PD-L1 regulates the development, maintenance, and function of induced regulatory T cells

Loise M. Francisco,1,2 Victor H. Salinas,4 Keturah E. Brown,1 Vijay K. Vanguri,1,2 Gordon J. Freeman,3 Vijay K. Kuchroo,3 and Arlene H. Sharpe1,2

1Department of Pathology, Harvard Medical School, Boston, MA 02115
2Department of Pathology and 3Center for Neurological Diseases, Brigham and Women’s Hospital, Harvard Medical School, Boston, MA 02115
4Harvard University, Cambridge, MA 02138
5Department of Medical Oncology, Dana-Farber Cancer Institute, Boston, MA 02115

Both the programmed death (PD) 1–PD-ligand (PD–L) pathway and regulatory T (T reg) cells are instrumental to the maintenance of peripheral tolerance. We demonstrate that PD–L1 has a pivotal role in regulating induced T reg (iT reg) cell development and sustaining iT reg cell function. PD–L1−/− antigen−presenting cells minimally convert naive CD4 T cells to iT reg cells, showing the essential role of PD–L1 for iT reg cell induction. PD–L1−coated beads induce iT reg cells in vitro, indicating that PD–L1 itself regulates iT reg cell development. Furthermore, PD–L1 enhances and sustains Foxp3 expression and the suppressive function of iT reg cells. The obligatory role for PD–L1 in controlling iT reg cell development and function in vivo is illustrated by a marked reduction in iT reg cell conversion and rapid onset of a fatal inflammatory phenotype in PD–L1−/−PD–L2−/− Rag−/− recipients of naive CD4 T cells. PD–L1 iT reg cell development is mediated through the down−regulation of phospho−Akt, mTOR, S6, and ERK2 and concomitant with the up−regulation of PTEN, all key signaling molecules which are critical for iT reg cell development. Thus, PD–L1 can inhibit T cell responses by promoting both the induction and maintenance of iT reg cells. These studies define a novel mechanism for iT reg cell development and function, as well as a new strategy for controlling T reg cell plasticity.

© 2009 Francisco et al. This article is distributed under the terms of an Attribution–Noncommercial–Share Alike–No Mirror Sites license for the first six months after the publication date (see http://www.jem.org/misc/terms.shtml). After six months it is available under a Creative Commons License (Attribution–Noncommercial–Share Alike 3.0 Unported license, as described at http://creativecommons.org/licenses/by-nc-sa/3.0/).
Figure 1. PD-L1 mediates Foxp3+ iT reg cell development. (A and B) Development of Foxp3+ iT reg cells was assessed by flow cytometric analysis of Foxp3-GFP expression after co-culture of naive CD4+CD62L+Foxp3.GFP− T cells with anti-CD3 and irradiated WT or PD-L1−/− APCs plus the indicated range of TGF-β concentrations for 3 d (A) or PD-L1−Ig or control Ig (human IgG1)-coupled beads (B). One representative experiment of at least three similar experiments is shown. (C) Analysis of Foxp3-GFP expression after culture of naive CD4+CD62L+Foxp3.GFP− T cells with PD-L1−Ig beads and over
immunity and the maintenance of self-tolerance (Keir et al., 2007a, 2008). PD-1 is up-regulated on T cells upon activation, and its ligands have distinct expression patterns, with PD-L1 being expressed much more broadly than PD-L2. PD-L1 is constitutively expressed on mouse APCs (DCs, macrophages, and B cells) and T cells and is further up-regulated upon activation. PD-L1 is also expressed on a wide variety of nonhematopoietic cell types, including vascular endothelial cells, pancreatic islet cells, and at sites of immune privilege including the placenta, testes, and eye (Keir et al., 2008). In contrast, PD-L2 is inducibly expressed primarily on DCs and macrophages.

PD-1–PD-L interactions regulate peripheral CD4 and CD8 T cell tolerance at multiple checkpoints. The PD-1–PD-L1 pathway exerts its effects during the initial phase of activation and expansion of autoreactive T cells by attenuating self-reactive T cell responses during presentation of self-antigen by DCs. For example, loss of PD-1 enhances the responses of naïve self-reactive CD8 T cells upon encounter with DCs bearing self-antigen (Probst et al., 2005; Keir et al., 2007b). In addition, PD-L1 has a novel role in inhibiting self-reactive T cell function. Bone marrow chimera studies have shown that PD-L1 on nonhematopoietic cells mediates tissue tolerance, controlling the intensity of T cell eff cell responses in nonlymphoid organs and shielding tissues from potentially pathogenic effects of self-reactive T cells and immune-mediated tissue damage (Keir et al., 2006).

T reg cells are also essential for the maintenance of peripheral tolerance, and roles for B7-CD28 family members during T reg cell development are emerging. CD28 is a critical regulator of T reg cell homeostasis and function (Tang et al., 2003; Liang et al., 2005). We demonstrate key roles for PD-L1 in promoting T reg cell development and function. Significantly, PD-L1 can promote differentiation and maintain the function of induced T reg cells by sustaining and enhancing Foxp3 expression in iT reg cells. PD-L1 induces iT reg cells by inhibiting the Akt/mTOR signaling cascade, thereby flipping the “molecular switch” in a naïve CD4+ T cell toward T reg cell development. The novel role for PD-L1 in the maintenance, as well as induction of iT reg cells, identifies PD-L1 as an attractive therapeutic target for controlling T reg cell plasticity.

**RESULTS**

**PD-L1 synergizes with TGF-β to promote iT reg cell conversion**

To investigate whether PD-L1 influences the development of iT reg cells, we cultured freshly isolated WT or PD-L1−/− APCs with naïve CD4+CD62L+Foxp3−T cells from Foxp3.GFP reporter mice in the presence of TGF-β and anti-CD3. WT APCs induced naïve T cell conversion into Foxp3+ iT reg cells over a range of TGF-β concentrations (0.5, 1, 2, 4, and 8 ng/ml; 39.8–42.9% CD4+Foxp3+; Fig. 1 A). In contrast, when naïve T cells were cultured with PD-L1−/− APCs, anti-CD3, and the same range of TGF-β concentrations, we observed a profound defect in conversion to Foxp3+ iT reg cells (17.4–22.1% CD4+Foxp3+; Fig. 1 A), suggesting a critical role for PD-L1 in regulating iT reg cell induction.

