Humanised IgG1 antibody variants targeting membrane-bound carcinoembryonic antigen by antibody-dependent cellular cytotoxicity and phagocytosis

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BACKGROUND: The effect of glycoengineering a membrane specific anti-carcinoembryonic antigen (CEA) (this paper uses the original term CEA for the formally designated CEACAM5) antibody (PR1A3) on its ability to enhance killing of colorectal cancer (CRC) cell lines by human immune effector cells was assessed. In vivo efficacy of the antibody was also tested.

METHODS: The antibody was modified using EBNA cells cotransfected with β-1,4-N-acetylglucosaminyltransferase III and the humanised hPR1A3 antibody genes.

RESULTS: The resulting alteration of the Fc segment glycosylation pattern enhances the antibody's binding affinity to the FcγRIIIa receptor on human immune effector cells but does not alter the antibody's binding capacity. Antibody-dependent cellular cytotoxicity (ADCC) is inhibited in the presence of anti-FcγRIII blocking antibodies. This glycovariant of hPR1A3 enhances ADCC 10-fold relative to the parent unmodified antibody using either unfractionated peripheral blood mononuclear or natural killer (NK) cells and CEA-positive CRC cells as targets. NK cells are far more potent in eliciting ADCC than either freshly isolated monocytes or granulocytes. Flow cytometry and automated fluorescent microscopy have been used to show that both versions of hPR1A3 can induce antibody-dependent cellular phagocytosis (ADCP) by monocyte-derived macrophages. However, the glycovariant antibody did not mediate enhanced ADCP. This may be explained by the relatively low expression of FcγRIIIa on cultured macrophages. In vivo studies show the efficacy of glycoengineered humanised IgG1 PR1A3 in significantly improving survival in a CRC metastatic murine model.

CONCLUSION: The greatly enhanced in vitro ADCC activity of the glycoengineered version of hPR1A3 is likely to be clinically beneficial.

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Antibodies are increasingly used for immunotherapy of cancers, with over 200 currently in trials and 12 licensed for use in treatment (Adams and Weiner, 2005; Carter, 2006; Reichert and Valge-Archer, 2007). There are now three antibodies approved by the US FDA for treatment of advanced colorectal cancer (CRC) (Goldenberg, 2005; Saltz et al, 2006; Reichert and Valge-Archer, 2007). The main potential advantages of this form of therapy lie in their specificity, lesser side-effects and their ability to elicit a tumour response by multiple mechanisms (Carter, 2006; Reichert and Valge-Archer, 2007). The outcomes of early clinical trials using murine unconjugated and radioconjugated monoclonal antibodies for cancer treatment were disappointing (Goldenberg, 2002; Reichert and Valge-Archer, 2007) probably because of immune responses to foreign proteins reducing serum half-lives, and the delivery of inadequate radiation doses from the radioconjugates to the tumour (Goldenberg, 2002; Sharkey and Goldenberg, 2005; Reichert and Valge-Archer, 2007). Responses were also severely limited by the attenuated ability of the Fc segment of murine antibodies to interact with the relevant receptors of the human immune system. These problems have now been largely overcome by the development of technologies for the production of chimeric, humanised and completely human antibodies (Clynes, 2006).

Two lines of evidence support the importance of immune function for naked antibody therapy. Firstly, Clynes et al (2000) showed that antibody-based killing of xenografted and syngeneic tumours was abrogated in FcγR knockout mice. This established that the ability of antibodies to reduce tumour burden was greatly reduced in the absence of immune cells expressing FcγRs. Secondly, colorectal, lymphoma and breast cancer patients carrying the polymorphic variants in the FcγRIIa and FcγRIIa genes that encode for high affinity binding receptors for the Fc antibody segments, have the best clinical outcomes following
antibody treatment (Weng and Levy, 2003; Zhang et al, 2007; Musolino et al, 2008).

