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RGMb is a novel binding partner for PD-L2 and its engagement with PD-L2 promotes respiratory tolerance

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Programmed death 1 (PD-1, CD279) and its ligands PD-L1 (B7-H1, CD274) and PD-L2 (B7-DC, CD273) are key inhibitory molecules in immune regulation (Keir et al., 2008; Pardoll, 2012). This pathway provides particularly promising targets for cancer immunotherapy (Topalian et al., 2012). There is considerable evidence that PD-L2 inhibits immunity by binding to the PD-1 co-inhibitory receptor (Latchman et al., 2001; Zhang et al., 2006). However, studies have shown that PD-L2 can function to stimulate T cell proliferation and cytokine production, even in PD-1–deficient T cells or with PD-L2 mutants that did not bind to PD-1 (Liu et al., 2003; Shin et al., 2003; Wang et al., 2003). These findings suggest that PD-L2 may function through a receptor other than PD-1. Most studies using blocking mAbs show a dominant role for PD-L1 in inhibiting immune responses; however, PD-L2 plays a dominant role in responses such as airway hypersensitivity, experimental allergic conjunctivitis and nematode infection (Ritprajak et al., 2012). In some situations, PD-L2 dominance may be explained by preferential PD-L2 up-regulation by IL-4, but other instances may be explained by the binding of PD-L2 to a receptor other than PD-1.

Here, we demonstrate that PD-L2 binds to a second receptor, repulsive guidance molecule b (RGMb). RGMb, also known as DRAGON, is a member of the RGM family which consists of RGMa, RGMb, and RGMc/hemojuvelin (Severyn et al., 2009). RGMs are glycosylphosphatidylinositol-anchored membrane proteins that bind bone morphogenetic proteins (BMPs) and neogenin (Conrad et al., 2010). RGMs do not bind to PD-1, which inhibits antitumor immunity, and to RGMb, which regulates respiratory immunity, targeting the PD-L2 pathway has therapeutic potential for asthma, cancer, and other immune-mediated disorders. Understanding this pathway may provide insights into how to optimally modulate the PD-1 pathway in cancer immunotherapy while minimizing adverse events.

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RESULTS

RGMb binds to PD-L2, but not PD-L1 or other related molecules

We identified RGMb as a novel binding partner for PD-L2 using COS cell expression cloning with PD-L2-Ig fusion protein. Using flow cytometry with stably transfected 300 cells and Ig fusion proteins, we found that mRGMb binds to mPD-L2 but not mPD-L1 or other proteins of the B7 family (Fig. 1, a and b). ELISA with purified proteins showed that mRGMb binds to mPD-L2 and hPD-L2, and that hRGMb binds to hPD-L2 and mPD-L2 (Fig. 1 c and not depicted). Thus, the RGMb–PD-L2 interaction occurs in both mice and humans. Further studies showed that PD-L2 does not bind to RGMa or RGMc (Fig. 1 d). Biacore data showed that PD-L2 bound to RGMb with a similar affinity as to PD-1, $K_d = 48.5$ and 58.8 nM, respectively (Fig. 1 e).

Figure 1. RGMb binds to PD-L2, but not PD-L1 or other related molecules. (a and b) mRGMb- or mPD-L2-transfected 300 cells, untransfected 300 cells, and PD-1-transfected Jurkat T cells were stained with the indicated Ig fusion proteins (red) or control-Ig (blue) and analyzed by flow cytometry. MFI, mean fluorescence intensity. (c and d) Binding of mPD-L2-Ig or control-Ig to plates coated with mRGMb-HIS, hRGMb-HIS, mRGMa-HIS or mRGMc-HIS, was analyzed by ELISA. (e) Biacore analysis of the interaction of RGMb with PD-L2 and of PD-1 with PD-L2. (f–k) Cell–cell binding of the indicated transfected cells was analyzed by cell conjugation assay. In f, h, and j, the top panels show the FSC-SSC plots and the bottom panels show the corresponding dot plots. g, i, and k show only the dot plots. All data are representative of 2–11 independent experiments.
To test the RGMb–PD-L2 interaction under more physiological conditions, we used a cell conjugation assay where one transfected cell was labeled with a red dye and the other transfected cell with a green dye. The binding of the two cells was assessed by flow cytometry and indicated by double positive events (yellow dots). As expected, mRGMb cells bound to mPD-L2 cells (Fig. 1 f), but not mPD-L1 cells (Fig. 1 g). As positive controls, mPD-1 cells bound to both mPD-L2 and mPD-L1 cells (Fig. 1, h and i). Negative control binding assays had few conjugates (<0.3%; Fig. 1, j and k; and not depicted). These results show that mRGMb binds specifically to mPD-L2, but not to mPD-L1, and the structural orientation is compatible with RGMb and PD-L2 cell surface-to-cell surface binding.

Characterization of RGMb and PD-L2 mAbs

We characterized a panel of mRGMb mAbs and identified 6 RGMb mAbs showing specificity by flow cytometry on mRGMb- and/or hRGMb-transfected 300 cells, ELISA, and Western blotting (unpublished data). We identified two mRGMb mAbs, 9D1 and 8B2, that blocked interaction of RGMb with PD-L2 (Fig. 2 a). These RGMb antibodies do not bind to mRGMa or mRGMc (unpublished data).

We also determined whether mPD-L2 mAbs would block PD-L2 interactions with RGMb. We identified one mPD-L2 mAb, 2C9, that blocked PD-L2–RGMb but not PD-L2–PD-1 interactions, and three mPD-L2 mAbs, 3.2, TY25, and MIH37, that blocked both PD-L2–RGMb and PD-L2–PD-1 interactions (Fig. 2, b and c). TY25 and 3.2 share the same epitope, whereas 2C9 recognizes a different epitope (unpublished data). The existence of both single and double blocker PD-L2 mAbs suggests that the PD-1- and RGMb-binding sites on PD-L2 are close but distinct. Properties of RGMb and PD-L2 mAbs are summarized in Table I.
(Fig. 2 d and not depicted). These data suggest that the binding sites on RGMb for PD-L2 and BMP are close but distinct.

To test if RGMb can bind both PD-L2 and BMP at the same time, we performed an ELISA to analyze the binding of PD-L2-Ig fusion protein to immobilized BMP-2/4 in the presence or absence of RGMb. PD-L2 could not directly bind to BMP-2/4, but in the presence of RGMb could form a complex with BMP when RGMb and PD-L2-Ig were added simultaneously or sequentially to BMP-2/4 (Fig. 2 e and not depicted). These data are consistent with RGMb having distinct sites for PD-L2 and BMP binding, and show that RGMb has the capacity to form a trimeric complex with BMP and PD-L2.

RGMb has also been reported to bind to neogenin (Conrad et al., 2010; Bell et al., 2013). Furthermore, neogenin has been shown to directly bind BMP-2/4/6/7, and to modulate BMP signaling (Hagihara et al., 2011; Tian and Liu, 2013). We found that soluble RGMb-mlgG2a bound to neogenin-transfected 300 cells (Fig. 2 f). RGMb mAbs weakly blocked RGMb-mlgG2a binding to neogenin (unpublished data). Surprisingly, cell conjugation assays showed that 300-mRGMb cells did not bind to 300-neogenin cells (Fig. 2 g), suggesting that the structural orientation of RGMb and neogenin binding is not compatible with cell to cell binding but can support binding in cis on the same cell surface or of soluble RGMb to cell surface neogenin.