To further interrogate the role of PD-L1 in iT reg cell development and circumvent potentially confounding factors (such as surface molecules or soluble factors) that might be differentially expressed by WT and PD-L1−/− APCs, we conducted further studies using epoxy beads covalently coupled with anti-CD3, anti-CD28, and either PD-L1-Ig or control Ig as artificial APCs (henceforth referred to as PD-L1-Ig beads or control Ig beads). Co-culture of PD-L1-Ig beads, but not control Ig beads, with naïve CD4+CD62L+Foxp3−T cells significantly enhanced iT reg cell development in the presence of TGF-β (58.9 vs. 32.5%; Fig. 1 B).

Because TGF-β has been shown to be necessary for iT reg cell generation (Chen et al., 2003; Fantini et al., 2004; Marie et al., 2005; Pyzik and Piccirillo, 2007), we questioned whether PD-L1 alone could drive iT reg cell development and override the need for TGF-β. We cultured naïve T cells with PD-L1−Ig beads or control Ig beads in the absence of exogenous TGF-β. PD-L1 beads alone could induce the conversion of naïve T cells to Foxp3+ iT reg cells without TGF-β (2.67 vs. 0.64% for control Ig bead). Moreover, even with increasing amounts of TGF-β, we could not augment iT reg cell development with control Ig beads to the extent observed with PD-L1−Ig beads (Fig. 1 C). Very low amounts of TGF-β (0.033–0.33 ng/ml) were sufficient for significant conversion of naïve CD4+ T cells into iT reg cells by PD-L1−Ig-coated beads (Fig. 1 D).

To determine the contribution of PD-L1 to iT reg cell development, we titered the amount of PD-L1−Ig on the surface of the beads and found a quantitative relationship
between the amount of PD-L1 and the frequency of induced Foxp3+ CD4 T cells. Greater numbers of Foxp3+ CD4 T cells could be induced with increasing amounts of PD-L1, particularly in the absence of exogenous IL-2 (Fig. 1, E and F); however, at the highest concentration, PD-L1 strongly inhibited overall CD4 T cell expansion and survival. IL-2 was critical for overcoming high-dose PD-L1–mediated inhibition of T cell expansion and rescued iT reg cell development at high concentrations of PD-L1 (PD-L1–Ig 60; Fig. 1, E–G). Using CFSE dye dilution in these assays, we found that PD-L1 simultaneously promoted the development of T reg cells and suppressed the development and activation of T eff cells. Furthermore, PD-L1 induced greater levels of Foxp3 expression per cell as indicated by the mean fluorescence intensity (MFI) of GFP expression (Fig. 1 G). Thus, PD-L1 and TGF-β have synergistic roles in regulating Foxp3+ iT reg cell development.

PD-L1–induced CD4+Foxp3+ T reg cells suppress CD4+ T eff cells

To assess whether PD-L1–induced iT reg cells not only express Foxp3 but also function as suppressor T cells, we treated naive T cells with TGF-β plus control Ig or PD-L1–Ig beads and sorted Foxp3.GFP+ iT reg cells after 3 d of culture. Sorted iT reg cells were then cultured in a standard suppression assay with CD4+CD25− responder T cells and bead-bound anti-CD3/anti-CD28 plus PD-L1–Ig for another 3 d. Both PD-L1 iT reg cells and control iT reg cells could suppress the proliferation of WT T eff cells to a similar extent as that measured by [3H]thymidine incorporation (Fig. 2 A). To evaluate whether PD-L1 iT reg cells or control iT reg cells affect the suppression of T eff cell proliferation on a per cell basis, we performed additional suppression assays measuring CFSE dilution of T eff cells. We cultured either CD45.1−PD-L1 iT reg cells or control iT reg cells with CD45.1+ T eff cells plus bead-bound anti-CD3/anti-CD28 and PD-L1–Ig for 3 d and analyzed CFSE dilution in CD45.1+ T eff cells by flow cytometry (Fig. 2 B). PD-L1 iT reg cells reduced T eff cell expansion at the single cell level to a greater extent than control iT reg cells (1.5-fold greater, P = 0.006), as measured by the division index (Fig. 2 C). These studies demonstrate that PD-L1 has a significant role in inducing the development of functional Foxp3+ iT reg cells.

PD-L1 enhances and maintains Foxp3 expression on iT reg cell and augments suppression at low T reg/T eff cell ratios

Our studies thus far could not discriminate whether PD-L1 only controls iT reg cell development or also has a role in maintaining iT reg cell function. Recent studies indicate that continued Foxp3 expression is necessary for sustaining T reg cell function (Kim et al., 2007; Williams and Rudensky, 2007). Therefore, we investigated whether PD-L1 influences the maintenance of Foxp3 expression and iT reg cell suppressive function. We induced T reg cell development by culturing naive CD45.1− T cells with either PD-L1–Ig beads or control Ig beads plus TGF-β. After 3 d of culture, PD-L1 iT reg cells or control iT reg cells were sorted by Foxp3.GFP+ expression and used in secondary cultures to evaluate the maintenance of Foxp3 expression and iT reg cell suppressive capacity. The sorted PD-L1 or control iT reg cell were cultured with CD4+CD25−CD45.1+ T eff cells, in the presence of either...
PD-L1–Ig beads or control Ig beads for 3 d (Fig. 3 A). Foxp3. GFP expression was then assessed by flow cytometry. When iT reg cells were stimulated with control Ig beads, Foxp3 was better maintained in iT reg cells originally induced with PD-L1–Ig beads as compared with control Ig beads (24.2% positive vs. 7.07% positive; Fig. 3 B, top). In addition, stimulation of control iT reg cells or PD-L1 iT reg cells with PD-L1–Ig beads significantly enhanced the percentage of iT reg cells maintaining Foxp3 expression (16.1% for control iT reg cells vs. 38% for PD-L1 iT reg cells; Fig. 3 B, bottom). Interestingly, iT reg cells that were both induced and restimulated in the presence of PD-L1–Ig maintained the greatest percentage of Foxp3–expressing cells (38%; Fig. 3, B and C). These studies demonstrate that PD-L1 strategically regulates Foxp3 expression in CD4 T cells at two stages: (1) during the induction phase of naive T cell conversion to iT reg cells and (2) during the effector phase of iT reg cell–mediated suppression. These findings point to a novel role for PD-L1 in regulating the stability of iT reg cells in the periphery.