Responses to antibody therapy for solid cancers are so far modest for cetuximab and trastuzumab monotherapy. Although cetuximab in combination with irinotecan improves the response rate from 10% to over 20% (Cunningham et al, 2004), monotherapy response rates in haematological malignancies are markedly higher than for solid cancers (McLaughlin et al, 1998). This may be due to the increased surface area of haematological tumours that is exposed to antibody binding and to NK cells (Olszewski and Grossbard, 2004). Emphasis is now being placed on techniques that ‘tune’ antibodies to make them more efficient at harnessing the immune system for killing solid cancers. Thus, technologies that ‘tune’ antibodies to make them more efficient at ADCC (Richman and Bodmer, 1987) were utilized to enhance the immune Fc segment (Lazar et al, 2004). One approach uses protein engineering to modify the Fc segment (Lazar et al, 2006). Another uses glycoengineering, which has been shown to improve in vitro ADCC by 10–100-fold (Umana et al, 1999; Niwa et al, 2004b; Schuster et al, 2005). In this study, we have used glycoengineering to enhance the immune effector function of an antibody that we have shown earlier to kill CRC-derived cell lines by ADCC (Conaghan et al, 2008).

The antibodies used for the treatment of CRC target EGFR and VEGF (Cunningham et al, 2004; Adams and Weiner, 2005; Goldberg, 2005; Carter, 2006; Saltz et al, 2006) and only appear to be effective in subsets of patients. Earlier work from our laboratory has shown that the antibody PR1A3, which is specific for membrane-bound carcinoembryonic antigen (CEA, formally designated CEACAM5), elicits ADCC against CEA-positive CRC cell lines even in the presence of super-physiological levels of free CEA (Durbin and Bodmer, 1987; Durbin et al, 1994; Conaghan et al, 2008). CEA is present on a high proportion of colorectal cancers, in which it is accessible to intravenously administered antibody, whereas in normal colorectal epithelium, due to the apical localisation of the antigen, it is not (Granowska, 1993). Radioconjugates targeting CEA have been used in early phase clinical trials in CRC (Wong et al, 2004; Liersch et al, 2005), but there is so far no un conjugated anti-CEA antibody licensed for treatment (Blumenthal et al, 2008). The pre-clinical studies reported here are directed at optimising PR1A3 for use in the treatment of colorectal cancer. This antibody has been shown to bind specifically to CEA in the R3-GPI domains. This epitope is at the point CEA anchors into the membrane, and this presumably explains why PR1A3 binds to membrane-bound CEA but not soluble CEA (Durbin et al, 1994; Conaghan et al, 2008).

The three classes of FcRs are expressed on NK cells, known to be potent activators of ADCC (Schmitz et al, 2002; Lazar et al, 2006; Conaghan et al, 2008), and on monocytes and granulocytes, which have also been identified as having at least some potential for antibody-dependent killing of target cells, for example, by antibody-dependent cellular phagocytosis (ADCP) (Munn et al, 1991; Huls et al, 1999; Dhodapkar et al, 2005; Lazar et al, 2006; Karagiannis et al, 2007; McEarchern et al, 2007). We describe here a modified phagocytosis assay, based on that developed by Munn et al (1991), and its application to the in vitro testing of the ability of humanised and glycoengineered PR1A3 to kill CRC-derived cell lines. We also present in vivo data that shows the efficacy of PR1A3 as a therapeutic agent in a mouse xenograft model.

MATERIALS AND METHODS

Cell lines

The SKCO-1 (from ATCC), LS174T (from BH Tom, NW University, Chicago), LoVo (from ATCC), HCT116 (from ATCC) and MKN45 (from Cell Services LIF, CRUK, London, UK) cell lines were cultured as described (Conaghan et al, 2008). For cytotoxicity and phagocytosis assays, cells were suspended in 2% RPMI 1640 medium with 1% glutamine and 10% FCS (RPMI complete medium).

Antibodies

PR1A3 The original murine IgG1k monoclonal antibody to CEA (Richman and Bodmer, 1987) was humanized by Stewart et al (1999). Murine (mPR1A3, IgG1) and unmodified humanized (uhPR1A3, IgG1) antibodies were acquired from the Biotherapeutics Development Unit, Clare Hall, CRUK, London, UK. Human IgG2a PR1A3 (hPR1A3, IgG2a) and a glycoengineered version of re-derived humanised IgG1 PR1A3 (ghPR1A3, IgG1) were made by joining the variable region of PR1A3 to murine IgG2a and the complementarity determining regions of PR1A3 to human IgG1 variable regions, respectively (Jones et al, 1986). Glycoengineering was as described (Umana et al, 1999; Schuster et al, 2005). Briefly, vectors containing IgG heavy and light chains, and (β(1,4)-N-acetylglucosaminyltransferase-III (GnT-III), were transfected into EBNA cells. The enzyme modifies the N-glycosylation pattern at Asn-297 of the IgG heavy chain resulting in a high degree of bisected, afucosylated oligosaccharides (Umana et al, 1999; Schuster et al, 2005; Ferrara et al, 2006), which has been shown earlier to increase the affinity to FcγRIIa (Ferrara et al, 2006). The modified antibody was purified from culture supernatants using Protein-A, cation-exchange chromatography and subsequent size-exclusion chromatography.

