD20 induced potent and long-lasting B-cell depletion in cynomolgus monkeys after IV or SC administration

As part of the non-clinical safety studies, DuoBody-CD3xCD20 was administered to cynomolgus monkeys, the selected toxicology species for which antigen binding characteristics and in vitro pharmacology of DuoBody-CD3xCD20 were comparable to humans (data not shown). A single injection of DuoBody-CD3xCD20, administered IV or SC, induced a dose-dependent depletion of B cells from peripheral blood (Fig. 5A, C) and lymph nodes (Fig. 5B, D). B-cell depletion was reversible at all dose levels, with time to recovery correlating with the treatment dose. Efficiency of B-cell depletion was comparable after SC and IV administration.

Analysis of DuoBody-CD3xCD20 levels in plasma demonstrated different PK profiles for the IV and SC administration routes. Following IV infusion, the peak plasma concentration ($C_{max}$) was reached at the end of the 30-min dosing period, then decrease in a generally bi-phasic manner (Fig. 5E). After SC dosing, $C_{max}$ was not reached until approximately three days after dosing and the plasma levels remained relatively steady up to seven days post dose. Thereafter,
Fig. 4. Anti-tumour activity of DuoBody-CD3xCD20 in vivo in humanized mouse models. A-B: Tumours were induced in NOD-SCID mice by SC injection of a 1:1 mixture of human PBMC and Raji-luc cells at day 0. Immediately after tumour inoculation, mice (5 per group) were treated with a single IV dose of DuoBody-CD3xCD20 (0.5, 0.05, 0.005 mg/kg) (A) or mice (10 per group) were given a single IV dose of IgG1-RTX-FEA (10, 1 or 0.1 mg/kg), followed by IV treatment with 0.05 mg/kg DuoBody-CD3xCD20 4 h later, treatment with DuoBody-CD3xCD20 was repeated on day 7 (B). Data shown are mean tumour volumes per treatment group ± SEM. Differences were analysed by Mann-Whitney two-tailed test versus PBS (A: day 21; B: day 25). Statistically significant differences are indicated as follows: *p < 0.05, **p < 0.005, ***p < 0.0005, ****p < 0.0001 (A-B).

C: Tumours were induced in BRGS-HIS mice by IV injection of Daudi-luc cells at day 0. Mice were treated IV with 1 mg/kg DuoBody-CD3xCD20 on day 3 and day 7. Tumour growth was evaluated by BLI and depicted as average radiance of BLI ± SEM (7 mice per group). Statistically significant differences compared to PBS-treated mice were determined by Kruskal-Wallis ANOVA test followed by Dunn’s multiple comparison post-test (**p < 0.01).

D-G: Tumours were induced in BRGS-HIS mice by SC injection of Raji-luc cells at day 0. Mice were treated IV with 1 mg/kg or 0.1 mg/kg bsAb or vehicle control on days 3, 7, 10, 14, and 17. Tumour growth is shown for each individual mouse, one graph per treatment group (D). Individual (circles) and average (black line) absolute numbers of circulating human CD19⁺ B cells determined by flow cytometry at indicated time points (E). T-cell activation was determined on indicated time points by flow cytometry. Individual (circles) and average (black line) percentages of CD69⁺ cells in CD4⁺ T-cell population (F) and CD8⁺ T-cell population (G) are shown. Statistically significant differences compared to PBS-treated mice at each time point were determined by Kruskal-Wallis ANOVA test with Dunn’s multiple comparisons test (*p < 0.05, **p < 0.01, ***p < 0.001).
concentrations decreased in a monophasic manner up to the end of the four-week sampling period. At doses of 0.1 and 1 mg/kg DuoBody-CD3xCD20, the peak plasma concentration was 7- to 17-fold higher after IV compared to SC dosing. Importantly, the exposure (area under the curve) of a single dose of 0.1 or 1.0 mg/kg was comparable after SC and IV injection.

Injection of DuoBody-CD3xCD20 was associated with a rapid and transient increase in plasma cytokine levels. Interleukin (IL)-2, IL-6, IL-8, IL-10, interferon (IFN)γ, tumour necrosis factor (TNF)α and monocyte chemotactic protein (MCP)-1 reached peak levels within 2–12 h after dosing (Fig. 5F) and had resolved to baseline after 24 h. Peak cytokine levels were considerably lower after SC compared to IV treatment. For IL-6, IL-8 and IFNγ peak cytokine levels were reached somewhat later after SC compared to IV dosing, although for both administration routes plasma levels had returned to baseline at 12–24 h.