We next tested whether the presence of PD-L1 could influence the efficiency of suppression by iT reg cells. Foxp3. GFP+ iT reg cells were sorted and cultured with naive CD4+CD25−CD45.1+ T eff cells plus either PD-L1–Ig or control Ig beads at a variety of iT reg/T eff cell ratios, as graphically depicted in Fig. 3 A (right). PD-L1–Ig enhanced iT reg cell suppressive function at a low T reg/T eff cell ratio of 1:4 (46% suppression using PD-L1 bead vs. 3% suppression using control bead, P = 0.0149; Fig. 4, A and B). The effect was less pronounced when T reg cells were more numerous, but PD-L1–Ig iT reg cells still showed improved suppressive capacity (Fig. 4 A). Because T eff cells may be directly inhibited by the presence of PD-L1 on beads during the suppression assay, we tested the suppressive capacity of PD-L1–Ig iT reg cells or control iT reg cells in the presence of TCR stimulation.
alone (i.e., control Ig beads) over a range of T reg/T eff cell ratios (Fig. 4 C). PD-L1 iT reg cells are far more effective than control iT reg cells at limiting the proliferative capacity of T eff cells at very low T reg/T eff cell ratios. Collectively, these results show that PD-L1 enhances the efficiency of suppression by iT reg cells.

**PD-L1 deficiency leads to impaired T reg cell conversion in vivo**

To investigate the role of PD-L1 in iT reg cell development and function in vivo, we compared iT reg cell conversion and function in Rag−/− mice because iT reg cells can spontaneously develop from naive T cells in a lymphopenic environment (Bloom et al., 2008; Calzascia et al., 2008; Winstead et al., 2008). We adoptively transferred naive CD4+CD62L+Foxp3.GFP+ T cells into PD-L1−/−PD-L2−/−Rag−/− or WT Rag−/− mice and analyzed the CD4+ T cells for Foxp3 expression at days 14–17 after transfer by flow cytometry. It should be noted that PD-L1−/− versus PD-L1−/−PD-L2−/− APCs are similarly deficient in their ability to induce Foxp3+ iT reg cells in the presence of TGF-β and anti-CD3 (unpublished data). There were ~10-fold fewer Foxp3.GFP+ cells in PD-L1−/−PD-L2−/−Rag−/− recipients compared with WT Rag−/− recipients (Fig. 5, A and B). These data demonstrate the critical role of PD-L in regulating the de novo generation and/or maintenance of Foxp3+ iT reg cell in vivo. We next compared Foxp3+CD4+ T eff cell responses in PD-L1−/−PD-L2−/−Rag−/− and WT Rag−/− recipients by measuring cytokine production by

---

**Figure 4. PD-L1 enhances the efficiency of iT reg cell-mediated suppression of T eff cells.** (A) Foxp3.GFP+ iT reg cells were sorted and cocultured with naive CD4+CD25−CD45.1+ T eff cells plus either PD-L1-Ig beads or control Ig beads (at various T reg/T eff cell ratios). 72 h later, cultures were pulsed with [3H]thymidine for 12–14 h. P < 0.0009 at a 1:4 ratio cultured with PD-L1 beads (comparing T eff + iT reg vs. T eff cells). Data represent the mean proliferation ± SD and are representative of at least four independent experiments. (B) Quantification of suppression at 1:4 ratio of T reg/T eff cells. P = 0.014. Data represent the mean ± SD and are representative of at least four independent experiments. (C) PD-L1 iT reg cells suppress T eff cells more effectively than control iT reg cells. CD4+CD62L+Foxp3.GFP+ naive T cells were induced with PD-L1 or control beads in the presence of TGF-β. GFP+ iT reg cells were sorted and co-cultured at the indicated T reg/T eff cell ratios with CFSE-labeled CD4+CD25−Thy1.1 T eff cells and beads coated with anti-CD3 and anti-CD28 (in the absence of PD-L1) for 3 d. Graphs are representative of three experimental replicates and data are representative of three independent experiments.
Figure 5. Attenuated iT reg cell development in PD-L1−/−PD-L2−/− mice in vivo. (A) CD4+CD62L+Foxp3.GFP− cells were adoptively transferred i.v. into the tail veins of WT Rag−/− or PD-L1−/−PD-L2−/−Rag−/− mice. Spleens and lymph nodes were analyzed for Foxp3.GFP expression 14–17 d after transfer. (B) Quantitation of Foxp3.GFP expression from independent mice depicted in A. Data represent the mean ± SE of five independent mice. (C and D) Analysis of IL-17+ and IFN-γ+ T eff cells by intracellular cytokine staining (C) and ratios of IL-17–producing T eff/T reg cells and IFN-γ–producing T eff/T reg cells from WT Rag−/− or PD-L1−/−PD-L2−/−Rag−/− mice 14–17 d after transfer (D). Data represent the mean ± SD of n = 5 mice per group and are representative of two independent experiments.
Foxp3–CD4+ cells 14–17 d after naive T cell transfer. There were comparable numbers of IL-17 and IFN-γ-producing cells generated in PD-L1−/−PD-L2−/−Rag−/− and WT Rag−/− recipients (Fig. 5 C). However, IL-17− and IFN-γ-producing T eff cells highly outnumbered the iT reg cells in PD-L1−/−PD-L2−/−Rag−/− recipients as a result of the paucity of iT reg cells in these mice (Fig. 5 D), resulting in a dramatic increase in the T eff/T reg cell ratio.