**Model of RGMb interactions**

Based on previous studies and our findings, we propose a model for RGMb–PD-L2 signaling (Fig. 2 h, left). RGMb directly binds to BMP-2 or BMP-4, which bind to type I BMP receptors (BMPR1a, BMPR1b, ACVR1, and ACVRL1) and recruit type II BMP receptors (BMPR2, ACVR2a, and ACVR2b; Corradini et al., 2009; Yoshioka et al., 2012). Then type II BMP receptors phosphorylate type I BMP receptors, which phosphorylate Smad1/5/8 or p38 mitogen-activated protein kinase (MAPK) and extracellular signal-regulated protein kinase (ERK), leading to downstream target gene transcription (Corradini et al., 2009; Xia et al., 2011). RGMs facilitate the utilization of ACVR2a by BMP-2/4. In the absence of an RGM, BMP-2/4 preferentially use BMPR2 (Corradini et al., 2009). RGMb may also signal through neogenin and downstream effector Rho, triggering cytoskeletal rearrangement (Conrad et al., 2007; Bell et al., 2013). We propose that RGMb can form a signaling supercomplex of BMP–BMP receptors–RGMb–neogenin (BBRN supercomplex). We would caution that this model pictures all the possible players and not all may be complexed simultaneously. PD-L2 may interact with this BBRN supercomplex by binding to RGMb, and modulate these signaling pathways. Fig. 2 h (right) shows PD-L2 binding to PD-1, which results in tyrosine phosphorylation of the PD-1 cytoplasmic domain, recruitment of tyrosine phosphatases, particularly SHP-2, and attenuation of antigen receptor signals. Thus, PD-L2 may participate in three important signaling circuits, the PD-1, BMP, and neogenin signaling pathways, by binding to either PD-1 or RGMb.

**RGMb expression in mouse macrophages**

RGMb mRNA has been reported in mouse lung macrophages, macrophage cell line RAW264.7 and myoblast cell line C2C12 (Xia et al., 2011), but protein expression was not determined. We confirmed RGMb mRNA expression (not depicted) and showed cell surface expression of RGMb protein on these cell lines by flow cytometry using RGMb mAb 9D1 (Fig. 3 a).

We also analyzed RGMb protein expression using RGMb mAb 1H6, which Western blots both mouse and human RGMb. One major band (~37 kD) and one minor band (~49 kD) were detected in RAW264.7, J774.1, and C2C12 cells as well as mRGMb-transfected 300 cells (Fig. 3 b). No such bands were observed in 300 cells (Fig. 3 b), consistent with the absence of RGMb mRNA (unpublished data). Western blotting demonstrated RGMb expression in mouse peritoneal macrophages and thioglycollate-induced peritoneal macrophages (Fig. 3 c).

Fig. 3 d shows how proteolytic cleavage of RGMb accounts for these multiple protein bands. RGMb contains a portion of a von Willebrand Factor type D domain with a proteolytic cleavage site between Asp171 and Pro172. After cleavage, the two fragments of RGMb remain connected by disulfide bonds (Severyn et al., 2009). The molecular weight of uncleaved RGMb is predicted to be 40 kD, and cleaved RGMb will have N- and C-terminal fragments of 13 and 27 kD, respectively. Each fragment has one predicted N-linked glycosylation site, which should increase the molecular weight of each fragment by 5–10 kD. The 1H6 mAb reacts with an epitope in the C-terminal fragment and recognizes the 37-kD cleaved form as well as the 49-kD uncleaved form. Our Western blotting analysis shows that most native RGMb is the cleaved form.

**RGMb can be expressed on the cell surface, but is primarily localized intracellularly**

Previous studies have reported RGMb in breast and prostate tumor cells (Li et al., 2012a,b). We detected RGMb protein on the cell surface of human breast cancer cell lines, MDA-231, SKBR-3, and MCF-7, nonsmall cell lung cancer cell line NCI-H226, and renal cancer cell line SN12C (Fig. 4 a), and this expression was confirmed by Western blotting (Fig. 4 b).
Because a previous immunohistochemical study showed RGMb primarily in the cytoplasm of prostate tumor cells (Li et al., 2012a), we also evaluated the cellular localization of RGMb in SKBR-3 cells. Confocal microscopy showed substantial amounts of RGMb inside SKBR-3 cells (Fig. 4c), consistent with the published study, but also revealed detectable RGMb on the cell surface (Fig. 4a).

Western blotting showed positive RGMb expression in cells from spleen, thymus, purified splenic CD4⁺, and CD8⁺ T cells from naive mice (Fig. 4d). Cell surface RGMb expression was not detectable in primary hematopoietic cells by flow cytometry with PE-conjugated RGMb mAb 9D1 (unpublished data). RGMb mRNA and protein levels were not up-regulated in T cells by CD3 and/or CD28 activation (unpublished data), suggesting that RGMb is not a T cell activation molecule. Intracellular flow cytometry staining using PE-conjugated RGMb mAb 9D1 did not detect RGMb expression in splenic T cells (CD3⁺), B cells (CD19⁺), or DCs (CD11c⁺) from naive mice (Fig. 4e, top), but detected low levels of RGMb expression in these cells from mice treated with FMS-like tyrosine kinase 3 ligand (FLT-3L) to expand DC populations (Fig. 4e, bottom). Our findings of intracellular RGMb expression are in agreement with the intracellular expression seen in confocal microscopy of SKBR-3 cells (Fig. 4c) and immunohistochemical staining of cancer cells (Li et al., 2012a). The intracellular localization of RGMb along with the variability in RGMb cell surface expression suggests additional levels of regulation of RGMb cell surface expression.

qRT-PCR showed positive RGMb expression in cells from lung, spleen, thymus, peripheral, and mesenteric LNs with much higher levels of RGMb mRNA in lung cells (Fig. 4f). Western blotting demonstrated RGMb expression in mouse lung alveolar macrophages (AMs) and interstitial macrophages (IMs; Fig. 4g).

Lung IMs, alveolar epithelial cells (AECs), and DCs may be involved in RGMb–PD-L2 interaction in the lung PD-L2 blockade has particularly strong effects on immune responses in the lung (Akkari et al., 2010; Singh et al., 2011), and RGMb is highly expressed in the lung (Fig. 4f). Notably, lung inflammation (pneumonitis) is the most severe adverse event reported in human clinical trials of PD-1 mAb (Topalian et al., 2012). Therefore, we investigated the roles of PD-L2 and RGMb, and of RGMb–PD-L2 interactions in a mouse model of OVA-induced respiratory tolerance. Lung IMs, but not AMs, have been reported to produce high levels of IL-10 and inhibit LPS-induced maturation and migration of DCs, thereby preventing airway allergy in mice (Bedoret et al., 2009). In addition, epithelial cells have been shown to interact with DCs to maintain respiratory tolerance through production of a diverse array of mediators that modulate the activity of DCs (Strickland et al., 2010).

