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Activating Fc γ receptors contribute to the antitumor activities of immunoregulatory receptor-targeting antibodies

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Fc γ receptor (FcγR) coengagement can facilitate antibody-mediated receptor activation in target cells. In particular, agonistic antibodies that target tumor necrosis factor receptor (TNFR) family members have shown dependence on expression of the inhibitory FcγR, FcγRIIB. It remains unclear if engagement of FcγRIIB also extends to the activities of antibodies targeting immunoregulatory TNFRs expressed by T cells. We have explored the requirement for activating and inhibitory FcγRs for the antitumor effects of antibodies targeting the TNFR glucocorticoid-induced TNFR-related protein (GITR; TNFRSF18; CD357) expressed on activated and regulatory T cells (Treg cells). We found that although FcγRIIB was dispensable for the in vivo efficacy of anti–GITR antibodies, in contrast, activating FcγRs were essential. Surprisingly, the dependence on activating FcγRs extended to an antibody targeting the non–TNFR receptor CTLA-4 (CD152) that acts as a negative regulator of T cell immunity. We define a common mechanism that correlated with tumor efficacy, whereby antibodies that coengaged activating FcγRs expressed by tumor-associated leukocytes facilitated the selective elimination of intratumoral T cell populations, particularly Treg cells. These findings may have broad implications for antibody engineering efforts aimed at enhancing the therapeutic activity of immunomodulatory antibodies.

Activating Fc γ receptors (FcγRs) stimulate immune cell effector mechanisms, such as antibody-dependent cell-mediated cytotoxicity (ADCC) and phagocytosis (ADCP), which combine to facilitate antibody-mediated tumor cell killing (Nimmerjahn and Ravetch, 2008; Hogarth and Pietersz, 2012). The importance of FcγR-mediated immune effector cell function has been demonstrated in preclinical efficacy studies for antibodies targeting a range of tumor cell–expressed receptors, including trastuzumab (HER2) and rituximab (CD20; Clynes et al., 2000; Nimmerjahn and Ravetch, 2012). The inhibitory FcγR, FcγRIIB, functions to modulate activating FcγR-mediated effector mechanisms in immune cells that coexpress both FcγR classes, such as macrophages and dendritic cells. FcγRIIB has recently been implicated in augmenting antibody-mediated receptor forward signaling through a mechanism of cross-linking in target cells expressing the TNF receptor (TNFR) family members TNFRSF10, TNFRSF10B (DR4 and DR5, respectively), and TNFRSF5 (CD40; Wilson et al., 2011; Li and Ravetch, 2012). It remains unclear what contribution FcγR biology has in the modality of antibody therapeutics that target other cell surface receptors. In particular, the emerging clinical benefit of agonistic antibodies targeting the T cell–APC interface raises the possibility that FcγR coengagement may contribute to their in vivo mechanism of action (Mellman et al., 2011).
Preclinical studies in mice using agonistic antibodies targeted to glucocorticoid-induced TNFR-related protein (GITR)—a costimulatory TNFR expressed by regulatory and activated T cells—have shown compelling antitumor activity in syngeneic mouse tumor models (Turk et al., 2004; Ko et al., 2005). In vitro, stimulation of GITR with agonist antibodies can induce forward signaling into T cells, which promotes proliferation and cytokine production (Kanamaru et al., 2004; Ronchetti et al., 2007). In vivo, several mechanisms have been proposed to contribute to the antitumor activity of antibodies targeting GITR; however, the current paradigm stipulates that agonist properties of these antibodies promotes cytotoxic effector T cell generation, while dampening the immunosuppressive effects by FoxP3+ CD4+ T reg cells (Ronchetti et al., 2012; Schaer et al., 2012). The recent findings that antibodies targeted to TNFR family members require FcγRIIB interaction for their in vivo activities led us to explore a common mechanism for antibodies targeting TNFRs expressed by T cells, using GITR to test this paradigm.