**PD-L1−/−PD-L2−/− Rag−/− mice develop fatal immune-mediated pulmonary damage after transfer of naive CD4+Foxp3− T cells**

To evaluate the in vivo consequences of the skewed T reg/T eff cell ratio in PD-L1−/−PD-L2−/−Rag−/− recipients, we monitored PD-L1−/−PD-L2−/−Rag−/− and WT Rag−/− mice after transfer of CD4+CD62L+Froxp3, GFP− T cells (Fig. 6 A). PD-L1−/−PD-L2−/−Rag−/− recipients exhibited rapid and dramatic weight loss beginning on day 6 after transfer, with a reduction of 17.6 ± 5.7% (P ≤ 0.0001) of the pretransfer body weight within 2 wk, compared with 2.65 ± 2.7% (P = 0.100) weight loss in WT Rag−/− recipients. There was a marked increase in cellularity of the lymph nodes and spleen in PD-L1−/−PD-L2−/−Rag−/− compared with WT Rag−/− adoptive transfer recipients on days 14–17 after transfer (Fig. 6 B).

Strikingly, all PD-L1−/−PD-L2−/−Rag−/− recipients were moribund within 17 d after transfer of naive T cells (n = 12) in marked contrast to the survival of all WT Rag−/− recipients (n = 8; Fig. 6 C). In a separate experiment, we examined PD-L1−/−PD-L2−/−Rag−/− and WT Rag−/− recipients histologically on days 14–17 after transfer (Fig. 6 D). The lungs of PD-L1−/−PD-L2−/−Rag−/− recipients showed widespread severe perivascular, peribronchial, and interstitial infiltrates consisting predominantly of mononuclear cells and some neutrophils. Alveolar walls were markedly thickened by the inflammation, and there was severe alveolar consolidation and edema. This potentially reduced gas exchange in these mice. In contrast, all the WT Rag−/− recipients displayed a minimal degree of inflammation and absence of alveolar consolidation in the lungs. Both groups also showed varying degrees of mild colitis on days 14–17 after adoptive transfer. Analysis of brain, heart, pancreas, kidney, esophagus, stomach, small intestine, and skin revealed only mild scattered inflammation with no significant differences between WT and PD-L1−/−PD-L2−/−Rag−/− recipients (unpublished data).

To ascertain the critical role for PD-L1 in vivo, we transferred naive CD4 T cells to Rag−/− recipients treated with anti–PD-L1 blocking antibody (Fig. S1) and monitored the mice for 3–4 wk. Mice were sacrificed to assess T reg cell development and immunopathology. A significant defect in de novo iT reg cell development was observed in Rag−/− mice given anti–PD-L1 mAb compared with isotype control in both the spleen (isotype = 6.35% vs. anti–PD-L1 = 2.73%, P = 0.0318) and mesenteric lymph nodes (isotype = 30.2% vs. anti–PD-L1 = 18%, P = 0.0219). The lungs of Rag−/− mice treated with anti–PD-L1 mAb showed moderate to severe perivascular, peribronchial, and interstitial inflammation, consisting of mononuclear cells and a few scattered neutrophils. Thus, similar to the PD-L1−/−PD-L2−/−Rag−/− recipients of naive CD4 T cells, WT Rag−/− mice given anti–PD-L1 mAb exhibited defects in iT reg cell generation and developed pulmonary pathology. Collectively, these studies demonstrate a key role for PD-L1 in iT reg cell development in vivo.

**PD-L1 antagonizes the Akt–mTOR signaling cascade during the induction of iT reg cells**

Recent studies have shown notable differences in signaling pathways used by CD4+ T eff cells compared with T reg cells. In particular, Akt signaling is essential for naive T cell activation and proliferation but dispensable for T reg cell development and function (Battaglia et al., 2006; Coenen et al., 2007; Gao et al., 2007; Qu et al., 2007; Strauss et al., 2007; Haxhinasto et al., 2008; Sauer et al., 2008). These findings led us to hypothesize that PD-L1 may mediate T reg cell conversion by antagonizing the Akt signaling pathway. To test this, we cultured naive T cells in the presence of PD-L1−Ig or control Ig beads for 18 h and then measured phosphorylation of Akt, mTOR, and S6. Intracellular staining for phospho-Akt and phospho-mTOR revealed significantly diminished levels of Akt and mTOR phosphorylation when naive T cells were cultured in the presence of increasing quantities of PD-L1 relative to control Ig (MFIs of phospho-Akt and phospho-mTOR were significantly down-regulated; Fig. 7, A–D). As a downstream target of the mTOR–regulated p70 S6 kinase, phosphorylation of S6 ribosomal protein reflects the sustained activation of the Akt–mTOR pathway. Upon culture of naive CD4 T cells with increasing amounts of PD-L1, we observed a marked decrease in phospho–S6 as compared with control (Fig. 7, E and F). Furthermore, PD-L1 up-regulated the expression of PTEN (phosphatase and tensin homologue deleted on chromosome 10), a phosphoinositol 3,4,5-triphosphatase important for antagonizing PI3K signaling, demonstrating that PD-L1 antagonizes the Akt pathway during T reg cell differentiating conditions (Fig. 7, G and H). Western blots assessing the specific down-regulation of phospho-Akt, phospho-mTOR, and phospho-S6 and up-regulation of PTEN confirmed phospho-flow cytometry data (unpublished data).

Down-regulation of the MAP kinase signaling cascade has also been implicated in TGF-β–mediated T reg cell development (Luo et al., 2008). We questioned whether PD-L1 might regulate T reg cell differentiation by modulating the MAP kinase pathway. Stimulation of naive T cells with increasing amounts of PD-L1 Ig attenuated the phosphorylation of ERK2/p42 (Fig. S2). These data further substantiate our hypothesis that the PD-L1–PD-1 pathway truncates signaling cascades downstream of TCR signaling, preferentially converting naive T cells toward a Foxp3+ T reg cell lineage.

