Increased in vivo effector function of human IgG4 isotype antibodies through afucosylation

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ABSTRACT
For some antibodies intended for use as human therapeutics, reduced effector function is desired to avoid toxicities that might be associated with depletion of target cells. Since effector function(s), including antibody-dependent cell-mediated cytotoxicity (ADCC), require the Fc portion to be glycosylated, reduced ADCC activity antibodies can be obtained through aglycosylation of the human IgG1 isotype. An alternative is to switch to an IgG4 isotype in which the glycosylated antibody is known to have reduced effector function relative to glycosylated IgG1 antibody. ADCC activity of glycosylated IgG1 antibodies is sensitive to the fucosylation status of the Fc glycan, with both in vitro and in vivo ADCC activity increased upon fucose removal (“afucosylation”). The effect of afucosylation on activity of IgG4 antibodies is less well characterized, but it has been shown to increase the in vitro ADCC activity of an anti-CD20 antibody. Here, we show that both in vitro and in vivo activity of anti-CD20 IgG4 isotype antibodies is increased via afucosylation. Using blends of material made in Chinese hamster ovary (CHO) and Fut8KO-CHO cells, we show that ADCC activity of an IgG4 version of an anti-human CD20 antibody is directly proportional to the fucose content. In mice transgenic for human FcγRIIIa, afucosylation of an IgG4 anti-mouse CD20 antibody increases the B cell depletion activity to a level approaching that of the mIgG2a antibody.

Introduction
The crystallizable fragment (Fc) imparts effector function to full-length antibodies. These activities include long serum half-life due to binding of the Fc portion to the recycling FcRn receptor, activation of complement-dependent cytotoxicity (CDC) initiated through C1q binding to the Fc, and antibody-dependent cell-mediated cytotoxicity (ADCC), which occurs through engagement of Fcγ receptors on effector cells. Notably, binding of antibodies to FcγRIIa on natural killer (NK) cells enables the immune system to target foreign cells for destruction. ADCC can be employed in therapeutic agents to remove pathologic cells including malignant tumor cells. For example, non-Hodgkin’s lymphoma (NHL) is effectively treated with a combination of a chemotherapy regimen and administration of the anti-CD20 antibody rituximab. Rituximab binds CD20 on normal and cancerous B cells, targeting them for destruction. Response to rituximab therapy is linked to a polymorphism in the FcγRIIIa receptor whereby patients having the higher Fc-affinity allele Val158, as compared with Phe158, show a more robust clinical response. These observations suggest that ADCC activity is required for the full effectiveness of rituximab therapy in patients afflicted with NHL. These results have prompted the development of antibodies such as obinutuzumab (Gazyva) that are engineered to have increased ADCC activity.

Both ADCC and CDC require glycosylation of the Fc residues Asn297 whereas FcRn interaction does not, and thus aglycosylated antibodies retain long serum half-life. ADCC activity is modulated by glycan structure and removal of the penultimate fucose (“afucosylation”) can increase FcγRIIIa affinity and ADCC activity. In addition to glycan structure, FcγRIIIa affinity and ADCC activity are sensitive to the subclass of the IgG heavy chain. Human IgG1 and IgG3 isoforms show good binding and activity whereas IgG2 and IgG4 isotypes have reduced binding and activity.

For some therapeutic applications, decreased or abolished ADCC activity is required. In cases where the antibody binds a cell surface receptor to inhibit a signaling pathway, it may be undesirable to cause depletion of cells that express the receptor. Decreased ADCC activity may be achieved by producing an IgG1 antibody in aglycosylated form or by changing the isotype to IgG2 or IgG4. Aglycosylation is accomplished through expression of the antibody in E. coli or through mutation of Asn297 to remove the glycosylation site for production in mammalian cells. However, aglycosylation can affect the conformation and stability of the Fc. Glycosylated IgG4 is an...
alternative with a preferred version having the hinge mutation S228P (Eu numbering) to stabilize the antibody against Fab arm exchange. Although in general IgG4 isotype antibodies have reduced capacity to support in vitro ADCC, in vivo target cell depletion activity in humans has been noted with an IgG4 version of an anti-CD52 antibody. As with IgG1 isotype antibodies produced in CHO cells, there could be variability in the fucose levels on IgG4 antibodies. Afucosylation has been demonstrated previously to increase the in vitro ADCC activity of an anti-CD20, IgG4 isotype antibody. Here, we show that afucosylation does increase the in vivo B cell depletion activity of an IgG4 isotype anti-CD20 antibody, suggesting that, for some IgG4 antibodies produced in CHO, it may be necessary to monitor fucosylation status in order to limit cell killing activity.