To identify the lung cell subsets involved in RGMb–PD-L2 interaction that regulate respiratory tolerance, we examined mRNA expression of RGMb, PD-L2, and related proteins in unstimulated lung IMs (F4/80⁺CD11c⁻), AMs (F4/80⁺CD11c⁻), DCs (F4/80⁺CD11c⁻), other cells (F4/80⁻CD11c⁻), as well as CD4⁺ T cells (TCRβ⁺CD4⁺), CD8⁺ T cells (TCRβ⁺CD8⁺), AECs, and tracheobronchial epithelial cells (TECs; Fig. 5, a–i). IMs and AECs expressed the highest levels of RGMb, BMP-2/4, BMP type I receptors (Bmpr1a, Bmpr1b, and Actvr1), BMP type II receptors (Bmpr2, Actvr2a, and Actvr2b), neogenin and its ligand netrin 1 (Fig. 5, a–e and i). IMs also expressed the highest levels of IL-10 (Fig. 5f). PD-L2 expression on IMs and AECs was barely detectable (Fig. 5g). DCs expressed...
with antigen (Fig. 6 a). We first compared the development of respiratory tolerance in WT and PD-L2–deficient mice. T cells from WT mice exposed to intranasal OVA were tolerized, as indicated by significantly reduced T cell proliferation and IL-4 responses compared with control mice that did not receive i.n. OVA (Fig. 6 b). Strikingly, PD-L2–deficient mice showed resistance to the development of respiratory tolerance. T cells from PD-L2–deficient mice that received OVA i.n. displayed similar levels of proliferation and IL-4 production as T cells from control PD-L2–deficient mice that received PBS i.n. Similarly, treatment of WT mice with a PD-L2 mAb (3.2) that blocks both PD-L2–RGMb and PD-L2–PD-1 interaction prevented the development of respiratory tolerance, resulting in increased proliferation and IL-4 production compared with tolerized mice treated with control mAb (Fig. 6 c). These results suggest that PD-L2 is critical for the development of respiratory tolerance.

Because PD-L2 can interact with both PD-1 and RGMb, we used specific mAbs to evaluate the contribution of RGMb to immune responses. RGMb is an essential receptor for PD-L2, 9D1, and B7-2, but low levels of most other molecules (Fig. 5, a–h). IMs had very different expression profiles than AMs (Fig. 5, a–h). Lung CD4+ and CD8+ T cells expressed very low levels of all the molecules examined except for PD-1 and PD-L1 (Fig. 5, a–h). These expression data support our proposed BBRN model with RGMb associating in a supercomplex with BMP receptors and/or neoegenin on IMs and/or AECs, then interacting with PD-L2 on DCs (Fig. 2 h).
Article

Airway hyperreactivity (AHR), a cardinal feature of asthma. Previous work has shown that tolerization via i.n. administration of OVA can protect from OVA-induced AHR. We exposed mice to OVA i.n. or PBS control, and then immunized with OVA in ALUM i.p., challenged with OVA i.n. on days 20–22, and assessed for the development of AHR on day 23 (Fig. 6 e). As expected, mice that received PBS i.n. (nontolerized) developed high levels of AHR measured as airway resistance (R_L) and dynamic compliance (C_{dyn}) in anesthetized, tracheostomized, and mechanically ventilated mice. Administration of OVA i.n. induced respiratory tolerance, such that the tolerized mice were protected and developed only mild levels of AHR. However, treatment of the mice with RGMb-1a mAb 9D1 (blocks RGMb–PD-L2 and RGMb–BMP) or with PD-L2 mAb 2C9 (blocks RGMb–PD-L2 but not PD-1–PD-L2) or with isotype control mAb. Administration of either 2C9 or 9D1 mAb inhibited the development of respiratory tolerance and led to higher levels of T cell proliferation and IL-4 production compared with control mAb-treated mice (Fig. 6 d). These results support a role for the RGMb–PD-L2 interaction in promoting the development of respiratory tolerance.

To further evaluate the effects of the RGMb–PD-L2 interaction in tolerance, we rechallenged mice that were exposed to antigen i.n. during RGMb or PD-L2 blockade, and evaluated airway hyperreactivity (AHR), a cardinal feature of asthma. Previous work has shown that tolerization via i.n. administration of OVA can protect from OVA-induced AHR. We exposed mice to OVA i.n. or PBS control, and then immunized with OVA in ALUM i.p., challenged with OVA i.n. on days 20–22, and assessed for the development of AHR on day 23 (Fig. 6 e). As expected, mice that received PBS i.n. (nontolerized) developed high levels of AHR, measured as airway resistance (R_L) and dynamic compliance (C_{dyn}) in anesthetized, tracheostomized, and mechanically ventilated mice. Administration of OVA i.n. induced respiratory tolerance, such that the tolerized mice were protected and developed only mild levels of AHR. However, treatment of the mice with RGMb

Figure 5. Expression of RGMb, PD-L2, BMPs, BMPRs, and related molecules in resting lung cell populations and airway epithelial cells. Expression of indicated mRNAs in resting lung cell populations and airway epithelial cells by qRT-PCR. (a–h) IMs (F4/80^+ CD11c^-), AMs (F4/80^+ CD11c^+), DCs (F4/80^- CD11c^-), and other cells (F4/80^- CD11c^-) were sorted from lung cells pooled from five to eight mice; CD4^+ T cells (TCR^β^+ CD4^+), CD8^+ T cells (TCR^β^- CD8^-) were sorted from lung cells pooled from two mice. (i) AECs and TECs were pooled from three and eight mice, respectively. Data are mean ± SEM; n = 3; *, P < 0.05; **, P < 0.01; ***, P < 0.005; ****, P < 0.001. Ordinary one-way ANOVA followed by Dunnett’s multiple comparison test. All data are representative of two or more experiments.
Figure 6. PD-L2 or RGMb blockade inhibits induction of respiratory tolerance. (a) Experimental protocol for induction of respiratory tolerance. (b–d) PD-L2–deficient or WT mice in b, and WT mice injected with the indicated mAb in c and d, were exposed to OVA i.n. (tolerized) or PBS (nontolerized) and subsequently received 50 µg OVA/ALUM i.p. T cell proliferation (top) and IL-4 production (bottom) in response to restimulation with OVA in vitro are shown. n = 2–3. Data are representative of two to five experiments. Data are mean ± SEM, *, P < 0.05; **, P < 0.01; ***, P < 0.005. (b, top) two-way ANOVA followed by Tukey’s multiple comparisons test. (c, top, and d) Two-way ANOVA followed by Dunnett’s multiple comparison test. (b and c, bottom) Two-tailed Student’s t test. (e) Experimental protocol used in f and g. (f) WT mice injected with the indicated mAb were exposed to OVA i.n. or PBS, immunized with 50 µg OVA/ALUM i.p. Mice were subsequently challenged with OVA i.n. on three consecutive days and assessed for AHR by measuring lung resistance (R_L) and dynamic compliance (Cdyn). Data are the mean ± SEM of 4 mice/group. (g) Lung histopathology of mice from panel f. Lung tissue was stained with H&E and analyzed for cell infiltration. Data in f and g are representative of two experiments.
mAb 9D1 or with PD-L2 mAb 3.2 abolished the induction of tolerance, and upon rechallenge with OVA, these mice developed severe AHR which was comparable to that of non-tolerized mice (Fig. 6 f). The lungs of tolerized mice showed low levels of cellular infiltration surrounding the airways in contrast to nontolerized mice, which showed substantial airway inflammation. Mice treated with PD-L2 mAb or RGMb mAb demonstrated even more extensive cellular infiltration than nontolerized mice (Fig. 6 g). Thus, respiratory tolerance promoted by RGMb–PD-L2 interaction protects against the development of airway hyperreactivity and lung inflammation after antigenic challenge.