**DISCUSSION**

In this paper, we demonstrate a novel function for PD-L1 in promoting the development and sustaining the function of
Figure 6. Dramatic weight loss, severe pulmonary inflammation, and fatal inflammatory disorder develop in PD-L1−/−PD-L2−/− Rag−/− mice after adoptive transfer of naive CD4+CD62LhiFoxp3.GFP− T cells. Sorted CD4+CD62LhiFoxp3.GFP− cells were adoptively transferred i.v. into WT Rag−/− or PD-L1−/−PD-L2−/− Rag−/− mice. (A and B) Clinical manifestations shown are the following: percentage of weight loss of mice after adoptive transfer of CD4+CD62LhiFoxp3.GFP− cells into WT Rag−/− or PD-L1−/−PD-L2−/− Rag−/− (P < 0.001; n = 5 mice per group; A) and quantified lymph node (axillary, brachial, and inguinal) cellularity (B). Data represent the mean ± SD and represent two independent experiments. (C) Survival of mice after adoptive transfer of naive CD4+CD62LhiFoxp3.GFP− T cells was monitored for 30 d. PD-L1−/−PD-L2−/− Rag−/−, n = 12; WT Rag−/−, n = 8. (D) Hematoxylin and eosin-stained paraffin sections of lung tissue obtained on days 14–17 after transfer of naive CD4+CD62LhiFoxp3− T cells. Bars: (top, 40x) 500 µm; (bottom, 400x) 50 µm. PD-L1−/−PD-L2−/− Rag−/−, n = 9; WT Rag−/−, n = 10. Data represent more than four independent experiments.
iT reg cells. There was a profound defect in conversion of naive CD4 T cells into Foxp3+ iT reg cells in the absence of PD-L1. Consistent with this observation, PD-L1 presented on beads (along with anti-CD3 and anti-CD28) could induce the development of functional Foxp3+ iT reg cells, demonstrating that PD-L1 is responsible for promoting iT reg cell development. Although TGF-β signaling is important for the conversion of naive CD4+ T cells toward Foxp3-expressing cells with suppressive capacity (Chen et al., 2003; Fantini et al., 2004), PD-L1 could induce T reg cells in the absence of exogenous TGF-β, suggesting that PD-L1 signaling alone can serve to promote iT reg cell development. Our signaling studies support the conclusion that PD-L1 reduces signaling of the Akt–mTOR pathway in naive T cells, which is critical for iT reg cell development. There was a profound defect in conversion of naive CD4 T cells into Foxp3+ iT reg cells in the absence of PD-L1. Consistent with this observation, PD-L1 presented on beads (along with anti-CD3 and anti-CD28) could induce the development of functional Foxp3+ iT reg cells, demonstrating that PD-L1 is responsible for promoting iT reg cell development. Although TGF-β signaling is important for the conversion of naive CD4+ T cells toward Foxp3-expressing cells with suppressive capacity (Chen et al., 2003; Fantini et al., 2004), PD-L1 could induce T reg cells in the absence of exogenous TGF-β, suggesting that PD-L1 signaling alone can serve to promote iT reg cell development. Our signaling studies support the conclusion that PD-L1 reduces signaling of the Akt–mTOR pathway in naive T cells, which is critical for iT reg cell development.
for their conversion into iT reg cells. Furthermore, we show that PD-L1 has a novel role in sustaining expression of Foxp3 in iT reg cells and in enhancing iT reg cell suppressive function. Thus, these studies reveal a new mechanism by which PD-L1 mediates T cell tolerance. PD-L1 can inhibit self-reactive T cell responses by promoting iT reg cell development and maintaining iT reg cell function.

Where does PD-L1 exert its critical effects on iT reg cell development and function? PD-L1 is widely expressed on hematopoietic and non-hematopoietic cells (Keir et al., 2008). There may be a critical interaction between PD-L1 expressing DC and T cells during the induction of T reg cell development from naive T cells (Brown et al., 2003). PD-L1 is also expressed on T cells; however, WT T cells transferred into PD-L1−/−/PD-L2−/− Rag−/− recipients could not convert to iT reg cells, suggesting that T cell–T cell interaction via PD-L1 is not sufficient to drive naive T cell conversion. Instead an interaction with the host environment is crucial for T reg cell conversion.

Our bone marrow chimera studies have demonstrated an important role for PD-L1 on non-hematopoietic cells in mediating tissue tolerance (Keir et al., 2006). Our findings of critical roles for PD-L1 in the development and maintenance of iT reg cell function suggest that PD-L1 may protect tissues from the potentially pathogenic self-reactive T eff cells not only by inhibiting the function of T eff cells but also by increasing the frequency and function of T reg cells in the target tissue. By promoting de novo generation of iT reg cells in situ, PD-L1 may play a role in mediating immune privilege, especially in environments where TGF-β is present (e.g., placenta and eye).

PD-L1 may also exert its inhibitory effects on anti-tumor and anti-microbial immunity, at least in part by inducing T reg cell development and maintaining iT reg cell function. PD-L1 is expressed on a wide variety of tumors, and high levels of PD-L1 expression strongly correlate with unfavorable prognosis in several cancers (Dong et al., 2002; Iwai et al., 2002; Strome et al., 2003; Konishi et al., 2004; Thompson et al., 2004; Blank et al., 2005; Hirano et al., 2005; Ohigashi et al., 2005; Dorfman et al., 2006; Wu et al., 2006; Inman et al., 2007; Nakanishi et al., 2007; Nomi et al., 2007; Zhang et al., 2008). The numbers of Foxp3+ T cells within tumors also correlates with a poor prognosis. Our work provides a mechanism by which high PD-L1 expression can lead to increased numbers of Foxp3+ T reg cells and thus, poor prognosis. Increased PD-L1 expression by tumor cells may induce and maintain iT reg cells in the periphery, thereby augmenting the suppression of anti-tumor T cell responses and allowing tumor progression.

Increased T reg cells are also seen during chronic infections (Belkaid, 2008), correlating with a lack of sterilizing immunity. During persistent infection by Helicobacter pylori, PD-L1 is up-regulated on gastric epithelial cells (Das et al., 2006). Blocking PD-L1 on gastric epithelial cells enhances CD4 T eff cell function and prevents the generation of CD4+CD25hiFoxp3+ T reg cells in vitro (Beswick et al., 2007). During chronic viral infections, PD-L1 has a key role in limiting the function of exhausted CD8 T cells (Barber et al., 2006; Sharpe et al., 2007). PD-L1 blockade reinvigorates the function of these T cells and enhances viral clearance. Thus, PD-L1 may exert inhibitory effects in chronic infections by promoting iT reg cell development and maintaining iT reg cell function, as well as by inhibiting anti-microbial T eff cell function.