SM3E This is a humanised IgG1 antibody that binds with extremely high affinity (Kd = 20 pM) to both soluble and membrane-bound CEA (Graff et al, 2004). A glycoengineered variant was generated using the methods described above.

Blocking antibodies Purified IgG1 anti-CD16 (FcγIII) (clone 3G8, which is unable to distinguish between FcγRIIIa and FcγRIIIb), purified IgG1 anti-CD64 (clone 10.1), and Fab, fragments targeting CD16 (clone 3G8), CD32 (FcγII, clone 7.3) and CD64 (FcγII, clone 10.1) were purchased from Ancell Corp, Bayport, MN, USA. Purified IgG2b anti-CD32 (clone IV.3) was obtained from Stemcell Technologies (Vancouver, Canada).

FITC-conjugated murine anti-human IgG (clone G18-145), CD3-FITC, CD11b-FITC, CD14-FITC, CD15-FITC, CD16-PE, CD19-PE, CD32-PE, CD45-PE, CD56-APC, CD64-APC, IgG1-FITC (isotype control), IgG1-PE (isotype control) and IgG1-APC (isotype control) were obtained from BD Biosciences, Oxford, UK. Anti-HLA class II (DA-2) was obtained from Monoclonal Antibody Services, CRUK (Brodsky et al, 1979).

Isolation of immune effector cells

Peripheral blood mononuclear cells (PBMCs) were isolated from healthy laboratory volunteers, having taken informed consent, or fromuffy coats obtained from single donors (National Blood Service, Bristol, UK). The Rosettesep Human NK isolation cocktail (Stemcell Technologies) was used to obtain an enriched population of human NK cells from blood, as described (Rodà et al, 2006). Granulocytes were separated using One-step Polymorph (Accurate Chemical and Scientific Corp, Westbury, NY, USA). Monocytes were enriched from blood using the Rosettesep Human Monocyte Enrichment Cocktail (Stemcell Technologies). Purification was verified by phenotypic analysis of surface markers (See Supplementary Materials for details).

Flow cytometric analysis to compare uhPR1A3 and ghPR1A3 binding to membrane-bound CEA

Cells from SKCO-1, a high CEA expressing CRC line used as target, were incubated in varying concentrations of uhPR1A3 or ghPR1A3
(0.001 – 100 μg ml⁻¹), and with anti-human IgG-FITC as secondary antibody. The cells were then washed once with FACS buffer (PBSA, 1% FCS, 1% Sodium Azide) and the supernatant removed before resuspending in 2% paraformaldehyde in PBSA. FACS analysis was carried out using a FACS Calibur Flow cytometer.

Fluorescence-based EuTDA cytotoxicity assay

As described (Conaghan et al, 2008), a fluorescent probe, BATDA (Blomberg et al, 1996) (Perkin Elmer, Boston, MA, USA), was used to label target cells. PBMCs, NK cells, monocytes and granulocytes were used as effectors with varying effector: target cell ratios and concentrations of antibodies. After appropriate incubation, supernatant was added to Europium in 96-well plates and the resulting fluorescence read in a time-resolved fluorometer.

The EuTDA assay was used to compare the ADCC activity of mPRA13 IgG1, mPR1A3 IgG2a and uhPR1A3 IgG1 on SKCO-1 cells, using antibody at a final concentration of 20 μg ml⁻¹ and either PBMCs or NK cells as effectors.

The effect of blocking FcγRs I, II and III using Fab2 or IgG antibodies was assessed using the EuTDA to measure uhPR1A3 mediated ADCC of SKCO-1. PBMC and enriched NK cells were used as effectors. Murine IgG prostate membrane specific antigen (PMSA, provided by Robert Vessella, University of Washington) was used as an isotype murine control.

The relative ability of PBMCs, NK cells, monocytes and granulocytes, enriched, as described earlier, to elicit ADCC was assessed using the EuTDA assay with uhPR1A3.