**Results**

**Production and receptor-binding measurements on anti-human CD20 IgG4 isotype antibody**

An anti-human CD20 antibody (x-huCD20) was chosen for tests of IgG4 isotype switch on activity because an assay of ADCC activity on huCD20 expressing cell lines is readily available, the level of activity for the IgG1 subclass antibody is sensitive to fucosylation status, and in vivo activity can be measured in pre-clinical models. An x-huCD20-IgG4 heavy chain construct was expressed in both Chinese hamster ovary (CHO) and Fut8-knock-out (KO) CHO cells. Antibody was easily purified from both expression hosts using standard methods and shown to bind CD20 in a WIL2 cell-based assay (Fig. S1). CD20-binding of the 2 preparations of x-huCD20. IgG4 was similar, but appeared to be slightly weaker than the binding of x-huCD20.IgG1 (CHO). The relative binding affinities compared to IgG1 were 0.5 and 0.75 for IgG4 produced in CHO and Fut8-KO CHO cells, respectively. As expected, 100% afucosylated glycans was observed for the antibody obtained from expression in the Fut8KO host. The percentage of afucosylated glycans on IgG4 antibody produced transiently in CHO was higher than for IgG1 or mlgG2a subclass antibodies purified from stably transfected CHO cells (Table 1). The reproducibility and generality of this effect has not been tested.

Binding of x-huCD20.IgG4 to a panel of human Fcγ receptors was reduced relative to the IgG1 isotype (Table 2). As shown in Fig. 1, binding of x-huCD20.IgG4 to human FcγRIIIa-V158 was very weak and did not approach saturation at the highest concentrations tested; however, increased binding was observed for the material isolated from the Fut8-KO host. Complete afucosylation increased binding to the F158 allotype was 3.8-fold and the increase for Val158 was > 15-fold (Table 2). As observed for IgG1 antibodies, the increase in binding upon afucosylation was specific to FcγRIIIa.

**ADCC activity of afucosylated x-huCD20.IgG4**

ADCC activity of the various x-huCD20 antibodies on WIL2S tumor cells was assessed using a NK cell line as effector (Fig. 2). x-huCD20.IgG4 (CHO) showed measurable ADCC activity, albeit much reduced relative to x-huCD20.IgG1 (CHO). A comparison with x-huCD20.IgG4 (Fut8KO) indicates that complete afucosylation increases the activity by nearly 50-fold, calculated from the ratio of EC50 values (Table 3), to within about 5-fold of the activity measured for x-huCD20.IgG1 (CHO).

We further probed the effect of IgG4 afucosylation on ADCC activity by testing preparations of varied fucose levels. It is difficult to produce preparations of defined but varied fucose levels unless procedures are used to modulate the activity of the fucosyl transferase. This approach generates a distribution of molecules with one or both heavy chains of the assembled antibody being afucosylated. Instead, we produced these materials by blending of CHO-produced and Fut8KO-CHO-produced antibodies in various ratios. In these blends, and since the heavy chains do not exchange, assembled molecules have essentially neither or both chains afucosylated. A similar approach has been used to establish a quantitative relationship between fucose levels and ADCC activity for the IgG1 version of x-huCD20. As shown in Fig. 3A, the ADCC activity of these samples increases as the fraction of Fut8KO-CHO-derived antibody is increased. ADCC activity is barely detectable if the afucosylation level is <20%. The ADCC activity as a percentage of the activity measured for the 100% Fut8KO-CHO antibody shows a linear dependence on % afucosylation within the range tested (Fig. 3B). The correlation coefficient of this linear dependence is quite good, suggesting that measurements of the % afucosylation of different lots of x-huCD20.IgG4 could be used to predict the expected ADCC activity. A linear relationship between afucosylation levels and ADCC activity has been previously observed in an IgG1 anti-CD20 antibody.