PD-L1−/−/PD-L2−/−Rag−/− recipients of Foxp3+ naïve T cells resemble scurfy mice, which have a defect in Foxp3 and die by 3 wk of age as a result of multi-organ infiltration of CD4+CD8+ T cells (Clark et al., 1999; Brunkow et al., 2001; Schubert et al., 2001). PD-L1−/−/PD-L2−/−Rag−/− recipients had a marked deficit in Foxp3+ T reg cell development, altering the T reg/T eff cell ratio. These findings illustrate the important role of PD-L1 in regulating the dynamic balance between T eff and T reg cells in vivo. PD-L1 may do the following: (a) control T reg cell development in lymphoid organs, which is important for immune homeostasis; (b) promote T reg cell development at target tissues, protecting against immune-mediated tissue damage, and (c) sustain and enhance T reg cell function within an inflammatory microenvironment, effectively counterbalancing the pathogenic T eff cells.

Foxp3 is a transcription factor only expressed in the T reg cell lineage (Fontenot et al., 2003; Hori et al., 2003; Vignali et al., 2008). Along with contributing a distinct genetic signature to T reg cells, Foxp3 conveys regulatory activity to nT reg cells, iT reg cells, and, upon ectopic expression, in conventional T cells (Schubert et al., 2001; Fontenot et al., 2003, 2005; Hori et al., 2003; Gavin et al., 2007; Hill et al., 2007). PD-L1 can induce and maintain the expression of Foxp3 in iT reg cells, suggesting that PD-L1 may function to stabilize and sustain T reg cell function in the periphery, similar to effects reported for TGF-β (Marie et al., 2005; Pyzik and Piccirillo, 2007). This maintenance of Foxp3 expression may explain both the increased efficiency of suppression seen for T reg cells cultured with PD-L1 and the lack of T reg cell conversion in the PD-L1−/−/PD-L2−/−Rag−/− adoptive transfer recipients. It has been suggested that although Foxp3 expression is a salient feature of the regulatory cell signature, it is not the master regulator of T reg cell development (Ramsdell, 2003; Collison et al., 2007; Hill et al., 2007). In fact, it is increasingly clear that other factors (such as TGF-β and IL-2; Ramsdell, 2003; Setoguchi et al., 2005) induce key changes within the regulatory cell transcriptome that contribute to the identity of T reg cells separate from, and in addition to, the effects attributed to Foxp3 (Lin et al., 2007). In vivo, TGF-β production is maintained at a basal level and up-regulated with inflammation. Interestingly, endogenous TGF-β was not sufficient to induce and/or maintain Foxp3+ iT reg cells in PD-L1−/−PD-L2−/− Rag−/− recipients. In complementary studies, PD-L1 blockade similarly impaired iT reg cell development in vivo, showing that loss of PD-L1 can diminish the effect of TGF-β, which is critical for iT reg cell identity and function. This indicates that PD-L1 contributes unique and essential signals that drive iT reg cell development and function.
We find that PD-L1 attenuates the Akt signaling pathway during the conversion of naive T cells to T reg cells by reducing the phosphorylation of Akt and its downstream substrates mTOR and S6 while simultaneously augmenting PTEN. Previous work has shown that truncation of TCR signaling and inhibition of the Akt–mTOR signaling axis is critical for T reg cell development (Qu et al., 2007; Strauss et al., 2007; Haxhinasto et al., 2008; Long and Buckner, 2008; Sauer et al., 2008). In these studies, drug inhibitors (FK506, rapamycin) or retrovirally modified Akt were used to demonstrate the role of Akt and mTOR in T reg cell development. We provide the first demonstration of a naturally occurring protein that can inhibit the Akt–mTOR cascade and regulate the development of T reg cells.

Similar to the Akt–mTOR pathway, recent data indicate an important role for the MAP kinase cascade in iT reg cell development (Adler et al., 2007; Huber et al., 2008; Luo et al., 2008). We found that PD-L1 attenuated the phosphorylation of p42/ERK2, suggesting that PD-L1 may mediate the induction of T reg cells by modulating ERK2 activity and, hence, the MAP kinase signaling cascade. No discernable effects of PD-L1–Ig on p38 were detected (unpublished data).

There is great interest in generating T reg cells ex vivo as a therapy for autoimmune diseases and transplant rejection (Roncarolo and Battaglia, 2007). However, recent studies indicate that T reg cells exhibit functional plasticity and can produce proinflammatory cytokines at the site of inflammation (Joetham et al., 2008; Yang et al., 2008). Thus, in order for T reg cell therapy to be a viable approach, it is critical to find ways to maintain and enhance the suppressive function of T reg cells. Our work suggests that administration of PD-L1–Ig or PD-1 agonists may harness the therapeutic potential of iT reg cells by providing a novel means of sustaining and enhancing their function in vivo while concomitantly suppressing the expansion and functions of activated T eff cells.

**MATERIALS AND METHODS**

*Mice.* 6–8-wk-old WT C57BL/6 and CD45.1 (B6.SJ–Pepc<sup>+/−</sup> Pepc<sup>+/−</sup>) mice were purchased from The Jackson Laboratory. PD-L1<sup>−/−</sup>/PD-L2<sup>−/−</sup> (Keir et al., 2006) and PD-L1<sup>−/−</sup> (Latchman et al., 2004) mice were generated in our laboratory. 2D2 TCR Tg mice Foss3–IRE-GFP knockin mice (Foss3;GFP; Bettelli et al., 2006) were generated in our laboratory by crossing 2D2 TCR Tg mice (Bettelli et al., 2003) with Foss3;GFP reporter mice. PD-L1<sup>−/−</sup>/PD-L2<sup>−/−</sup> Rag 2<sup>−/−</sup> mice were generated by breeding PD-L1<sup>−/−</sup>/PD-L2<sup>−/−</sup> with Rag 2<sup>−/−</sup> mice (B6.129S6-Rag2<sup>−/−</sup>N12; Taconic). Genotypes were verified by PCR and flow cytometry. Harvard Medical School is accredited by the American Association of Accreditation of Laboratory Animal Care. Mice were maintained in a pathogen-free facility and used according to the Harvard Medical School Standing Committee on Animals and National Institutes of Health Animal Care Guidelines. Animal protocols were approved by the Harvard Medical School Standing Committee on Animals.