RESULTS AND DISCUSSION
Activating, but not inhibitory, FcγRs are necessary for antitumor activity of a GITR-targeting antibody
To evaluate the contribution of activating or inhibitory FcγRs in the mechanism of tumoricidal activity of an agonist antibody targeting GITR (clone DTA-1, rat IgG2b), Colon26 colorectal cancer cells were implanted subcutaneously in wild-type, FcγRIIB−/−, or Fc common γ chain−/− mice. The common γ chain cofactor is required for assembly and membrane expression of the activating FcγRs I, III, and IV (Nimmerjahn and Ravetch, 2008). Mice with preformed tumors (∼70 mm³) were treated with a single dose of the anti-GITR antibody (clone DTA-1) or a rat IgG2b isotype control. As previously shown for this tumor model, DTA-1–mediated single dose regressions in 100% of wild-type mice (Fig. 1 A; Zhou et al., 2007). In contrast to recent reports studying anti-TNFR antibodies targeting DR4, DR5, or CD40, the antitumor efficacy of DTA-1 was independent of FcγRIIB expression (Fig. 1 B; Wilson et al., 2011; Li and Ravetch, 2012). Instead, activating FcγRs were required for the tumoricidal activity of a GITR-targeting antibody (Fig. 1 C).

Co-engagement of FcγRs by DTA-1 is required for optimal antitumor activity
To further examine the contribution of activating FcγRs for the tumoricidal activity of antibodies to GITR, we generated two chimeric antibodies from the parental DTA-1 rat IgG2b: a murine IgG2a (mIgG2a), and mIgG2a with a N297A mutation that eliminates binding to all murine FcγRs (not depicted; Shields et al., 2001; Chao et al., 2009; Wilson et al., 2011). Binding to in vitro–stimulated splenic T cells was conserved for the murine IgG2a DTA-1 variants and was comparable to parental DTA-1 rat IgG2b (Fig. 2 A and B). All versions of DTA-1 showed a similar ability to induce NF-kB signaling in a reporter cell assay (Fig. 2 C). Moreover, the three DTA-1 variants similarly enhanced anti-CD3-mediated T cell proliferation and cytokine production in vitro (Fig. 2 D and not depicted). Both DTA-1 rIgG2b and mIgG2a showed potent antitumor efficacy, which was abrogated in absence of activating FcγRs but was not impacted by the absence of FcγRIIB expression (Fig. 1 and Fig. 2 E). Eliminating FcγR interaction through the N297A mutation abrogated the antitumor activity of DTA-1 (Fig. 2 F). The three versions of DTA-1 showed similar GITR saturation properties in the tumor and draining lymph, supporting common pharmacokinetic/pharmacodynamic properties.
Figure 2. Engagement of FcγRs by DTA-1 is required for antitumor activity. (A and B) GITR-binding assay. Primary splenocytes stimulated with CD3- and CD28-specific antibodies served as targets. DTA-1 variants were detected using rat IgG2b-specific (A) or murine IgG2a-specific (B) PE-conjugated antibodies. (C and D) In vitro activity of GITR-specific antibodies at various concentrations tested on GITR-expressing NF-κB 293 reporter cell line (C) and splenocytes incubated with suboptimal doses of anti-CD3 and anti-CD28 antibodies (D). The in vitro data are derived from triplicates and are a
of these variants (Fig. 2 G). Collectively, our results support that agonist antibodies targeting GITR require co-engagement with activating FcγRs for their tumoricidal activities in this model.