To determine if RGMb blockade would impact systemic immunization, mice treated with RGMb mAb were immunized in the footpad with OVA in IFA, and T cell responses in the draining lymph nodes were evaluated 9 d later. There were no differences in proliferation, IL-4 or IFN-γ production between mice treated with RGMb mAb 9D1 or control mAb (unpublished data). These results demonstrate that treatment with RGMb mAb does not have a global impact on in vivo immune responses.

**Tolerized mice have fewer OVA-responsive T cells than control mice or RGMb mAb-treated mice**

To further understand the mechanisms underlying this model of respiratory tolerance induction and how blockade of RGMb–PD-L2 interaction abrogates tolerance in this model, we compared the number and phenotype of T cells responding to challenge with OVA after tolerance induction in the presence or absence of RGMb mAb. Splenocytes from RGMb mAb-treated and control mice were labeled with CFSE before culture with OVA. Tolerized mice had a lower percentage of proliferating OVA-specific CD4+ T cells (CFSE low cells) than nontolerized mice (Fig. 7, a and b). Treatment with RGMb mAb prevented tolerance induction, as shown by a restoration of the percentage of CD4+ T cells proliferating to OVA to levels observed in nontolerized mice. Cells from tolerized mice produced less IL-4, IL-5, and IFN-γ than nontolerized mice, whereas cells from tolerized, RGMb mAb-treated mice had the same high levels of cytokine as nontolerized mice (Fig. 7 c). Cytokine production per cell was calculated by dividing total IL-4 or IFN-γ production by the number of responding (CFSE low) T cells. No significant differences were found in cytokine per cell between nontolerized, tolerized, and tolerized plus RGMb mAb–treated groups at 250 µg/ml OVA in culture (Fig. 7 d). Thus, tolerization primarily affects the quantity of responsive T cells. These data suggest that the primary mechanism of tolerance in this model is deletion or anergy of OVA-specific cells and that treatment with RGMb mAb during tolerance induction impairs this process.

**Lung macrophages and DCs are not depleted by treatment with PD-L2 or RGMb mAbs**

To determine if PD-L2 mAb 2C9 inhibited tolerance induction by depleting PD-L2–expressing cells, mice primed with OVA in ALUM were treated with mAb 2C9 on the day of the second administration of OVA i.n. Previous work had shown that OVA–primed mice express high levels of PD-L2 on lung DCs and macrophages. The number and intensity of PD-L2–positive cells in the lung were examined 48 h later using a PD-L2 mAb (TY25) that recognizes a different epitope than mAb 2C9. Under these inflammatory conditions, PD-L2 was expressed by alveolar macrophages (F4/80CD11c+) and about half of dendritic cells (F4/80CD11c+) and interstitial macrophages (F4/80CD11c+b). The PD-L2 intensity and number of cells was not different in mice treated with mAb 2C9 (Fig. 8, b and c), indicating that treatment with mAb 2C9 did not deplete PD-L2–expressing cells. Similarly, treatment of mice with RGMb mAb 9D1 did not deplete these populations (not depicted).

**Blockade of RGMb–PD-L2 interaction impairs T cell expansion to antigen**

Previous studies of respiratory tolerance showed that intranasal administration of antigens such as house dust mite (Hoynes et al., 1996) or OVA (Tsitoura et al., 1999; Albacker et al., 2012) resulted in a strong transient CD4+ T cell response, followed by deletion of most of the antigen–specific T cells, with a small population of unresponsive T cells remaining. Inhaled antigen is sampled by immature DCs in the lung, which migrate to the draining mediastinal LNs, where they encounter antigen-specific T cells. In the absence of inflammatory stimuli, DCs induce transient antigen–specific T cell activation followed by T cell deletion and unresponsiveness (Hawiger et al., 2001). Respiratory tolerance involves multiple mechanisms including deletion of antigen–specific T cells, the development of anergy and regulatory T (T reg) cells and these processes may occur concurrently (Holt and Upham, 2004; GeurtsvanKessel and Lambrecht, 2008; Bedoret et al., 2009; Albacker et al., 2012). To explore the mechanism whereby RGMb and PD-L2 interaction enhances respiratory tolerance, we examined the effect of blocking RGMb–PD-L2 interaction on the activation and expansion of transferred OVA-specific DO11.10 CD4+ T cells. Recipient mice were treated with RGMb mAb 9D1 or control mAb, subsequently given OVA or PBS i.n. on days 0, 1, and 2, and the fate of DO11.10 T cells was monitored using the clonotypic mAb KJ1-26 (Fig. 9 a). As expected, exposure of the mice to i.n. OVA resulted in marked expansion of the DO11.10 cells in the mediastinal LN on day 7 compared with mice that received i.n. PBS. Notably, this expansion of KJ1–26+ T cells was greatly diminished in tolerized mice treated with anti-RGMb compared with the control mAb treated mice (Fig. 9 b, left). The number of OVA-specific T reg cells (KJ1–26+ Foxp3+) was similarly reduced in RGMb mAb–treated mice, indicating that the reduced expansion of KJ1–26+ T cells was not due to increased numbers of OVA-specific T reg cells (Fig. 9 b, right).

We next examined the effect of RGMb mAb on the expansion of KJ1–26+ T cells at earlier time points. Similar numbers of KJ1–26+ cells were detected in LNs of RGMb mAb and control treated mice on day 3. However, by day 5 the number of KJ1–26+ cells increased in the control mice, but showed
little expansion in the RGMb mAb–treated group (Fig. 9 c).
These findings suggest that blockade of RGMb inhibits the 
induction of tolerance by impairing the expansion of T cells 
that normally occurs after respiratory administration of OVA.

To determine if treatment with PD-L2 mAb would similarly 
inhibit the response of OVA–specific CD4+ T cells after 
activation to OVA i.n., mice were treated with PD-L2 mAb, 
RGMb mAb, or control mAb and the number of KJ1–26+ cells 
examined on day 5. Reduced numbers of KJ1–26+ cells were 
detected in both PD-L2 mAb and RGMb mAb–treated 
groups compared with control mAb–treated mice, indicating 
that expansion of KJ1–26+ cells was inhibited (Fig. 9 d). To 
determine if anti–PD-L2 mAb was altering the time course 
of the T cell response, we examined the in vivo response of 
DO11.10 T cells to i.n. OVA over a 9-d period in mice treated 
with PD-L2 mAb or control mAb (Fig. 9 e). In control mAb–
treated mice, the number of KJ1–26+ cells in mediastinal LNs 
increased substantially after day 3, peaking at day 5 and de-
clining thereafter with a small but detectable population 
remaining on day 9, as previously described (Tsitoura et al., 1999).