Monocyte-derived macrophages for phagocytosis assays

Enriched monocytes were cultured on Lumox Petraperm Plates (Greiner, Bio-One, North America, Monroe, NC, USA), for 8–12 days at 37°C in a 5% CO₂ incubator, in X-VIVO 15 medium (BioWhittaker, Walkersville, MD, USA), supplemented with M-CSF, GM-CSF and γIFN (Peprotech, UK) (see Supplementary Materials for details). Target cells (SKCO-1) were labelled with CellTracker Green CMFDA (Invitrogen, San Diego, CA, USA). Free label was washed off before incubation with varying concentrations of uhPR1A3 or ghPR1A3 (0.1 –10 μg ml⁻¹). Effector monocyte-derived macrophages were added (3 : 1–8 : 1 effector : target ratio), the cells were incubated for 1 h at 37°C and analysed using FACS Calibur to measure ADCC.

An alternative ADCP assay was developed using an automated microscope (Ikoniscope, Ikonisys, New Haven, CT, USA) (Ntourophi et al, 2008). Target cells and macrophages, stained as described above, were deposited on to a poly-L-lysine coated slide, fixed with 2% formaldehyde. The slides were washed with PBS containing Tween (PBS-Tween), covered with blocking solution and Goat anti-mouse–HRP (1 : 100 dilution in blocking solution) added as a secondary antibody. The slides were washed with PBS-Tween, before adding Tyramide-647 (1 : 100 dilution).

Phagocytosis events were detected using the Ikoniscope imaging system (Ntourophi et al, 2008). Slides are first scanned at low magnification (× 10) to detect green label and then revisited at high magnification (× 100) to identify red labelled macrophages with ‘engulfed’ green targets.

In vivo testing of PR1A3

Six to 12-week-old SCID/Beige mice were used for animal experiments. All experiments were performed after ethical approval from the Swiss Veterinary Office. LS174T cells were maintained in DMEM medium with 1% glutamine and 10% FCS (E4 complete medium). A murine model for CRC tumours was set up by intra-splenic injection of LS174T (3 × 10⁶ cells per mouse) under aseptic conditions (day 0). This resulted in the development of liver metastasis. Passive antibody protection was measured by comparing the survival of mice (n = 10 for each group) after intravenous injection of either glycoengineered humanised IgG1 PR1A3, or glycoengineered high affinity IgG1 anti-CEA antibody (clone SM3E) or vehicle PBS control. The SM3E antibody, unlike PR1A3, is not membrane CEA specific. Antibody injections were performed at days 7, 14 and 21 with a treatment dose of 25 mg kg⁻¹ of bodyweight of mouse. Termination criteria were in accordance with the Swiss Veterinary Office.

ADCC and ADCP data analysis

Percentage cell lysis in the cytotoxicity assays was calculated as [experimental release—background release]/[maximum release—background release] × 100. Antibody-dependent (specific) lysis was calculated as [experimental release—antibody-independent release]/[maximum release—antibody-independent release] × 100. The standard error of the mean of multiple experiments was calculated using Graphpad Prism software, San Diego, CA, USA. Percentage cell phagocytosis was calculated using the formula: number of dual-stain positive target cells (cells engulfed by macrophages) divided by the total number of target cells. Standard normal distribution tests were performed to test the significance of differences found.

RESULTS

The binding of uhPR1A3 and ghPR1A3 to SKCO-1, based on FACS analysis, was similar over a wide range of concentrations (0.001 – 100 μg ml⁻¹) (Figure 1; see Supplementary Figure S1). Glycoengineering the Fc segment of PR1A3 has, therefore, no effect on the binding efficiency of the antibody to its epitope on a CEA-expressing CRC line. The affinity of PR1A3 Fab for cell-bound CEA is approximately 10 nM.

The ADCC mediated activities of mPR1A3IgG1, mPR1A3IgG2a and uhPR1A3IgG1, using human effector cells and SKCO-1 as targets, are illustrated in Figure 2A and B. The data show that uhPR1A3IgG1 kills much more actively than either mPR1A3IgG1 or mPR1A3IgG2a. PR1A3 does not mediate any killing in the absence of immune cells (data not shown). These data confirm that hPR1A3 mediated killing depends on the appropriate Fc–FcγR interaction.