*Reagents.* The following anti–mouse antibodies were used in cell surface staining, intracellular cytokine staining, and epoxy bead conjugation: anti–CD16/CD32 (Fc Block), CD4 PerCP-Cy5.5 (clone RM4-5), CD62L PE (clone ME1-14), IL-2 APC (clone JES6-5H4; eBioscience), CD45.1 APC (clone A20), IL-17 PE (clone TC11-18H10), and IFN-γ PE (clone XM1G12; BD). Anti–CD3 (clone 2C11) plus anti–CD28 (clone 37.51) were used for bead conjugation and were purchased from Bio X Cell. Cells were sorted on a FACSAria cell sorter (BD). Cell surface staining was performed at 4°C in FACS Buffer (1% FCS, PBS, 2 mM EDTA; Invitrogen). CFSE was purchased from Invitrogen.

**Cell purification.** Naive CD4<sup>+</sup>CD62L<sup>+</sup>Foxp3.GFP<sup>−</sup> T cells were isolated from the spleen and lymph nodes (axillary, brachial, and inguinal) of male C57BL/6 Foxp3.GFP or 2D2 Foxp3.GFP reporter mice. Single cell suspensions were made by mechanical dissociation. After red blood cell lysis with ACK buffer (Invitrogen), cells were washed and isolated by incubation with CD4 microbeads, positively selected through LS columns (Miltenyi Biotec), and stained with anti–CD4 PerCP-Cy5.5 (clone RM4-5; eBioscience) and anti–CD62L APC (clone MEL-14; eBioscience) before cell sorting on a FACSAnA cell sorter. Naive CD4<sup>+</sup>CD62L<sup>+</sup>Foxp3.GFP<sup>−</sup> T cells were always >99.2% pure.

**In vitro iT reg cell development.** Anti–CD3 (clone 2C11; Bio X Cell) plus anti–CD28 (clone 37.51; Bio X Cell) were covalently attached to Dynabead M450 glycidyl ether beads according to the manufacturer’s directions (Invitrogen). We ensured equal loading of proteins during preparation by keeping constant the total amount of protein (antibodies and fusion proteins) at 5 µg per 10<sup>6</sup> beads as previously described (Broeren et al., 2000; Riley et al., 2002). In general, 10<sup>5</sup> beads were coated with 1 µg of anti–CD3 (20% of total protein), 1 µg of anti–CD28, and either 60% control human IgG1 (referred to control Ig beads; Bio X Cell) or 40% PD-L1–IgG1 Fc fusion protein (referred to as PD-L1–Ig beads; 40; R&D Systems) plus 20% control human IgG1 Fc. In some experiments, increasing amounts of PD-L1–Ig was used to coat the epoxy beads (20, 40, and 60% of total protein per 10<sup>5</sup> beads = 1, 2, and 3 µg of PD-L1–Ig per 10<sup>5</sup> beads). In these cases, the remaining surface of the beads were coated with control human IgG1. Covalent attachment of the proteins to the beads was performed in NaPO<sub>4</sub> buffer for 24 h at room temperature on a Nutating Mixer (Lab-Tech Incorporated). Beads were then washed three times in PBS over a magnetic column and suspended in complete media before use.

CD4<sup>+</sup>CD62L<sup>+</sup>Foxp3<sup>−</sup> naive T cells were cultured with beads at a fixed ratio of 1:5 (T cells/beads). In brief, 1–2 × 10<sup>5</sup> T cells were plated at 10<sup>6</sup> cells/ml in a 24-well flat-bottom tissue culture plate with beads in complete media consisting of RPMI-1640 with L-glutamine (Invitrogen) supplemented with 10% FCS (Sigma-Aldrich), penicillin-streptomycin (100 U penicillin and 100 µg streptomycin; Invitrogen), 12 mM HEPES (Invitrogen), and 50 µM β-mercaptoethanol (Sigma-Aldrich) plus 2 ng/ml TGF-ß (R&D Systems) and 20 U/ml IL-2 (R&D Systems) for 3 d at 37°C with 5% CO<sub>2</sub>. In some experiments, sorted CD4<sup>+</sup>CD25<sup>+</sup>CD62L<sup>+</sup>Foxp3<sup>−</sup> naive T cells were labeled with 1 µM CFSE before culture with beads and TGF-ß for 3 d at 37°C with 5% CO<sub>2</sub>. T cells were then stained for Foxp3 expression (eBioscience). To further evaluate the effect of IL-2 on PD-L1–mediated T reg cell development, some experiments were conducted in the absence of exogenous rh-IL-2 (as noted in the Fig. 1 legend). To analyze the effect of TGF-ß on PD-L1–mediated T reg cell development, we performed some experiments using a range of TGF-ß concentrations (0.03–8 ng/ml).

In some experiments, APCs were isolated from WT or PD-L1<sup>−/−</sup> mice. In brief, T cells were depleted from spleens of WT or PD-L1<sup>−/−</sup> mice with CD4 and CD8 microbeads (Miltenyi Biotec). Remaining splenocytes were irradiated and co-cultured with naive CD4<sup>+</sup>CD62L<sup>+</sup>Foxp3<sup>−</sup> T cells plus anti–CD3 and a range of TGF-ß concentrations (0.5–8 ng/ml).

**In vitro suppression assays.** Naive T cells were induced toward T reg cell development in vitro using PD-L1 or control beads in the presence of 20 U/ml TGF-ß and IL-2 (R&D Systems) for 3 d, at which time Foxp3<sup>−</sup> T cells were sorted on a FACSAria (BD) cell sorter. Foxp3<sup>−</sup> T reg cells were then co-cultured with sorted CD4<sup>+</sup>CD25<sup>+</sup>CD45.1<sup>−</sup> naive T eff cells and stimulated with PD-L1 beads (containing anti–CD3 (20%), anti–CD28 (20%), PD-L1–Ig (40%), and control Ig (20%)) for 3 d. Proliferation of T cells was determined by incorporation of [H]thymidine (1 µCi/well) for 12–14 h. Suppression assays were performed using a constant number of T eff cells (10<sup>4</sup>) and by addition of decreasing numbers of Foxp3<sup>+</sup> T reg cells plus a 5:1
ratio of beads to T eff cells. The percentage of suppression of effector cell proliferation was calculated based on the proliferation of T eff cells with control versus PD-L1–Ig beads in the absence of T reg cells.