**Co-engagement of FcγRs by DTA-1 results in intratumoral loss of T cells**

To further understand the underlying FcγR-dependent mechanism of DTA-1, we analyzed the immune cell populations in tumor and draining lymph node after administration of the DTA-1 variant antibodies, focusing on early time points before tumors typically begin to regress. First, we confirmed that GITR was expressed on T cells in the draining lymph node and tumor (Fig. 3 A; Shimizu et al., 2002). The highest expression of GITR was by T cells in the tumor compared with the draining lymph node, with T reg cells in the tumor showing approximately fourfold stronger signal than the equivalent lymph node population. Next, we profiled myeloid and NK cells for expression of FcγRIII and IV (Fig. 3, B and C). Tumor-associated myeloid and NK cell populations were abundant in Colon26 tumors (>50% of all CD45+ leukocytes) and expressed activating FcγRIII/FcγRIV and FcγRIII, respectively. FcγRIII was distinguished from FcγRIIB using mice deficient for FcγRIIB in combination with the 2.4G2 antibody, which cross-reacts with FcγRIIB and III. In contrast, the same innate immune cell populations were underrepresented in the tumor-draining lymph node (<0.5% of all leukocytes; Fig. 3 B). Treatment with the parental or mlgG2a DTA-1 variant resulted in a strong reduction in the percentage of intratumoral FoxP3+ T reg cells (Fig. 3 D). This effect was specific to the tumor, with no significant change in the T reg cell population in draining lymph nodes. In contrast, treatment with the DTA-1–N297A variant did not alter intratumoral T cell populations, supporting dependence on FcγR interactions for this effect. To quantify the change in T cell populations after DTA-1 variant treatment, we monitored the density of T cells in the tumor over 5 d. By 24 h, the parental rat IgG2b or DTA-1 mlgG2a mediated a dramatic reduction in the density of FoxP3+ T reg cells in the tumor, which did not occur in the DTA-1–N297A–treated cohort (Fig. 3 E). Although the loss of intratumoral T cells was most dramatic at early time points in the T reg cell population, CD4+ T cells were also significantly reduced on days 3 and 5 after antibody treatment (Fig. 3 F). A slight reduction in the density of CD8+ T cells was also observed, particularly in the DTA-1 mlgG2a cohort. The depletion of T reg cells was most pronounced, which resulted in a significant shift in the ratio of CD8+ T cells to T reg cells in the tumor early after treatment (Fig. 3 G). This finding correlated with previous reports showing an obvious shift in the ratio of CD8+ T cells to T reg cells in the tumor, although in the previous reports, tumors were extracted at later time points after the onset of regression (Ko et al., 2005; Sharma et al., 2008; Cohen et al., 2010). Importantly, DTA-1 treatment did not affect the overall density of tumor-associated leukocytes, supporting that T cell elimination was a specific event (Fig. 3 H). Again, the cellularity of T cells in the tumor-draining lymph node, as well as the ratio of CD8+ T cells to T reg cells, remained mostly unchanged (Fig. 3 I and not depicted). Together, our data reveal that DTA-1–mediated an FcγR–dependent loss of T cells in the tumor, with the magnitude of depletion correlating with cell surface expression of GITR (T reg→FoxP3→CD4+→CD8+). Furthermore, we demonstrate that the loss of intratumoral T cells precedes tumor regression, supporting that this depletion mechanism may serve as an initial event that culminates in tumor rejection.

**Activating, but not inhibitory, FcγRs are required for intratumoral T reg cell depletion**

To evaluate if the depletion of intratumoral T cells required activating FcγR expression, we monitored T cell populations in wild-type, FcγRIIB–, and Fc common γ chain–deficient mice after treatment with DTA-1–mlgG2a. Murine IgG2a DTA-1 mediated potent T reg and non–T reg cell elimination in both wild-type and FcγRIIB–deficient animals, which resulted in a shift in the ratio of CD8+ T cells to T reg cells in the tumor, whereas the ratio remained unchanged in the draining lymph node (Fig. 4, A and B). In contrast, no significant depletion or shift in T cell ratio was observed in mice devoid of activating FcγR expression (Fig. 4 C). GITR expression on T cell populations isolated from tumors and lymph node was conserved between the three strains and was highest among intratumoral T reg cells (Fig. 4, D and F). These data support the notion that treatment with antibodies targeting GITR–expressing tumor-infiltrating T cells leads to activatory FcγR–dependent elimination, the degree of which correlates with levels of GITR expression.

**Activating FcγRs are also required for the antitumor activities of an antibody targeting the non–TNFR antigen CTLA-4**

With the unexpected dependence on activatory FcγRs for DTA-1–mediated antitumor activity, we next sought to understand if this finding was restricted to a T cell–expressed immunostimulatory TNFRs. To evaluate this, we chose an antagonist antibody (clone 9D9, murine IgG2b) targeting the T cell antigen CTLA-4, which functions to negatively regulate T cell proliferation upon signaling (Leach et al., 1996; Curran et al., 2010). Consistent with earlier reports, a single representative of two or more independent experiments. (E) Efficacy study of 5 mg/kg DTA-1 mlgG2a in wild-type, FcγRIIB−/−, and Fc common γ chain−/− mice bearing Colon26 tumors (n = 7–10 mice per treatment group). Mean and standard errors are based on triplicates, and the data is a representative of two or more independent experiments. (F) In vivo efficacy study after treatment with the DTA-1 variant antibodies (n = 7). The efficacy data are a representative of two or more independent experiments. (G) Saturation of GITR on T reg cells in the tumor and draining lymph node by the three versions of DTA-1.
intrapertitoneal injection of 9D9 mediated tumor regressions in 100% of animals (Fig. 5 A; Leach et al., 1996; Curran et al., 2010). To explore the involvement of FcγR in anti–CTLA-4–mediated antitumor efficacy, we evaluated tumoricidal activity of anti–CTLA4 in mice devoid of activating FcγRs (Fig. 5 A). Similar to what we observed in DTA-1–treated animals, 9D9 efficacy was strictly dependent on activating FcγRs. Similar to the expression profile of GITR, CTLA-4 was expressed by intratumoral CD4+ T cells, most notable the T reg cell population (Fig. 5 B). 9D9 administration resulted in the
specific depletion of tumor-associated T cells and a shift in the ratio of CD8+ to T reg cells, which was dependent on activating FcγR expression (Fig. 5 C). Together, these data support a common antitumor mechanism shared between antibodies targeting receptors highly expressed at the surface of intratumoral CD4+ T cells, which requires the function of activating FcγRs.