**Depletion of B cells in mouse model using anti-mouse CD20.IgG4 antibody**

Given that afucosylation increased the in vitro ADCC activity of IgG4 antibody, we tested whether this translated to in vivo activity in an animal model relevant to human disease. A model of B cell depletion in mice has been used previously to compare therapeutics intended for treatment of B cell malignancies. In mice, a change in B cell population in the blood and tissues upon antibody treatment is easily monitored using fluorescence-activated cell sorting (FACS) analysis. For our purposes, we used mice having the murine FcγRIIa knocked out and transgenic for human FcγRIIa-Val158. However, since these mice are not transgenic for human CD20, and because x-huCD20 doesn’t bind murine CD20, it was necessary to

| Table 1. Fucose levels on mAbs. |
|------------------------|
| mAb                    | % Afucosylated Glycan |
| x-huCD20.IgG1 (CHO)    | 6                      |
| x-huCD20.IgG4 (CHO)    | 11 ± 0.9               |
| x-huCD20.IgG4 (Fut8KO) | 100 ± 0.2              |
| 5D2.mlglG2a (CHO)      | 4 ± 0.8                |
| 5D2.hlglG4 (CHO)       | 29 ± 2.2               |
| 5D2.hlglG4 (Fut8KO)    | 100 ± 0.4              |

IgG4 antibodies were produced by transient transfection of CHO cells; IgG1 and mlglG2a antibodies were expressed by stable transfection of CHO cells. Values for % afucosylated glycans are shown ± standard error.
change to an antibody specific for murine CD20. Mouse monoclonal antibody 5D2.16.8 was chosen, reformatted as a mouse/human IgG4 chimera, and expressed in both CHO and Fut8KO cells. As shown by FACS analysis (Fig. S2), the IgG4 antibody is functional for binding mCD20. The IgG4 version of 5D2 showed very weak binding across a panel of mouse FcR; however, afucosylation did increase apparent binding affinity to mFcRIV (Supplemental Table 1). Since the fucosylated antibody produced in CHO had very weak binding to mFcRIV, it was not possible to calculate the fold improvement in binding accompanying afucosylation.

Administration of 5D2.mIgG2a at either 12.5 \( \mu g \) or 125 \( \mu g \) led to a rapid depletion of peripheral blood B cells (Fig. 4A) relative to the isotype control antibody, anti-ragweed. 5D2.hIgG4 (CHO – 29% afucosylated) at either dose did not show significant activity for B cell depletion. In contrast, 5D2.hIgG4 (Fut8KO-CHO – 100% afucosylated) showed activity comparable to 5D2.mIgG2a, even at the lower dose of 12.5 \( \mu g \).

Depletion of B cells in compartments other than peripheral blood, for example in the marginal zones (Fig. 4D), is known to be more challenging. Nonetheless, 5D2.mIgG2a even at the lower dose of 12.5 \( \mu g \) showed good B cell depletion activity against spleen (Fig. 4B), follicular (Fig. 4C), marginal zone

### Table 2. Relative human FcγR binding measured by ELISA.

| FcγRI | FcγRlla (H131) | FcγRlla (R131) | FcγRllb | FcγRlla(F158) | FcγRlla(V158) |
|-------|----------------|---------------|---------|--------------|--------------|
| x-huCD20.IgG1 |               |               |         |              |              |
| mid-OD       | 0.87           | 1.76          | 1.79    | 1.68         | 1.59         | 1.60         |
| ng/ml at mid-OD | 120            | 980           | 1700    | 3000         | 2300         | 670          |
| Relative binding | 1.0           | 1.0           | 1.0     | 1.0          | 1.0          | 1.0          |
| x-huCD20.IgG4 (CHO) |       |               |         |              |              |
| Top OD       | 0.06           | 2.86          | 3.29    | 2.20         | 0.12         | 0.18         |
| Relative binding | <0.02\(^a\)  | 0.071         | 0.20    | <0.02\(^a\) | <0.003\(^a\) |
| x-huCD20.IgG4 (Fut8KO) |     |               |         |              |              |
| Top OD       | 0.15           | 2.75          | 3.29    | 2.69         | 1.46         | 1.99         |
| Relative binding | <0.04\(^a\) | 0.069         | 0.28    | 0.21         | 0.076        | 0.045        |

\(^a\)The top OD of the sample titration curve was much lower than the mid-OD for the x-huCD20-IgG1 control. To estimate the relative binding, the x-huCD20-IgG1 concentration corresponding to the top OD of the sample was calculated and divided by 500 ng/ml (for FcγRII binding) or 25000 ng/ml (for FcγRII binding).