Using enriched NK cells as effectors and SKCO-1 as targets, the effects on hPR1A3 mediated ADCC of blocking with Fab2 and IgG antibodies against the three known human classes of FcγRs, CD64 (FcγRI), CD32 (FcγRII) and CD16 (FcγRIII), are shown in Figure 3. CD16 Fab2 blocks the ADCC somewhat more than CD16 IgG, whereas CD32 Fab2 had no significant effect. However, CD32 IgG
Figure 2 (A) Comparison of the ADCC activity of PR1A3 antibodies with differing Fc portions (murine IgG1, murine IgG2a and humanised IgG1). PBMC were used as effectors and SKCO-1 as the target (100:1 effector:target ratio). Antibodies were used at a final concentration of 10 μg ml⁻¹. The control was targets and effectors with no antibody. The P-values are for the significance of the differences between controls and the hlgG1PR1A3 results, based on t-tests. (B) Comparison of the ADCC activity of PR1A3 antibodies with differing Fc portions (murine IgG1, murine IgG2a and humanised IgG1) over a range of antibody concentrations from 1 to 50 μg ml⁻¹. PBMC were used as effectors and SKCO-1 as target (50:1, effector:target ratio).

Figure 3 Effect of FcγR blocking on unmodified humanised PR1A3-induced ADCC, using anti-CD16, 32 and 64 Fab or IgG. SKCO-1 was used as the target and NK cells as the effectors (effector:target ratio = 8:1). The P-values are for the significance of the differences between the result using hlgG1PR1A3 and no blocking antibody, based on t-tests.

Figure 5A shows that glycoengineered antibody, ghPR1A3, elicits ADCC at much lower concentrations than the unmodified form. Thus, just 1 μg ml⁻¹ of ghPR1A3 kills over 40% of SKCO-1 target cells, using unfractionated PBMC as effectors, whereas 1 μg ml⁻¹ of uhPR1A3 kills only about 10% of the targets, suggesting 10–100-fold increased effectiveness of the glycoengineered antibody. Figure 5B shows that 1 μg ml⁻¹ of ghPR1A3 is significantly more effective at mediating killing than is 1 μg ml⁻¹ uhPR1A3 over a range of effector to target ratios from 25:1 to 100:1, again using PBMC as effectors. Figure 5C shows that the difference between the killing efficiency of ghPR1A3 and uhPR1A3 is similar using effector cells from three different PBMC donors whereas Figure 5D shows that using enriched human NK cells as effectors, ghPR1A3 is significantly more effective than uhPR1A3 at killing SKCO-1 cells over a range of concentrations (1–20 μg ml⁻¹), as expected if NK cells are the main cell type mediating ADCC in PBMC. Figure 5E and F shows that ghPR1A3 was also significantly more effective than uhPR1A3 in killing the intermediate CEA-expressing cancer cell lines MKN45, and, particularly, LoVo, the latter even at an effector:target ratio of 25:1. As expected, no enhancement was seen for HCT116, a cell line that does not express CEA (data not shown).

To investigate macrophage-based ADCP (Munn et al, 1991; Watanabe et al, 1999; Lazar et al, 2006; McEarchern et al, 2007), monocytes were cultured to give rise to macrophages, which by FACS analysis were shown to be negative for CD3, CD15, CD19 and CD56, positive for CD11b, CD14, CD32, CD64 and HLA class II and weakly positive for CD16 (see Supplementary Figure S2a). The cultured monocytes adhere to the bottom of the plates and have an enlarged granular appearance, as expected for macrophages (see Supplementary Figure S2b). Freshly harvested macrophages were incubated with green CMFDA labelled SKCO-1, and a range