Here, we have shown that antibodies targeting two functionally distinct immunoregulatory receptors on T cells require the coengagement of activatory FcγRs to mediate their antitumor effect. A single dose of the GITR targeting antibody DTA-1 mediated the rapid and selective elimination of T cells within the tumor microenvironment, particularly those of the T reg cell lineage, as defined by intracellular FoxP3 expression. The dependence on the expression of activating FcγRs for T cell depletion appears consistent with involvement of the immune effector cell mechanisms ADCC or ADCP. Indeed, tumor-associated immune effector cells expressing activating FcγRIII and IV were abundant in Colon26 tumors and are known to mediate ADCC or ADCP effects in other model systems (Nimmerjahn and Ravetch, 2005; Albanesi et al., 2012). In general, the depletion of T cell populations correlated with the overall level of target antigen expression, although with anti–CTLA-4 treatment this observation was less clear with an overall reduction of intratumoral CD4+ T cells on day 5. We reconcile these observations with the ability of activatory FcγR-expressing immune effector cells to elicit ADCC, based on a receptor threshold model (Lewis et al., 1993; Niwa et al., 2005). In contrast, few myeloid or NK cells were present in the draining lymph nodes, and the expression of GITR and CTLA-4 on FoxP3+ T reg cells in this compartment was lower than in the tumor. An alternative hypothesis is that agonist GITR signaling mediated by DTA-1 could alter the stability of the transcription factor FoxP3 expressed by intratumoral CD4+ T cells, which requires the function of activating FcγRs.

Figure 4. Activating, but not inhibitory, FcγRs are required for intratumoral T reg cell depletion by antibodies targeting GITR. Intratumoral T cell density and CD8+ T cells to T reg cells ratios 5 d after treatment with 5 mg/kg DTA-1-mIgG2a using wild-type (A), FcγRIIB−/− (B), or Fc common γ chain−/− (C) mice bearing Colon26 tumors. (D–F) Cell surface expression of GITR on T cells. Mean and standard errors are based on triplicates from two independent experiments. P-values were calculated using an unpaired Student’s t test (**, P < 0.001).
The requirement for activating FcγR coengagement of antibodies targeting tumor associated targets, such as HER2 or CD317 (Junttila et al., 2010; Nordstrom et al., 2011; Tai et al., 2012). Interestingly, ipilimumab, an antagonist antibody targeting CTLA-4 with human IgG1 framework, has shown efficacy in metastatic melanoma patients. Human IgG1 antibodies have an intrinsically higher activating to inhibitory (A/I) FcγR ratio, which is proposed to favor ADCC effector mechanisms (Hodi et al., 2010; Robert et al., 2011; Hogarth and Pietersz, 2012). In contrast, another antagonist antibody targeting CTLA-4, tremelimumab, is a human IgG2 antibody and has a lower A/I ratio. This antibody failed to reach its endpoints in a clinical trial (Chung et al., 2010). Although these two antibodies remain to be compared directly in patients, it is tempting to speculate that the varying ability of IgG1 versus IgG2 to bind human activating FcγRs may explain, at least in part, the apparent difference in clinical response. Accordingly, Fc-optimized variants of ipilimumab, such as mutations designed to enhance ADCC or ADCP, could also be considered for clinical evaluation. In parallel, a prospective or retrospective study to analyze patients carrying the allelic variants of the activating FcγRIIA and III could be valuable, particularly given the striking correlation with progression-free survival in patients treated with Rituximab (Weng and Levy, 2003). In these settings, a challenge will be to design relevant preclinical models to evaluate the efficacy of immunomodulatory antibodies that better correlate clinical response, and our findings provide a framework to test this with future immunomodulatory antibodies. For instance, mice engineered to express the human FcγR system might be of great utility (Smith et al., 2012).
sequences of the parental antibody were cloned from the hybridoma and inserted into a pR5sA derivative vector containing publicly available sequences for the constant domain region of murine IgG2a to create the chimeric antibody DTA-1-mlgG2a. The N297A mutation was introduced into the DTA-1-mlgG2a-encoding vector by site-directed mutagenesis to generate DTA-1-N297A. DTA-1 antibodies were produced from HEK 293 FreeStyle cells (Invitrogen) transfected with appropriate expression vectors and purified using Fast Flow rProtein A Sepharose (GE Healthcare), followed by size exclusion chromatography. The integrity of the antibodies was verified by SDS-PAGE and analytical SEC. The LTF-2 (control; rat IgG2b, C1.18.4 (control; mouse IgG2a), DTA-1 (GITR; rat IgG2b), and 9D9 (CTLA-4; mouse IgG2b) antibodies used for in vivo studies were purchased from BioXCell. Before injection in vivo, antibodies were cleared from precipitates by centrifugation and confirmed to contain endotoxin levels below 1 EU/ml of antibody (LAL assay). For in vivo studies, tumor-bearing mice were typically treated 8–9 d after tumor inoculation (~70 mm3) by intraperitoneal injection of antibodies at indicated doses.