Figure 1. Binding to human FcγRlla-Val158 determined by ELISA. Antibody concentration curves shown for x-huCD20.IgG1-CHO (diamonds), x-huCD20.IgG4-CHO (circles), and x-huCD20.IgG4-Fut8KO (squares). Readings from duplicate wells are shown and the lines are the result of a 4 parameter fit.

Figure 2. ADCC activity of x-huCD20 antibody on WIL2-S target cells using NK cell line as effector cells. Antibody concentration curves shown for x-huCD20.IgG1-CHO (diamonds), x-huCD20.IgG4-CHO (circles), and x-huCD20.IgG4-Fut8KO (squares). The results shown are the values of duplicate wells from one representative plate of an ADCC assay with the lines representing the result of a 4 parameter fit.
ADCC activity (EC50) of x-huCD20.IgG4 blends of CHO and Fut8KO-CHO produced antibody.

| Target %Afucosylation |Measured %Afucosylation | ADCC EC50 (ng/mL) |
|-----------------------|-------------------------|-------------------|
| 11                    | 16                      | 937 ± 82          |
| 20                    | 24                      | 185 ± 43          |
| 30                    | 32                      | 96 ± 28           |
| 40                    | 42                      | 58.8 ± 9.0        |
| 50                    | 51                      | 42.3 ± 11.7       |
| 75                    | 75                      | 24.5 ± 4.9        |
| 100                   | 98                      | 19.7 ± 2.9        |

Actual afucosylation levels determined using HILIC. ADCC EC50 values shown as mean with standard error.

Discussion

ADCC is elicited when NK cells expressing FcγRIIIa encounter target cells coated with antibody. FcγRIIIa binds the homodimeric Fc in an asymmetric fashion having a stoichiometry of 1 receptor per Fc fragment. The interaction between soluble FcγRIIIa and Fc is low affinity such that activation of effector function requires the multivalent, antibody-mediated interaction between target and effector cells. Affinity of the soluble receptor for Fc is increased if the core fucose of the Fc glycan is removed. Structures of glycosylated FcγRIIIa bound to fucosylated and afucosylated antibody suggest this reflects a direct interaction between the Fc glycan and a carbohydrate chain attached to Asn-162 of the receptor. Removal of the core fucose enables a closer interaction of the 2 carbohydrate chains with an associated increase in affinity. Since other Fc receptors lack the Asn-162 glycosylation site, the effect of fucose removal on Fc receptor affinity is restricted to human FcγRIIIa and human FcγRIII. This increased affinity ultimately translates to increased ADCC activity for afucosylated antibodies.

Human IgG1 and IgG3 antibodies bind FcγRIIIa with modest affinity and are generally capable of eliciting ADCC activity whereas IgG2 and IgG4 isotypes show reduced binding to FcγRIIIa and weak capacity to support ADCC. Interaction of FcγRIIIa with Fc involves mostly the CH2 domain of the Fc. Human IgG4 isotype differs from human IgG1/3 isotype at 4 positions in the CH2 domain – H268Q, Y296F, A327G, and A330S – that contact human FcγRIIIa in the structure determined for the IgG1- FcγRIIIa complex. These changes contribute to the decreased FcγRIIIa binding and ADCC activity of IgG4 isotype antibodies. However, this deficit in binding and activity can be partially recovered upon afucosylation of IgG4. For the x-huCD20 antibody we tested here, in vitro ADCC activity of the IgG4 isotype antibody becomes appreciable if the afucosylation level is greater than about 20%. Since cytotoxic activity can be highly dependent on epitope and binding site density on the target cell, the level of afucosylation required to produce significant activity may be different for other anti-CD20.IgG4 antibodies, and also for other IgG4 antibodies directed against different membrane targets. In addition, there can be differences between methods and labs for afucosylated glycan quantitation, as we observed for HILIC (16%) and mAb-Glyco chip (11%) analysis of CHO produced x-huCD20. IgG4. We presume the ADCC enhancement is mediated by a carbohydrate-carbohydrate interaction as observed for the binding of human IgG1 isotype to FcγRIIIa. Likely only one of the heavy chains needs to be afucosylated to produce the full effect for increased ADCC activity. Since the activity increase for the blends reflects mostly the activity of antibodies with both chains afucosylated, and because variation in fucose content resulting from cell culture conditions will produce a distribution of molecules with 0, 1, and 2 chains afucosylated, total afucosylation levels required to produce significant activity increase may be higher than 20%.