In mice treated with PD-L2 mAb, a limited increase in the 
number of KJ1–26+ cells was observed between days 3 and 5 
that was significantly lower than the increase observed in control 
mAb–treated mice. Numbers subsequently declined, indicat-
ing that blocking PD-L2 inhibited the expansion of OVA–specific 
T cells throughout the response.

We then determined if expression of PD-L2 on T cells, non–T 
cells, or both was required for T cell expansion. We compared 
expansion of WT KJ1–26+ T cells after transfer to PD-L2–
deficient or WT mice and exposure to OVA i.n. Between days 3 
and 5, the WT KJ1–26+ cells underwent a 72-fold expansion in 
WT recipients but only a 9.7-fold expansion in PD-L2–/– 
recipients (Fig. 9 f), indicating a requirement for PD-L2 ex-
pression on non–T cells. Consistent with this, DO11.10 T cells 
from PD-L2–/– and WT mice expanded similarly when trans-
ferred to WT mice (Fig. 9 g). These experiments demonstrate 
the importance of PD-L2 expression on non–T cells for the 
initial expansion of T cells to OVA i.n.

To determine the involvement of the PD-1–PD-L2 versus 
RGMb–PD-L2 pathways in this process, mice were treated 
with PD-L2 mAb 2C9, which blocks only the RGMb–PD-L2 in-
teraction or with control mAb. The expansion of KJ1–26+ cells 
on day 5 was significantly reduced in mice treated with mAb 
2C9 compared with control–treated mice (Fig. 9 h), suggest-
ing that PD-1 is not required for this effect because 2C9 does 
not block the interaction of PD-L2 with PD-1. Together, these 
data indicate that blocking RGMb–PD-L2 interaction prevents 
the induction of respiratory tolerance, and does so by reduc-
ing the initial expansion of CD4+ T cells in response to OVA 
by a pathway that does not involve PD-1.

We then examined the effects of blocking PD-L1 or PD-1 
compared with PD-L2 on the initial activation and expand-
ion of transferred DO11.10 OVA–specific CD4+ T cells. On 
day 5, mice treated with the blocking PD-L1 mAb 9G2 or 
PD-1 mAb IA2 had substantially higher numbers of KJ1–26+ 
cells in the mediastinal LNs than control mAb–treated mice.

Figure 7. Tolerized mice have fewer OVA– 
responsivce T cells than control mice or RGMb 
mAb–treated mice. B cell–depleted splenocytes 
prepared on day 19 from mice treated as in Fig. 6 a 
were labeled with CFSE and cultured with OVA for 
4 d. Cells were stained for CD3 and CD4 and sub-
gated on CD3+CD4+ cells for analysis of CFSE 
dilution. (a) Flow cytometry plots show data from 
one representative mouse and (b) shows mean ± 
SEM of three mice/group. (c) On day 4, cultures 
were harvested as in panel a and supernatants 
were collected and assayed for cytokine produc-
tion. Total IL-4, IL-5, and IFN-γ in culture super-
natants is shown. (d) Cytokine per cell was 
calculated as total cytokine amount in superna-
tant divided by the number of CFSE low cells. 
Data are mean ± SEM; **, P < 0.01; ***, P < 0.005; 
n = 3, Student’s t test. Data are representative of 
two experiments.

952
(a–c) Mice immunized with OVA

There is now substantial agreement that the PD-L2 and RGMb interaction reduces the initial T cell expansion involved in tolerance induction. BMP signaling could be involved in the induction of respiratory tolerance, as the RGMb mAb 9D1 that blocks both the RGMb–PD-L2 and RGMb–BMP-2/4 interactions impaired respiratory tolerance. Our data on RGMb and PD-L2 pathway expression in lung cell populations, together with previous findings (Bedoret et al., 2009; Strickland et al., 2010) suggest that IMs, AECs, and DCs are involved in the induction of respiratory tolerance. We propose that RGMb on IMs and/or AECs forms a BBRN supercomplex with BMP-2/4, BMP receptors, and neogenin and interacts with PD-L2 on DCs. This may signal directly or induce the production of mediators such as IL-10 in IMs or AECs to inhibit the maturation and migration of DCs, thereby promoting respiratory tolerance. In support of this, our data show that PD-L2 expression on non-T cells provides the required signal for the induction of respiratory tolerance because WT T cells transferred into PD-L2–deficient mice do not undergo the initial T cell expansion involved in tolerance induction. BMP signaling could be involved in the induction of respiratory tolerance, as the RGMb mAb 9D1 that blocks both RGMb–PD-L2 and RGMb–BMP-2/4 interactions impaired respiratory tolerance.

Components of the BBRN complex have been shown to have immune function. Previous studies have shown that BMP signaling has immunoregulatory effects on T cells (Graf et al., 2002; Hager-Theodorides et al., 2002; Lu et al., 2010; Yoshioka et al., 2012). BMP-7 was recently shown to be an instructive factor for Langerhans cell differentiation (Yasmin et al., 2013). Elevated BMP/Smad signaling has been found during airway inflammation (Rosenfeld et al., 2002). RGMb-mediated BMP signaling was reported to inhibit IL-6 production by lung macrophages (Xia et al., 2011). Neogenin has been shown to be involved in pulmonary inflammation and acute inflammatory peritonitis (König et al., 2012; Mirakaj et al., 2012), so neogenin signaling could also be involved in the induction of respiratory tolerance. RGMb interaction with neogenin has been shown to control aggregation and migration of neogenin-positive neuroepithelial cells (Conrad et al., 2010). Another member of the RGM family, RGMa, was also highly expressed in IMs and AECs (Fig. 5, a and i), and was shown to be involved in immune responses (Muramatsu et al., 2011).
molecular details of the cross-talk between the components of the BBRN signaling supercomplex remain to be determined. Though early work emphasized the restricted expression of PD-L2 on DCs and macrophages, recent work has identified PD-L2 expression on human primary nasal epithelial cells (Kim et al., 2005), Th2 cells (Lesterhuis et al., 2011), activated T cells (Messal et al., 2011), and some B cell subsets (Zhong and Rothstein, 2011). In the BBRN model, PD-L2 on any of these cells might interact with RGMb on various cells and signal through BMP and/or neogenin pathways.

Antibodies blocking the PD-1 pathway have shown promising results in cancer immunotherapy clinical trials. The most severe adverse event with PD-1 mAb treatment is pneumonitis (3%) leading to 3 deaths (1%; Topalian et al., 2012). The role of the different PD-1 ligands in pneumonitis is not yet understood. PD-1 blockade may shift the balance in PD-L2 interaction with its binding partners, increasing PD-L2 availability for binding to RGMb, leading to pneumonitis. Additional work will need to be done to distinguish which therapeutic approaches are beneficial.

Because our mRGMb mAb does not bind to mRGMa, the effects of our RGMb mAb on respiratory tolerance are independent of RGMa.