In summary, DuoBody-CD3xCD20 induced profound and long-lasting depletion of B cells from peripheral blood and lymph nodes in cynomolgus monkeys, with comparable efficacy between IV and SC administration. Peak plasma levels of DuoBody-CD3xCD20 were reduced after SC administration compared to IV administration and SC administration was associated with reduced plasma cytokine levels.

### 3.6. Discussion

DuoBody-CD3xCD20 (GEN3013) is a new bsAb recognizing CD3 and CD20, that triggers highly potent T-cell-mediated lysis of CD20-expressing cells. DuoBody-CD3xCD20 is a full-length IgG1 bsAb generated by cFAB of a humanized version of mouse anti-human CD3 mAb SP34 and human anti-CD20 mAb 7D8. Fc-mediated effector functions were silenced by the introduction of three amino acid mutations (FEA) in the
Fc region. In vitro, DuoBody-CD3xCD20 induced potent and specific activation and cytotoxicity of both CD4+ and CD8+ T cells. Although activation of CD4+ and CD8+ T cells by DuoBody-CD3xCD20 was detected at similar bsAb concentrations and in a comparable timeframe, CD8+ T cells induced cytotoxic activity faster than CD4+ T cells. This has been described previously [32] and most likely reflects an intrinsic difference between T-cell subtypes; whereas CD8+ T cells are harnessed with granules containing cytotoxic molecules that can be released immediately upon activation, CD4+ T cells are not primarily cytotoxic and first need to produce cytotoxic molecules in order to exert cytotoxic activity [33]. It would be highly interesting to gain further understanding of the contribution of CD4+ and CD8+ T-cell populations in the clinical setting.

Intriguingly, DuoBody-CD3xCD20 induced T-cell activation and cytotoxicity with EC50 values in the low ng/mL (low picomolar) range, whereas EC50 values for binding to CD20 on B cells and CD3 on T cells were roughly 1000-fold higher (μg/mL range). This shows that efficient T-cell-mediated cytotoxicity can be achieved when TAA binding is far from saturated, and is in line with studies demonstrating efficient CD3 bsAb-induced T-cell-mediated killing, even if antigen expression levels are very low [34–37]. This might be promising for treatment of patients with malignancies with lower CD20 expression levels, such as ALL. A low antigen expression threshold for CD3 bsAb is also in agreement with lack of correlation between CD20 expression levels and DuoBody-CD3xCD20-induced T-cell-mediated cytotoxicity, an observation that has previously been described for CD3 bsAb targeting other TAAs [34,36].

The cytotoxic capacity of CD3 bsAb is determined by the characteristics of its two Fab arms and by characteristics of the TAA. However, the exact characteristics determining CD3 bsAb potency are poorly understood. To understand the highly potent cytotoxicity of DuoBody-CD3xCD20, we looked at the relation between the expression of a diverse panel of B-cell antigens, and the cytotoxic activity of CD3 bsAb targeting those B-cells. Differences in expression levels between the different B-cell antigens could not predict the potency of CD3 bsAb, in line with the lack of correlation between antigen expression and CD3 bsAb potency for specific target antigens ([34,36] and the present study). For example, CD3 bsAb targeting HLA-DR and CD138, both of which show expression levels sufficient for cytotoxic activity of CD3 bsAb targeting the other B-cell antigens, did not induce killing of B cells. This lack of cytotoxicity can likely be attributed to other target antigen characteristics. For HLA-DR it can be speculated that the immune synapse formation between B and T cells is disrupted by the bsAb due to steric hindrance. Moreover, it cannot be excluded that lack of cytotoxicity for HLA-DR-specific CD3 bsAb might be caused by expression of HLA-DR on activated T cells [38], causing fratricide [39].