Ultimately, this increase in in vitro binding and activity is translated to in vivo activity, resulting in greater depletion of targets cells when the IgG4 isotype antibody is fully afucosylated. Although we have not tested in vivo a full range of afucosylation levels, we can conclude that B cell depletion activity of the anti-CD20 IgG4 antibody is absent when the afucosylation level is below 30%. Similarly, Scallon et al. found that a chimeric hamster/human IgG4 antibody with 10% afucosylation was essentially inactive in a mouse model of T cell depletion relative to the 10% afucosylated IgG1 antibody. A limitation of effector function studies in the mouse is that the binding abilities and expression patterns of the panel of Fcγ receptors differ from human. Mice have a unique activating receptor, FcγRIV, not present in human that is expressed on mouse monocytes, macrophages and neutrophils. We used mice having mFcγRIII knocked out and expressing the human FcγRIIIa transgene on natural killer (NK) cells. These mice may have abnormal expression of the transgene on other cell types. It is unclear from the results in the mouse model whether the increased cell killing observed for afucosylation of 5D2.IgG4 involves endogenous mFcγRIV or the transgenic hFcγRIIIa receptor. In general, human IgG4 binds much more weakly to mouse Fcγ receptors than endogenous mlgG2a, but we did observe increased binding to mFcγRIV for fully afucosylated 5D2.IgG4 (Table S1).

Although human IgG4 isotype antibodies have reduced ADCC activity compared with human IgG1 isotype antibodies, effector function both in vitro and in vivo is increased through afucosylation. If reduced effector function is required in an IgG4 therapeutic antibody that targets membrane bound receptors, then fucosylation status of the
antibody may need to be monitored. As shown here for anti-CD20, if the level of IgG4 afucosylation is less than about 30%, then in vivo cell killing should be minimal and further work to increase fucose levels may not be warranted. Also, amino acid substitutions in the IgG4 Fc domain may be employed to further reduce engagement of Fcγ receptors.

Alternatively, use of an afucosylated IgG4 isotype antibody may enable a strategy to increase the therapeutic index of an antibody if the IgG1 version has unacceptable toxicity related to effector function. An intermediate intensity of cell killing (Figs. 2–3) may be obtained with the IgG4 subclass and varied afucosylation levels.

Figure 3. Dependence of ADCC activity of x-huCD20.hIgG4 on afucosylation levels. (A) ADCC curves observed for samples prepared by blending CHO-produced and Fut8KO-CHO-produced antibody. The results shown are mean values of duplicate wells from one representative plate of an ADCC assay. (B) Relationship of ADCC activity calculated from ADCC curves to the % afucosylation. Data presented are normalized mean ADCC activity values, assigning the 98% afucosylated sample as ADCC activity of 100%. The line is the result of a linear least squares analysis of the relative % ADCC data with correlation coefficient $R = 0.99$; error bars represent corresponding standard error of the mean and in most cases are smaller than the data point symbol.
Material and methods

Reformatting of hu-xCD20 into human IgG4 isotype

A pRK plasmid containing DNA for a humanized IgG4 isotype antibody, having the S228P hinge stabilizing mutation, was used as the source of IgG4 constant domains. A fragment obtained upon ApaI/HindIII digestion of this DNA encodes a CH1-CH3 fragment and was used to replace a similar fragment in a pRK-based plasmid designed for expression of x-huCD20. IgG1 antibody. A clone with correct sequence for the x-huCD20.IgG4 (S228P) construct was identified by DNA sequencing. The x-huCD20.IgG4 plasmid, and also the pRK plasmid containing the light chain of x-huCD20, were produced at larger scale and purified (Qiagen maxi-prep) for transient expression of the IgG4 isotype antibody.