Our model suggests that PD-L2 binding to RGMb could signal through BMP and/or neogenin signaling pathways. This hypothesis may offer insight into the signaling conundrum of RGMb and PD-L2, i.e., RGMb and PD-L2 do not appear to be able to signal directly because RGMb is a glycosphingolipid-anchored protein and PD-L2 has only a short cytoplasmic tail without obvious signaling motifs. To signal, RGMb–PD-L2 may need to form a complex with BMPs, BMP receptors, and/or neogenin. BMP and neogenin signaling may act together because neogenin can directly bind BMP-2/4/6/7 and modulate BMP signaling (Hagihara et al., 2011; Tian and Liu, 2013). Nevertheless, these two signaling pathways may also function separately because a recent study showed that RGMb induces apoptosis in mouse renal tubular epithelial cells through neogenin and not the BMP pathway (Liu et al., 2013). RGMb is the pivot joining all these proteins, but the molecular details of the cross-talk between the components of the BBRN signaling supercomplex remain to be determined.

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Figure 9. Blockade of PD-L2–RGMb interaction reduces the initial expansion of antigen-specific T cells after intranasal OVA administration. (a) Experimental protocol for Fig. 9. Mice received D011.10 T cells on day 0 and OVA i.n. The number of KJ1-26 CD4+ cells in mediastinal LNs was determined on the indicated days. (b) Mice treated with RGMb mAb 9D1 or control mAb on day −1 received 5 × 10^6 D011.10 T cells. Left, TCRβ+ KJ1-26 CD4+. Right, FoxP3+ TCRβ+ KJ1-26 CD4+ cells. For each group, mediastinal LNs of 3–5 mice were combined; data are representative of two experiments. (c–e) Mice received 2 × 10^6 D011.10 T cells after (c) RGMb mAb 9D1 or control mAb; (d) RGMb mAb 9D1, PD-L2 mAb 3.2, or control mAb; (e) PD-L2 mAb 3.2 or control mAb. (f) WT D011.10 T cells were transferred into WT or PD-L2−/− mice; (g) WT or PD-L2−/− D011.10 T cells were transferred into WT mice. (h–j) Mice received 2 × 10^6 D011.10 T cells after (h) PD-L2 mAb 2C9 or control mAb; (i) PD-L1 mAb 9G2 or control mAb; (j) PD-L2 mAb 3.2, PD-1 mAb 1A12, or control mAb. The number of KJ1-26 CD4+ T cells in mediastinal lymph nodes on day 5 (h–j) was determined by flow cytometry. (b–d) Each point represents the mean and SEM of KJ1-26 CD4+ T cells in LNs of two or three groups of two mice each. Data are representative of two to four experiments. **, P < 0.01; ***, P < 0.005, two-tailed Student’s t test.
modality has the optimal balance of efficacy and minimal adverse events. Treatment with PD-L2-Ig decreased murine TC-1 lung tumor burden and increased survival, possibly by depletion of T cells expressing high levels of PD-1 (Mkrtichyan et al., 2012); however, these PD-L2-Ig effects could also be through RGMb and the BBRN complex. Further studies are needed to determine if RGMb–PD-L2 interaction serves as a target for immunotherapy.

In summary, we discovered that RGMb is a binding partner for PD-L2. RGMb may coordinate with multiple receptors in the BMP and neogenin pathways to form a BBRN signaling supercomplex. Engagement of PD-L2 with RGMb in this supercomplex may impact BMP and neogenin signaling. We detected high miRNA levels of RGMb and components of the BBRN complex in resting lung IMs and AECs. Our data suggest that the novel engagement of RGMb and PD-L2 promotes the development of respiratory tolerance by facilitating the initial T cell expansion in draining LNs. Targeting the PD-L2 pathway has therapeutic potential for asthma, cancer, and other immune disorders, and may help better use the PD-1 pathway in cancer immunotherapy while minimizing adverse events.

MATERIALS AND METHODS

Mice. WT BALB/c or BALB/cByJ mice were purchased from The Jackson Laboratory. PD-L2+/− mice on BALB/c background have been previously described (Keir et al., 2008). OVA TCR-transgenic DO11.10 mice and PD-L2+/− mice crossed to OVA TCR-transgenic DO11.10 mice were used as donors of OVA-specific CD4+ T cells (Tsitoura et al., 1999). Age-matched female mice were used at 6−12 wk. Animal protocols were approved by The Animal Care and Use Committees at Boston Children’s Hospital, the Dana-Farber Cancer Institute, and Harvard Medical School.

Cells and culture media. The mouse 300.19 pre-B cell line was transfected by electroporation with mRGMb, hRGMb, or mNeogenin cDNA in the pE-Puro expression vector. Cells were selected in media containing puromycin, sorted, and subcloned. Cell surface expression of mRGMb, hRGMb, or mNeogenin was verified by flow cytometry using an mRGMb polyclonal antibody (R&D Systems), an hRGMb mAb (R&D Systems), or mRGMb-Ig, respectively. Other transfected cells, such as 300-mPD-L2, 300-mPD-L1, 300-mPD-1, and Jurkat-hPD-1, have been made previously using similar methods. Cells were cultured in RPMI-1640 (Mediatech) supplemented with 10% heat-inactivated FBS (Invitrogen), 1% streptomycin/penicillin, 15 µg/ml gentamicin (Invitrogen), 1% glutamax (Invitrogen), and 50 mM β-mercaptoethanol (Sigma-Aldrich) at 37°C with 5% CO2.

The cell lines used in this study were purchased from American Type Culture Collection, except for SN12C which was obtained from the National Cancer Institute. Mouse spleen or LN cells for tolerance experiments were cultured in high-glucose DMEM (Invitrogen) supplemented with 10% heat-inactivated FBS (Invitrogen), 1% streptomycin/penicillin, 15 µg/ml gentamicin (Invitrogen), and 1% Hepes (Invitrogen) in a round-bottom 96-well plate for 45 min at 37°C. Conjugate formation was analyzed immediately by flow cytometry using the PE channel for the red dye and the FITC channel for the green dye. The automated plate harvester of the FACS Canto (BD) was used for uniformity of cell processing. The instrument settings were as follows: throughput mode, standard; loader settings, sample flow rate at 0.5 µl/second; sample volume and mixing volume at 100 µl; mixing speed at 50 µl/second; number of mixes at 2; and wash volume at 400 µl. When the cell conjugation assay was used to test the blocking capacity of antibodies, the relevant transfected cell was preincubated 30 min at room temperature with antibodies before adding candidate binding cells.

Flow cytometry. Cells were stained with target antibodies and isotype controls using standard flow cytometry procedures, and analyzed on a FACS Canto (BD) using FlowJo 9.2 software (Tree Star).

To initially verify RGMb expression on mRGMb- or hRGMb-transfected cells, sheep anti-mRGMb (R&D Systems) or sheep IgG (SouthernBiotech) plus donkey anti-sheep IgG-PE (Jackson ImmunoResearch Laboratories) and mouse anti-hRGMb (mAb; R&D Systems) plus goat anti-mouse IgG-PE (SouthernBiotech) were used, respectively, all at 10 µg/ml.

To test the binding specificities of mRGMb antibodies, mRGMb- or hRGMb-transfected 300 cells were incubated with serial dilutions of sera, culture supernatants, or purified antibodies, and binding was detected with 5 µg/ml of goat anti-rat IgG-PE (SouthernBiotech). For biotin-conjugated mRGMb mAb 9D1, 1.4 µg/ml of streptavidin-PE was used. For receptor-ligand binding assay, mRGMb-, mPD-L2-, or mNeogenin-transfected 300 cells, 300 cells, and hPD-1–transfected Jurkat T cells were stained with serial dilutions of specific protein–Ig or control-Ig plus 5 µg/ml of Fab2 goat anti-hlgG-PE (mouse-absorbed; SouthernBiotech), or 10 µg/ml of goat anti-mlgG2a-PE (SouthernBiotech), or 5 µg/ml of Fab2 goat anti-mlgG2a-PE (SouthernBiotech).