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of antibody concentrations using macrophage:target ratios from 3:1 to 8:1. A representative FACS analysis of the mixture of macrophages and target SKCO-1 cells after 1-h at 37° in the presence of an isotype control (PMSA), uhPR1A3 and ghPR1A3 antibodies (5.0 μg ml⁻¹) is shown in Figure 6A. The green-labelled target cells are in the right lower quadrant and the effectors,
labelled with PE conjugated anti-CD11b/CD14, in the left upper quadrant of the dot plots. After one hour’s culture in the presence of either uhPR1A3 or ghPR1A3 there were substantial numbers of red and green positive signals in the upper right quadrant, presumably representing macrophages containing engulfed target cells, in contrast to the results for the PMSA control (Figure 6A). The proportion of such signals is essentially the same for both uhPR1A3 and ghPR1A3, and is concentration dependent, as shown...
Figure 6  (A) Flow cytometric analysis of ADCP. SKCO-1 target cells were stained green with CMFDA and are present in the right lower quadrant of the dot plots. Macrophages were stained with anti-CD11b and CD14 conjugated with PE. They appear in the left upper quadrant of the dot plots. The left hand dot plot is from a representative 1-h culture of macrophages and target cells (SKCO-1) in the presence of IgG isotype control. The middle and right plots are from representative 1-h cultures of macrophages and target cells in the presence of uhPR1A3 and ghPR1A3, respectively (5 μg ml⁻¹). The effector: target ratio used was 5:1. (B) Effect of increasing concentrations of uhPR1A3 and ghPR1A3 on phagocytosis. Tumour targets were pre-incubated with an isotype control antibody (IgG, 10 μg ml⁻¹) or the variants of hPR1A3 at concentrations of 0.1–10 μg ml⁻¹. ADCP was determined by flow cytometric analysis as the percentage of targets in the upper right hand quadrant (see Figure 6C). The four graphs represent responses from four separate donors. (C) Effect of FcγR blocking on ADCP. Flow cytometry was used to calculate the percentage of tumour cell engulfment by cultured macrophages in the presence of 10 μg ml⁻¹ of uhPR1A3. Fab2 fragments were used to block either FcγR I (CD64), FcγR II (CD32) or FcγR III (CD16) (each antibody concentration was 1 μg ml⁻¹). The effector: target ratio used was 3:1, and the targets were SKCO-1. (D) Fluorescent images of macrophages phagocytosing SKCO-1 using the Ikoniscope. The macrophages (red) have been stained with anti-CD11b and anti-CD14 primary antibodies followed by goat anti-mouse-HRP and Tyramide 647. The target cell line (SKCO-1) was stained green with CMFDA (Celltracker probe). The left hand panels show microscope composite images viewed with FITC (green), Cy5 (for tyramide 647) and DAPI (blue) channels. The second, third and fourth panel column show the same cells viewed separately with the DAPI, green and Cy5 channels. Each represents a different phagocytic event.
in Figure 6B for four different donors. This indicates that, in contrast to ADCC, glycoengineering does not enhance ADCP. The apparent lower level of expression of FcγRIII (CD16) as compared with FcγRII (CD32) (see Supplementary Figure S2a) suggests that the density of surface membrane CD16 on macrophages may not be enough for glycoengineering of PR1A3 to enhanced ADCP.

The effects on FACS assayed ADCP of blocking with Fab2 fragments each of three classes of FcγRs receptor are shown in Figure 6C. Blocking FcγRII (CD32) or FcγRIII (CD16) reduced ADCP to some extent, whereas blocking FcγRI (CD64) had no effect. This indicates that both CD16 and CD32 are involved in ADCP, whereas previously only CD32 was thought to be mainly involved (Richards et al, 2008). The CD16 antibody (3G8 clone) used for blocking does not, however, distinguish between FcγRIIIa and FcγRIIIb. This suggests that the lack of an effect of glycoengineering on ADCP may be due to a lower concentration of FcγRIIIa than FcγRIIIb on macrophages, and the fact that the glycoengineering only enhances the binding to FcγRIIIa.

For the Ikoniscope-based ADCP assay target, SKCO-1 cells were labelled green with CMFDA and the macrophages red by combining anti-CD11a and anti-CD14 as primary antibodies followed by goat anti-mouse HRP and Tyramide 647 staining. Examples of fluorescent images of macrophages clearly engulfing the target cells, are shown in Figure 6D. In some cases, the macrophages clearly contain two nuclei, one of which is associated with the green stain of a target cell. Using the Ikoniscope it is therefore possible to count, separately, engulfed targets, conjugated targets, free targets and macrophages. Table 1 shows the relative percentage of presumed phagocytosis events seen with the FACS analysis compared with the percentage of confirmed phagocytosis events and percentage of conjugating events seen with the Ikoniscope. Combining engulfed and conjugated targets, as estimated using the Ikoniscope, there is reasonable agreement with the FACS analysis, which cannot distinguish simple attachment from actual engulfment. This suggests that the Ikoniscope-based ADCP assay gives more accurate results than that based on FACS analysis, which is the basis of previous publications reporting ADCP (Munn et al, 1991; Huls et al, 1999; Akewanlop et al, 2001; Lazar et al, 2006; McEarchern et al, 2007). The quantitative data of Table 1 also suggest that the glycoengineered

|                  | PMSA control | ghPR1A3 | unPR1A3 |
|------------------|--------------|---------|---------|
| Total targets    | 1336.00      | 1284.00 | 1428.00 |
| Targets engulfed by macrophages | 27.00        | 122.00  | 213.00  |
| Conjugates between targets and macrophages | 38.00        | 79.00   | 116.00  |
| % of targets engulfed | 2.02         | 9.50    | 14.92   |
| % of targets conjugated | 2.84         | 6.15    | 8.12    |
| % phagocytosis+% conjugated | 4.87         | 15.65   | 23.04   |
| % ADCP using FACS | 10.22        | 22.11   | 28.56   |