Anti-mCD20 antibody reformatting into human IgG4 isotype

Heavy and light chain DNA of mouse anti-mouse CD20 monoclonal antibody 5D2.16.8 (previously cloned from a hybridoma cell line into mouse IgG2b and mouse kappa) was diluted to less than 500 ng/μl and amplified through PCR with a premix Taq polymerase kit (Takara) using forward primers corresponding to antibody framework region and human vector signal sequence and reverse primers corresponding to antibody framework region and human IgG4 or kappa CH1 (primers shown below). The resulting PCR products were purified using a PCR product cleanup kit (Qiagen). The heavy chain PCR product and pRK human IgG4 S228P vector were digested using ApaI and EcoRI restriction enzymes (NEB) according to manufacturer recommendations. The light chain PCR product and pRK human kappa LPG3 vector were digested using KpnI and EcoRI restriction enzymes (NEB) according to manufacturer recommendations. Digestion products were purified by gel extraction (Qiagen) and vectors were treated with shrimp alkaline phosphatase (Promega) to prevent self-ligation. Inserts and vectors were ligated using T4 DNA Ligase (NEB) at a 1:1 ratio. Ligated vectors were used to transform XL-1 Blue competent E. coli cells (Stratagene); bacteria were spread onto agarose plates with 50 μg/ml carbenicillin and allowed to grow overnight at 37°C. 48 colonies were picked from each of the heavy and light chain transformation plates into 2YT media containing 50 μg/ml carbenicillin and incubated 15 hours at 37°C. Plasmids were purified from bacterial pellets by miniprep (Qiagen) and sequenced using forward and reverse primers corresponding to 3' and 5' regions of the pRK vectors near the multiple cloning site. Subclones with the fewest nucleotide errors from original

Figure 4. Depletion of B cells upon single intravenous injection of 5D2.mIgG2a, 5D2.hIgG4 (CHO), or 5D2.hIgG4 (Fut8-KO-CHO) relative to isotype control (anti-ragweed. mIgG2a) in human FcγIIIa transgenic mice. Two doses were tested (12.5 μg and 125 μg) with the B cells determined by FACS analysis. B cell levels are shown for (A) blood, (B) total spleen, (C) splenic follicular B cells, and (D) splenic marginal zone B cells. B cell levels shown are for 5 d post antibody injection except for blood B cells where levels at 2 d are shown.
Antibody samples were injected onto a HPLC-mAb-Glyco chip (Agilent, #G4240-64025), which was used within the Agilent 1260 Chip Cube coupled to a quadrupole time-of-flight mass spectrometer (Agilent 6520 Q-TOF). Upon injection, antibodies were incubated on the mAb-Glyco chip (Agilent, #G4240-64025), which was used within the Agilent MassHunter Qualitative Analysis software. The software's algorithm utilizes a combination of accurate mass with a mass tolerance of 10 ppm and expected retention time for glycan identification.

Measurement of Fcγ-R-binding and CD20-binding by ELISA

Binding of anti-human CD20 human IgG antibodies to FcγRs was measured by ELISA. MaxiSorp 384-well plates (Thermo Scientific, Nunc, Roskilde, Denmark) were coated with soluble human FcγRs at 1 μg/mL. Serially diluted antibodies (0.0085–500 ng/mL IgG for FcγRI binding and 0.42–25000 ng/mL IgG complexed with goat F(ab’)2 anti-human κ for FcγRII and FcγRIII binding) were added in duplicates to the ELISA plates as described by Lu et al.32 Bound IgG was detected using biotinylated Fab anti-idiotype antibody (Genentech). This antibody detected x-huCD20.IgG1 and x-huCD20.IgG4 equally (data not shown). The ELISA signal was generated by adding streptavidin-horseradish peroxidase (HRP) (GE Healthcare, Piscataway, NJ) followed by 3,3’,5,5’-tetramethyl benzidine (TMB) (Kirkegaard & Perry Laboratories, Gaithersburg, MD) as substrate. The reaction was stopped by adding 1 M phosphoric acid. Absorbance was read at 450 nm on a Multiskan Ascent reader (Thermo Scientific, Hudson, NH). Binding of anti-mouse CD20 chimeric mouse-human IgG mouse FcγRs by ELISA was carried out similarly using ELISA plates coated with soluble mouse FcγRs (Genentech) and serially diluted antibodies (0.42–25000 ng/mL complexed IgG). Bound IgG was detected with goat F(ab’)2 anti-human F(ab’)2-HRP (Jackson Immunoresearch, West Grove, PA). This detection antibody detected IgG4 at least 50% as efficient as IgG1. For data analysis, the absorbance at the midpoint of the x-huCD20-IgG1 titration curve (mid-OD) was calculated as the average OD reading of the highest and lowest standards. The corresponding concentrations of x-huCD20-IgG1 and samples were determined using a 4-parameter nonlinear regression curve-fitting program (XLfit, Guildford, Surrey, UK or KaleidaGraph, Synergy Software, Reading, PA). Relative affinities were calculated by dividing the midpoint concentration of x-huCD20-IgG1 by the midpoint concentration of sample.