To test the capacities of RGMb mAbs to block mRGMb binding to mNeogenin, mRGMb-mlgG2a at 5 µg/ml was incubated with serial dilutions...
of RGMb mAbs, and then added to neogenin-transfected 300 cells. Binding was detected using 10 µg/ml of goat anti-mlgG2a-PE (SouthernBiotech). To determine the background binding, 300-mNeogenin cells were stained with serial dilutions of mlgG2a isotype control.

To assess cell surface expression of RGMb and PD-L2 on mouse cell lines RAW264.7, J774.1, and C2C12, cells were preincubated with mouse Fc receptor mAb (2.4-G2), and then stained with PE-conjugated RGMb mAb 9D1, mPD-L2-mAb TY25, or rat IgG2a at 5 µg/ml.

To analyze cell surface expression of RGMb on human cancer cell lines, RGMb mAb 1H6 or rat IgG2a at 10 µg/ml plus goat anti-rat IgG-PE (SouthernBiotech) at 5 µg/ml were used.

For intracellular flow cytometry analyses of RGMb expression in primary cells from spleen, cells were first stained with LIVEDEAD Fixable Near-IR (Invitrogen) at 1:1,000. After preincubcation with mouse Fc receptor mAb (2.4-G2), cells were stained with CD3-Pacific blue, CD19-PE-cy7 and CD11c-APC (BioLegend). Then cells were permeabilized using BD Cytofix/Cytoperm Fixed/Permeabilization kit, preincubated with mouse Fc receptor mAb (2.4-G2), and stained with PE-conjugated RGMb mAb 9D1 or rat IgG2a at 5 µg/ml. PBS containing 3% BSA and 0.1% Triton X-100 was used for wash and antibody incubation. 300-mPD-L2 cells were used as negative controls.

DO11.10 T cells were identified by staining with TCRβ-APC-eFlour 780, CD4-PerCP, and KJ1–26-APC (eBioscience). DO11.10 T reg cells were identified using CD25-FITC (BioLegend), followed by intracellular staining with Foxp3-PE (BioLegend).

ELISA. To examine specificity of mRGMb mAbs, 96-well plates were coated with 2 µg/ml of recombinant mRGMa-HIS, mRGMb-HIS, or mRGMc-HIS (R&D Systems). Then serial dilutions of mRGMb mAbs and isotype controls were added and incubated for 1 h at 37°C. Mouse anti-rat IgG (y-specific) HRP (Jackson ImmunoResearch Laboratories) at 1:2,500 was used for detection.

To examine RGMa/RGMb/RGMc and PD-L2 interaction, 96-well ELISA plates were coated with 2 or 5 µg/ml of recombinant mRGMa-HIS, mRGMb-HIS, mRGMc-HIS, or hRGMc-HIS (R&D Systems). Then serial dilutions of mPD-L2-hlgG1/IgA, mPD-L2-hlgG2a/IgA, hpD-L2/mlgG2a, or control-Ig fusion proteins were added and incubated for 1 h at 37°C. Fab3 goat anti-hlgG-HRP (Jackson ImmunoResearch Laboratories) or rat anti–mlgG2a-HRP (BD) at 1:1,000 or 1:10,000 were used for detection.

To test the capacities of mRGMa antibodies and mPD-L2 fusion proteins to block RGMb binding to BMP-2/4, 96-well ELISA plates were coated with 1 µg/ml of recombinant mouse BMP-2 (Invitrogen) or BMP-4 (R&D Systems). mRGMa antibodies, isotype controls, mPD-L2-hlgG1/IgA, mPD-L2-hlgG2a/IgA, or control Ig fusion proteins at the indicated concentrations were preincubated with 20 µg/ml mRGMb-HIS (R&D Systems) for 45 min at 4°C, then added to the plates and incubated for 1 h at 37°C. Anti–penta-HIS-HRP (Qiagen) at 1:1,000 was used for detection.

To determine the capacities of mPD-L2 and BMP-2/4 simultaneously, 96-well ELISA plates were coated with BMP-2/4 as above; mPD-L2-hlgG1/IgA, mPD-L2-hlgG (R&D Systems), mPD-L1-hlgG (R&D Systems), or control-Ig fusion proteins at 10 µg/ml were preincubated with 10 µg/ml mRGMa-HIS (R&D Systems) or buffer alone for 15 min at room temperature, and then added to the plates and incubated for 1 h at 37°C. Alternatively, 10 µg/ml mRGMa-HIS (R&D Systems) or buffer alone was added first to the plates and incubated for 1 h at 37°C. After wash, mPD-L2-Ig or control-Ig fusion proteins were added and incubated for 1 h at 37°C. Fab3 goat anti–hlgG-HRP (Jackson ImmunoResearch Laboratories) at 1:10,000 was used for detection.

The substrate for HRP was TMB using the microwell peroxidase substrate system (KPL).

Biacore. Interactions were determined by analysis of surface plasmon resonance on a Biacore 3000 instrument. Flow cells within a CMS biosensor chip (GE Healthcare) were activated with N-hydroxysuccinimide (NHS) in the presence of 1-ethyl-3-[(3-dimethylaminopropyl)carbodiimide hydrochloride, generating an NHS ester that bound to free amines on RGMb or PD-L2-hlgG passed over the surface (R&D Systems; 25 µg/ml in 100 mM sodium acetate, pH 4.5), followed by binding of free NHS ester with 1 M ethanolamine. A further cell was exposed to immobilizing reagents and blocking reagents in the absence of protein as a control surface. After extensive washing of the surfaces with binding buffer (10 mM Hepes, 150 mM NaCl, pH 7.0, and 0.05% Tween 20), PD-L2-hlgG fusion protein binding was assessed by injecting varying concentrations (0–658 nM) simultaneously over the control and RGMb flow cell surfaces. PD-1-hlgG (R&D Systems; 4.8–308 nM) was similarly passed over immobilized PD-L2 and control surfaces. Between each cycle, the surface was regenerated using 20 mM glycine-HCl, pH 2.8. Values shown in RU have been corrected for nonspecific binding by subtracting the SPR of the control flow chamber exposed to the same injected material, followed by subtraction of buffer alone passing over the RGMb or PD-L2 surface, respectively. Data were analyzed using BIAevaluation 3.2 software and fitted to a 1:1 Langmuir binding model with separate kₐ and kᵦ determinations. The dissociation constant (Kᵦ) was determined as kᵦ/kₐ, and confirmed by a linear transformation of the binding isotherms. For PD-1 binding to PD-L2 surface, the kinetic parameters were derived for nonmonovalent binding of the Fc dimer at higher concentrations (red bar in Fig. 1 e), but data contributing to high-affinity bivalent interactions at lower concentrations were not included in the analysis.

Cell isolation and stimulation. Spleen and thymus cells were isolated by mechanically disrupting the tissues. For FACS analysis, splenocytes were treated with red blood cell lysing buffer (Sigma–Aldrich).