The number of target cells that were engulfed by macrophages was counted directly using the Ikoniscope. Targets that were conjugated to macrophages were defined as having cell-to-cell membrane contact but no envelopment. Targets and macrophages could readily be identified separately. The percentages of engulfed and conjugated cells are compared with the percentage of targets classified as having undergone ADCP using FACS analysis of the same preparation of cells.
Efficacy of glycoengineered IgG1 PR1A3

SQ Ashraf et al

Antibody engineering to enhance ADCC can be achieved either by altering the amino acid structure of the Fc backbone or by modifying the carbohydrate structures at the hinge region (glycoengineering) to improve the binding affinity to Fcγ receptors (Umana et al., 1999; Niwa et al., 2004b; Lazar et al., 2006). Three glycoengineering approaches have been used (i) over expression of the enzyme GntIII in the antibody producing cells, which results in the Fc segment containing an increase in bisected non-fucosylated oligosaccharides, (ii) producing antibody in CHO cells that lack the transferase enzyme involved in core-fucosylation and (iii) using siRNA to knockdown fucosyl transferase activity. Over expression of GntIII has been used successfully to increase the in vitro ADCC potency of antibodies IGN311 (Lewis Y-specific) and chCE7 (anti-neuroblastoma) (Umana et al., 1999; Schuster et al., 2005). Using this technique, we have shown that the ADCC activity of the humanised IgG1 anti-CEA antibody PR1A3 can be increased more than 10-fold. This effect is seen in both intermediate and high expressing CEA cell lines. The level of enhancement may be pronounced even at low effector:target ratios in intermediate expressing CEA cell lines. This underlines the potency of ghPR1A3, which may be particularly relevant considering that antibody penetration into tumours may limit therapeutic capacity. Therefore, it is of paramount importance to show efficient antibody killing by human effector cells. This ADCC activity is still dependent on CD16 (FcγRIIIa) on the NK cells, as shown by Fab2 anti-CD16 blocking and enhanced ADCC of ghPR1A3 over uhPR1A3 in the presence of enriched NK cells. The improved ADCC activity of the glycoengineered antibody is achieved without affecting PR1A3’s binding activity, leaving it membrane specific, as described earlier (Durbin et al., 1994; Stewart et al., 1999). This is an important property of PR1A3, given the finding that soluble CEA can accumulate in lymph nodes and lead to false positive detection of cancers in lymph nodes when using other anti-CEA antibodies for immunoscintigraphy (Goldnowski et al., 1989). In contrast to NK cells, neither freshly isolated granulocytes nor monocytes show any significant ADCC activity at comparable antibody concentrations even with high effector:target ratios. The lack of monocyte-based ADCC activity may be due to the low level of CD16 found in our preparations. We have shown, however, using two different assays, that macrophages kill target tumour cells in vitro by ADCP. Both of these mechanisms are therefore likely to have a significant function in in vivo patient responses to naked antibody therapy. Glycoengineering substantially improves antibody PR1A3’s NK cell mediated ADCC activity, but apparently has no effect on in vitro macrophage-based ADCC. This suggests that the FcγRIIIa receptor is not the effective Fc receptor on macrophages for ADCC and opens up the possibility of alternative engineering of antibodies to enhance their ADCP activity (Richards et al., 2008). However, the expression of FcγRs on macrophages is dependent on the cytokines in the tumour microenvironment, and so the in vivo assay may not reflect the in vivo situation, and this needs further investigation. The in vivo data presented here show that glycoengineered humanised IgG1 PR1A3 is effective in prolonging survival in a murine CRC metastatic model. This is thought to work by interaction between the Fc segment of the antibody and FcγRIV, the murine homologue of human FcγRIIIa. These latter receptors are found on macrophages and granulocytes in mice, emphasising the subtle differences between the mouse and human immune systems. Interestingly, despite the hugely lower affinity for CEA of PR1A3 as compared with SM3E, the improvement in survival was at least as great, if not greater with PR1A3 as with SM3E. This contrasts to the findings in vitro, in which glycoengineered SM3E has a much higher ADCC capacity than PR1A3. The in vivo finding may be explained by the distinctive specificity of PR1A3 for binding to membrane-bound CEA and scavenging of SM3E by shed CEA. Binding to soluble CEA, leading to sequestration of antibody–antigen complexes away from the tumour, is a major obstacle for any potential therapeutic anti-CEA antibody.