Zooming in on CD3 bsAb targeting CD20, CD37 and CD70, no correlation was observed between antigen binding affinity and (fine) binding domain specificity of CD3 bsAb and cytotoxic activity. This was unexpected, since previous studies demonstrated a correlation between CD3 bsAb efficacy and localization of the binding epitope on the TAA [40,41]. While the CD20, CD37 and CD70 bsAbs studied here showed diversity in binding epitopes, the panel of CD3 bsAbs explored for each antigen was small and more detailed studies may provide better insight in the binding epitope requirements for CD3 bsAb targeting these antigens. In addition to the characteristics studied here, other factors may contribute to CD3 bsAb potency, including the bsAb format [42], and TAA characteristics such as molecular size [41], flexibility [40], mobility [43], macromolecular organization [43], accessibility, ease of formation of an immunological synapse, and expression of costimulatory and coinhibitory receptors on the tumour cells, making it nearly impossible to rationalize target antigen and epitope choices. In summary, the high potency of DuoBody-CD3xCD20 in T-cell-mediated targeting of B cells, which was also illustrated by the comparison with four other CD3xCD20 bsAbs that are in clinical development, could not have been predicted based on the known characteristics of the 7D8 Ab clone and CD20.

DuoBody-CD3xCD20 showed potent anti-tumour activity in lymphoma xenograft models in humanized mice, also in the presence of a large excess of circulating rituximab, which competes with 7D8 for CD20 binding [31]. Again, this illustrates that probably only a very small amount of available antigen is required for CD3 bsAb to exert optimal activity. From a clinical perspective, efficacy of DuoBody-CD3xCD20 in the presence of rituximab may indicate that patients that still have CD20 mAb in the circulation when they are no longer responding to CD20 mAb-containing treatments, could be eligible for treatment with DuoBody-CD3xCD20. Surprisingly, we also observed some in vivo anti-tumour activity of a control bsAb containing the CD20-binding Fab arm and a non-binding Fab-arm instead of the CD3-binding Fab arm (bsAb-ctrlxCD20). This is remarkable, because of the inert Fc region, but a similar effect was observed using a rituximab variant carrying the FEA mutations. Any residual interaction of human IgG1 molecules carrying the FEA mutations with mouse FcγR, which has been shown to be the dominant mouse FcγR in antibody-mediated anti-tumour activity in vivo [44], was excluded (Genmab, unpublished data). Although the mechanism behind this CD20-specific Fab-mediated effect on tumour growth needs further investigation, it may contribute to the potent anti-tumour activity of DuoBody-CD3xCD20.

Blinatumomab, the only FDA-approved CD3 bsAb, is a fragment-based bsAb targeting CD19, currently approved for the treatment of B-cell precursor ALL. Because of the lack of an Fc region, the half-life of blinatumomab in humans is short, necessitating continuous IV infusion [45]. DuoBody-CD3xCD20 has a regular IgG1 structure and biochemical characteristics typical of a human IgG1 antibody, including a relatively long plasma half-life. The Fc region of DuoBody-CD3xCD20 contains three mutations (L234F, L235E, D265A) to abrogate FcγR and complement binding, thereby preventing the induction of T-cell activation through antibody-mediated, FcγR-dependent CD3 crosslinking and ensuring that T-cell activation is strictly dependent on simultaneous binding of CD3 and CD20. Binding to FcRn is not affected by the mutations. Neurotoxicities, as observed for blinatumomab, have been described for other CD19-targeting T-cell therapies and are thought to be class-related [45]. It would be interesting to see comparison of the clinical safety profile of blinatumomab and DuoBody-CD3xCD20, but at present insufficient clinical data is available.

In nonclinical safety studies in cynomolgus monkeys, DuoBody-CD3xCD20 induced profound depletion of B cells from both peripheral blood and lymph nodes, which was comparable after IV and SC administration. While bioavailability was comparable between the administration routes, peak plasma levels were lower and delayed after SC administration. Importantly, peak cytokine levels were consistently lower after SC administration. In the clinic, cytokine release and associated adverse events are an important concern for CD3 bsAb, therefore the SC administration route could provide a method to reduce cytokine release in patients.

In conclusion, DuoBody-CD3xCD20 (GEN3013) is a new bsAb that showed specific and highly potent preclinical anti-tumour activity in lymphoma models in vitro and in vivo. DuoBody-CD3xCD20 induced profound depletion of B cells from peripheral blood and lymphoid organs in cynomolgus monkeys upon IV or SC administration, with lower levels of plasma cytokines observed after SC compared to IV administration. These preclinical studies provided the rationale to assess the clinical safety and efficacy of subcutaneous administration of DuoBody-CD3xCD20 (GEN3013) in patients with relapsed, progressive or refractory B-cell lymphoma (GCT3013-01, NCT03625037).

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Declaration of competing interest

(former) Genmab employees own Genmab warrants and/or stock.
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Supplementary materials

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