To obtain lung cells or splenocytes to analyze splenic DCs, the lung (perfused with PBS) or spleen was cut into small pieces, digested in RPMI 1640 with 15% FBS, 1 mg/ml collagenase IV (Sigma–Aldrich), and 200 U/ml DNase I (Roche), and then treated with red blood cell lysing buffer (Sigma–Aldrich).

Lung cell populations from naive mice were sorted by flow cytometry. Whole lung cells were isolated as above, and stained with surface markers. Cell populations were defined as follows: IMs (F4/80⁺CD11c⁺), AMs (F4/80⁺CD11c⁻), DCs (F4/80⁺CD11c⁺; Bedoret et al., 2009), other cells (F4/80⁻CD11c⁻), CD4⁺ T cells (TCRβ⁺CD4⁺), CD8⁺ T cells (TCRβ⁺CD8⁺), CD4⁺CD8⁻CD16⁻15⁻ cells from OVA primed and challenged mice were identified as F4/80⁺CD11c⁺low. AECs and TECs were isolated from naive mice as described previously (Lam et al., 2011; Chuquimia et al., 2012). Thioglycollate-induced peritoneal macrophages were obtained from mice on day 4 after i.p. injection with 3% thioglycollate (DIFCO). CD4⁺ and CD8⁺ cells from mouse splenocytes were purified using CD4⁺ T cell isolation kit and CD8⁺ T cell isolation kit (Milenyi Biotec). FLT-3L–stimulated splenocytes were obtained from mice 2 wk after s.c. injection of FLT-3L–transfected RENCA tumor line.

qRT-PCR. Total RNA samples were isolated using the RNeasy mini kit (Qiagen). Reverse transcription was performed using the QuantiTect reverse transcription kit (Qiagen). qPCR using Taqman gene expression assays (Applied Biosystems) was performed in a 7300 Real-Time PCR system (Applied Biosystems). Fold change compared with GAPDH was calculated using the ΔΔt method.

Western blotting. Cell lysates were prepared using RIPA buffer with complete ULTRA protein inhibitors tablets (Roche). Lysates (60–80 µg/lane for cell lines and primary cells, and 0.5 or 1 µg/lane for RGMb-transfected 300 cells) were run on SDS-PAGE under reducing conditions. Western blotting was performed using RGMb mAb 1H6 (10 µg/ml) plus goat anti-rat IgG-HRP (Santa Cruz Biotechnology Inc.; 1:5,000). To blot for loading control, the membranes were treated with Restore Plus Western Blot Stripping Buffer (Thermo Fisher Scientific) and blotted with mouse anti-mouse β-actin (Abcam; 1:5,000) plus goat anti-mouse IgG-HRP (Santa Cruz Biotechnology Inc.; 1:4,000). Protein bands on the membranes were visualized using standard chemiluminescent techniques.

Immunohistochemistry staining and confocal microscopy. Cells were seeded on covergrips in culture medium the day before staining. Cells were fixed with 3.7% formaldehyde for 15 min at room temperature. After
three washes with PBS containing 0.5% BSA, cells were permeabilized in PBS containing 0.5% BSA and 0.5% Triton X-100 for 30 min. Endogenous biotin was blocked with Endogenous Biotin-Blocking kit (Molecular Probes), followed by blocking in PBS containing 3% BSA and 0.1% Triton X-100 for 30 min at room temperature and three washes in PBS containing 0.5% BSA and 0.1% Triton X-100. Cells were then incubated with biotin-conjugated RGMb mAb 1H6 or biotin-conjugated rat IgG2a at 0.5 µg/ml in PBS containing 1% BSA and 0.1% Triton X-100 for 2 h at room temperature, followed by three washes as above. Subsequently, cells were incubated with Alexa Fluor 488–conjugated streptavidin (Jackson ImmunoResearch Laboratories) at 0.5 µg/ml in PBS containing 1% BSA and 0.1% Triton X-100 for 1 h at room temperature, followed by three washes as above. Finally, cells were stained with Phalloidin–TRIC (Sigma–Aldrich) at 50 µg/ml to label F-actin, and coverslips were mounted on slides with mounting medium. Images were taken using a Nikon e–Eclipse Ci confocal microscope equipped with a Melles Griot 488 Ion Laser (with a 515/30 emission filter), a Melles Griot 543 Laser (with a 590/50 emission filter), a Melles Griot 540 Laser, and the Confocal Acquisition Software Nikon EZ–C1 version 3.90.

Respiratory tolerance and mAb treatment. To induce tolerance, lightly anesthetized WT BALB/cByJ or PD-L2−/− mice received 100 µg of LPS–free OVA (Worthington Biochemical) in PBS or PBS alone (control) by intranasal instillation on days 0, 1, and 2. In some experiments, mice were treated with RGMb, PD-L2, PD-L1, or PD-1 blocking mAb or control antibody i.p. on day −1 (450 µg/mouse) and day 2 (200 µg/mouse). On day 12, mice were immunized with 50 µg OVA (ICN Biomedical) adsorbed in 2 mg ALUM. Splenocytes were harvested on day 19 and spleenocytes or B cell–depleted spleenocytes were restimulated in vitro with OVA. Cultures were pulsed with 1 µCi of [3H]–thymidine for the final 17 h and harvested at 96 h. Culture supernatants were harvested at 96 h. IL-4, IFN-γ, and IL-5 were quantitated by ELISA. In some experiments, cells were labeled with CFSE before culture, stained on day 4 with CD3–PerCP/Cy5.5, and CD4–Alexa Fluor 700, and analyzed for CFSE dilution.

Transfer of DO11.10 T cells and respiratory tolerance. T cells from DO11.10 or DO11.10 PD-L2−/− mice were positively selected from spleens by incubating with CD4+ magnetic beads and sorting using MACS columns (Miltenyi Biotec). Each recipient received 2 × 10^5 or 5 × 10^5 DO11.10 T cells i.v., followed by 3 i.n. doses of 100 µg LPS–free OVA on days 0, 1, and 2 and mAbs as indicated. Cells from mediastinal LNs were analyzed by flow cytometry on days 3, 5, and/or 7.

Evaluation of antibodies for in vivo cell depletion. To determine if PD-L2 or RGMb mAbs inhibited tolerance induction by depleting cells, mice immunized on day 0 with OVA in ALUM received OVA i.n. on days 7–9, and were treated with PD-L2 mAb 2C9 or isotype control (500 µg i.p.) on day 8 and dispersed lung cells were examined by FACS on day 10. Expression of PD-L2 was analyzed on AMs (F4/80+CD11c−), IMs (F4/80+CD11c+) and DCs (F4/80−CD11c+), and cell numbers of PD-L2−/− and total cells were enumerated. In a separate experiment, mice were treated with RGMb mAb 9D1 or isotype control and the same analysis was performed.

Statistical analysis. Two-tailed Student’s t test, two-way ANOVA followed by Tukey’s or Dunnett’s multiple comparisons test and ordinary one-way ANOVA followed by Dunnett’s multiple comparisons test were performed using Prism version 6.00 for MacOS X (GraphPad Software). P < 0.05 was considered as significant. Data are mean ± SEM.

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