The glycoengineered variant antibodies have been shown to have substantially improved binding to the lower affinity FcγRIIIa receptor polymorphic variants, which have phenylalanine instead of valine at amino acid position 158 (Ferrara et al., 2006). In addition, the carbohydrate residue at Asn-162 of FcγRIIIa has also been shown to have an important function in the binding of glycoengineered FcγRIIIa (Richards, 2006). In contrast to NK cell receptors, this does not greatly attenuate affinity for these antibody variants (Ferrara et al., 2006). As only 10–15% of patients have the high affinity FcγRIIIa receptor polymorphic variants, use of the glycoengineered variant antibody should make this therapy accessible to all patients, whatever their genetic constitution with respect to these FcγRIIIa receptor polymorphic variants (Cartron et al., 2002; Weng and Levy, 2003; Niwa et al., 2004a,b; Schuster et al., 2005).

Earlier studies on macrophage mediated in vitro phagocytosis have relied heavily on FACs analysis using PKH lipid linkers to stain target cells (Munn et al., 1991; Huls et al., 1999; Akewalonp et al., 2001; Lazar et al., 2006; McEarchern et al., 2007; Richards et al., 2008). These linkers can, however, diffuse non-specifically from cell to cell, giving rise to false positive signals and thus limiting...
their use for phagocytosis assays. The use of a cytoplasmic stain, coupled with the Ikoniscope, shows unequivocally that cultured macrophages are able to phagocytose tumour cells in the presence of PR1A3, and enables satisfactory quantitation of the ADCP assay.

The low relative ADC activity of the murine IgG2a or IgG1 Fc domains on human cells is consistent with the dependence of ADC on the specificity of the binding of the Fc portion of an antibody to the relevant Fc receptor (FcγRIIIa) on NK cells. This may, in part, explain the poor results obtained in early oncology trials using naked murine monoclonal antibodies. On the basis of the blocking experiments with Fab2 of CD32 and CD64, these receptors have little if any role in NK mediated ADC. However, intact anti-CD32 antibody (clone IV.3; IgG2b), in contrast to the Fab2 fragment (clone 7.3), is as at least as potent as intact anti-CD16 in inhibiting both ADC and antibody-independent killing. This may be explained by low levels of CD32c on NK cells (present in 40% of donors; Metes et al, 1998) that may lead to self-killing or a secondary cross-linking effect between FcγRs.

NK-based ADC and macrophage-based ADCP are the most likely mechanisms for naked anti-CEA antibody therapy (Munn et al, 1999; Huls et al, 1999; Watanabe et al, 1999; Akewanlop et al, 2001; McEarthern et al, 2007), as there is no obvious basis for a functional blocking effect involving CEAs. In addition, macrophages are known to be potent antigen-presenting cells and so, following antibody stimulated engulfment of whole tumour cells, macrophages may digest the engulfed cells and then re-present peptide fragments to T cells. This may then enable the activation of the adaptive immune system against tumour cell products (Adams and Weiner, 2005; Carter, 2006). It is also likely that dendritic cells, which are super antigen presenters and express FcγRs, are stimulated by antibodies to engulf target cells, which would significantly enhance their ability to present tumour antigens to the adaptive immune system (Kalergis and Ravetch, 2002; Benitez-Ribas et al, 2006; Melief, 2008). Glycoengineering and other modifications of antibodies to enhance their immune effector functions appear, therefore, to be a most effective way to improve the efficacy of naked antibody therapy by a variety of mechanisms.

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Conflict of interest

PU, EM and PB are employed by Glycart, Roche. Glycart have provided CIL with antibodies and a grant partly covering laboratory expenses. There is a commercial agreement between CIL (WFB) and Glycart, Roche, for the development of PR1A3.

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**Efficacy of glycoengineered IgG1 PRA13**

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