Binding to human CD20 expressed on WIL2S cells was performed as described in Hong et al.33 using a competitive format. Serially diluted antibody (11–25000 ng/mL) and biotinylated IgG1 (0.67 μg/mL) were added to WIL2 cells. Streptavidin-HRP was added followed by TMB to generate the ELISA signal. Relative binding was calculated using the mid-OD concentrations as described above.

ADCC measurements

% ADCC activity was determined using an engineered NK cell line derived from NK92 (Glycart Biotechnologies/Roche) as effector cells and DELFIA BATDA (Perkin Elmer) labeled WIL2-S cells as target cells, essentially as described previously.24 The NK cell line stably expresses human FcγRIIA-F158. A mixture of labeled WIL2-S target cells (2 × 10⁶) and NK cells (1.0 × 10⁵; 5:1 effector:target) prepared in assay medium (RPMI 1640 with 10% heat inactivated-FBS, 2 mM L-glutamine and 20 mM HEPES, pH 7.2) was added to each well of a round-bottom, 96-well tissue culture plates. Targeted dilutions of antibodies (10,000 to 0.5 ng/mL) were added to the
plates, and the plates were incubated for 3 hours. Cell lysis was measured using time-resolved fluorescence in relative fluorescence units (RFU) by excitation at 345 nm and quantifying emission at 615 nm using a microplate reader SpectraMax® 190 (Molecular Devices). Absorbance of wells containing only labeled target cells served as the control for background (Background), whereas wells containing labeled target cells with DELFIA BATDA lysis buffer provided the maximum available signal (Maximum Release). Spontaneous release (SR) was measured from wells containing only target cells, whereas SR + NK was measured from wells containing both target and effector cells without antibody.

The % ADCC was determined according to the formula below

\[
% \text{ADCC} = \frac{(\text{Specific Release} - \text{Background}) - (\text{Spontaneous Release} - \text{Background})}{(\text{Maximum Release} - \text{Background}) - (\text{Spontaneous Release} - \text{Background})} \times 100
\]

The dose response curves were generated by plotting the mean % ADCC from duplicates against the concentration of antibody sample in ng/mL using a 4-parameter fit. Data analysis was performed using parallel line analysis curve fitting software (PLA) from SOFTmaxPRO™, generating separate lines for the reference material and test material to estimate potency of the test material relative to the reference material.

**x-huCD20.IgG4 of varied fucose through blending**

Samples of varied fucose level were prepared by mixing CHO-produced and Fut8KO-CHO-produced x-huCD20. IgG4 in various ratios. Five samples were prepared targeting 20, 30, 40, 50, and 75% afucosylation. Actual afucosylation levels determined by hydrophilic interaction liquid chromatography (HILIC) were 24, 32, 42, 51, and 75%. This can be compared to the afucosylation levels determined by HILIC for the non-blended samples of Fut8KO-CHO-produced antibody, 98%, and CHO-produced antibody, 16%. ADCC activity of these samples was determined as described above. Dependence of percent activity on afucosylation levels was determined using the 98% sample as standard and assigning the activity of that sample as 100%. Percent activity relative to the 100% sample was determined from the EC50 value obtained from analysis of the antibody concentration dependence of lysis.

**B cell depletion in hFcγRIIIα-transgenic mice**

In these experiments, mice were a C57BL6 strain expressing transgenic human CD16 and deficient in murine FcγRIII. Mice expressing human CD16 and deficient in murine FcγRIII were obtained from Dr. Jeffrey Ravetch (The Rockefeller University). They were fully backcrossed to C57BL6. Mice were intravenously administered isotype control (Ragweed, mIgG2a), anti-mCD20 mAb (5D2.mIgG2a), anti-mCD20 mAb (5D2.hIgG4) or afucosylated anti-mCD20 mAb (5D2.hIgG4). There were 5 mice in each treatment group. B cells from tail bleeds were assessed 2 d after dosing, and FACS analysis was performed another 5 d later on B cells from blood, spleen and mesenteric lymph nodes. B cells from blood and lymph nodes were defined as CD21+CD23+, while splenic follicular and marginal zone B cells were gated as CD21+CD23hi and CD21hiCD23int, respectively.

**Disclosure of potential closure of interest**

All authors are current or former employees of Genentech, Inc., a member of the Roche group, and may own stock or stock options of Roche.

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