Tumor dissociation and analyses by flow cytometry. CD45−, CD25−, CD4−, CD8−, and FcγRIII/IV (clone 2.4G2)-specific and matching isotype control antibodies were purchased from BD.BIOTEK-, CD11b−, DX5−, CTLA-4−, and Foxp3−specific antibodies were purchased from eBioScience, CD3−specific antibody from BioLegend, and FcγIV-specific antibody from Sino Biologicals. To obtain single cell suspensions from excised tumors, the tumors were first minced, dissociated with collagenase (Life/Roche) and DNase I (Life Technologies), filtered on a 70-µM sieve, and then treated with RBC lysis buffer (eBioScience) before washes with PBS + 2% FBS. For most stainings, cells were incubated with saturating doses of anti-CD16/32 FcγR block (BD) before incubation with fluorochrome-conjugated antibodies. For FcγRII/III staining, the directly conjugated 2.4G2 antibody was incubated before FcγR block. For staining of intracellular markers, extracellular markers were stained first, before fixation/permeabilization of the cells (eBioScience), followed by staining of intracellular proteins using primary-conjugated antibodies. During staining procedure, cells were maintained on ice. Acquisition was performed on an LSR-II flow cytometer (BD). The machine performances were verified daily using Cytometer Setup and Tracking beads (BD), and weekly using Sphero Rainbow Fluorescent I Peak particles (BD) and AccuCount Blank beads (Spherotech, Inc). For each analysis, the population of interest was gated on live leukocytes using a combination of morphological parameters, CD45−specific labeling and dead cells exclusion using 7AAD (BD Biosciences), or Live/Dead yellow (Life Technologies). The cell density in the tumor was calculated by dividing the number of cells of interest to the total number of cells extracted. Before each run, a compensation matrix was generated from primary cells stained separately with individual fluorochrome-conjugated antibodies. P-values were calculated using an unpaired Student’s t test.

NF-κB activation reporter assay. The 293-NFκB-luc cell line was generated by stable transfection of HEK293 cells with pNFκB-luc plasmid (Takara Bio Inc.), which had been modified to express the Zeocin resistance gene. GITR sequence was cloned from cDNA (Life Technologies), introduced into a pCDNA3 expression plasmid (Invitrogen), and transfected into 293-NFκB-luc cell line to generate the 293-GITR-NFκB-luc reporter cell line. 105 cells/ml cell cultures were incubated with indicated concentrations of GITR−specific antibodies and incubated at 37°C for 24 h. NF-κB activation levels were indirectly determined by measuring NF-κB−induced luciferase activity using CellBright-Glo (Promega).

In vitro T cell stimulation assays. CD3− and CD28−specific antibodies were purchased from R&D Systems. 5 × 105 splenocytes were stimulated with 0.1 µg/ml CD3− and 0.2 µg/ml of CD28−specific antibodies together with indicated concentrations of GITR−specific antibodies in round-bottom culture−treated 96-well plates. After 72 h of incubation, cell proliferation was indirectly determined using CellTiter-Glo (Promega).

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