For CFSE dilution experiments, CD4+CD25−CD45.1+ naive T eff cells were labeled with 1 nM CFSE for 10 min in RPMI-1640 (serum free) and washed twice with 100% FBS and twice with complete media before culture. 10^5 T eff cells were cultured with 10^5 iT reg cells and PD-L1–Ig beads (5:1 bead/T eff cell ratio) in 96-well flat-bottom plates (BD). 72 h later, CD4+CD45.1+ T eff cells were gated and analyzed for CFSE dilution. Divison index (defined as the mean number of divisions that a cell has undergone) was calculated using FlowJo. Proliferation analysis software (Tree Star, Inc.). For CFSE dilution experiments interrogating the suppressive capacity of PD-L1–Ig reg cells versus control iT reg cells, naive T eff cells were induced toward T reg cell development in vitro using PD-L1 or control beads in the presence of TGF-β and IL-2 for 3 d, as described in the previous section, FlowJo. G. F. T cells were sorted on a FACSAna cell sorter. Control and PD-L Fospx3.GFP+ iT reg cells were then co-cultured with CFSE-labeled CD4+CD25−Thy1.1+ T eff cells and stimulated with control beads (containing anti-CD3, anti-CD28, and control Ig). 10^5 CD4+CD25−Thy1.1+ T eff cells were cultured with varying numbers (3 × 10^4–10^5) of PD-L1 (iT reg cells or control iT reg cells plus control Ig beads (5:1 bead/T eff cell ratio) in 96-well flat-bottom plates (BD)). 4 d later, CD4+Thy1.1+ T cells were gated and analyzed for CFSE dilution.

**In vivo adoptive transfer.** Naive CD4+ T cells were isolated from spleens and lymph nodes of C57BL/6 mice. CD4+CD25−Fospx3.GFP+ T cells were sorted on a FACSAna as described in the previous section, and 1–1.5 × 10^6 CD4+CD25−Fospx3.GFP+ iT reg cells were i.v. injected into the tail veins of PD-L1−/−PD-L2−/− Rag−/− or WT Rag−/− mice. Mice were monitored and weighed for 14–17 d and euthanized for histological and cellular analysis. In some experiments, surviving mice were monitored for 30 d after T cell transfer. Organs were fixed in formalin, dehydrated, and embedded in parafin. 5-µm sections stained with hematoxylin and eosin were independently evaluated by two pathologists in a blinded fashion. Digital photomicrographs were acquired using DP Controller software (Olympus) driving a DP71 camera (Olympus) mounted on a BH-2 light microscope (Olympus). Image sizes were reduced using Photoshop CS3 software (Adobe).

**Intracellular cytokine staining.** Spleen and lymph node (auxillary, brachial, inguinal, and mesenteric) cells were isolated and re triturated with 50 ng/ml PMA and 500 ng/ml ionomycin (Sigma-Aldrich) for 4 h with Golgistop (BD) being added during the last 3 h of stimulation. After Fc block, cells were stained with anti-CD4-PE-Cy5.5, fixed with 4% paraformaldehyde, and permeab lized with Cytotox/Cytoperm solution (BD). Intracellular staining with IL-17 PE, IFN-γ PE, or APC and IL-2 APC was performed in Cytoperm buffer (BD) according to the manufacturer’s protocol. Cells were washed twice in Cytoperm buffer and twice in FACS buffer before data acquisition on a FACSCalibur (BD) and analysis by FlowJo software.

**Phospho-flow cytometry.** Naive CD4+ T cells were sorted from 2D2 Fospx3.GFP reporter mice and cultured with either PD-L1–Ig beads or control beads in the presence of 2 ng/ml TGF-β and 20 U/ml IL-2 for 18 h. Signaling molecules were assessed with antibodies against phospho-Akt Ser473 Alexa Fluor 647 (clone D9E), phospho-mTOR Ser2448 (clone D57.2.E), and phospho-S6 Ser235/236 Alexa Fluor 647 (clone D57.2.E), phospho-mTOR Ser2448 (clone D9E), phospho-mTOR Ser2448 (clone D57.2.E), and PTEN Alexa Fluor 647 (clone 138G6; Cell Signaling Technology). Isotype control staining was performed using rabbit IgG isotype mAb Alexa Fluor 647 (DA1E; Cell Signaling Technology). p-mTOR was detected with anti- rabbit Alexa Fluor 647 secondary (Invitrogen). Intracellular staining was performed as described in the manufacturer’s protocol. In brief, T cells were collected and washed thoroughly with PBS in 96-well V-bottom plates. Cells were then fixed with 2% paraformaldehyde for 10 min at 37°C. After fixation, plates were prechilled on ice for 1 min before permeabilization by slowly adding ice-cold methanol to a final concentration of 90% methanol. Cells were then incubated on ice for 30 min for permeabilization before being washed with 1% FCS/PBS (incubation buffer). Cells were blocked with 10% FCS/PBS for 10 min at room temperature and subsequently stained with the antibodies listed in this section for 1 h at room temperature. After incubation, cells were washed four times with incubation buffer and brought up in PBS before analysis.

**Statistical analysis.** Statistical analysis of Fospx3+ T reg cell development, T eff cell proliferation, intracellular cytokine production, and phospho-flow cytometry was performed using Student’s t-tests in StatView (SAS Institute Inc.). PD-L1–Ig titration, TGF-β titration, and percentage of weight loss were analyzed by ANOVA using StatView. Prism (GraphPad Software, Inc.) was used for Log-rank tests comparing WT Rag−/− and PD-L1−/−PD-L2−/− Rag−/− survival after transfer of naive T cells. P-values <0.05 were considered statistically significant.

**Online supplemental material.** Fig. S1 shows that blockade of PD-L1 attenuates T reg cell development in vivo. Fig. S2 shows that PD-L1–Ig attenuates the MAPK signaling pathway during iT reg cell differentiation. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20090847/DC1.

The authors thank Yin Wu and Robert Ortega for technical support, Dr. Sun J. Lee for western blotting assistance, Peter T. Sage for critical reading of the manuscript, and Dr. Marish J. Butte for technical advice, statistical analysis, and critical reading of the manuscript.

This work was supported by grants from the National Institutes of Health (AI38310 and AI40614 to A.H.; Sharp; PI0 56299 to A.H. Sharp, V.K. Kuchroo, and G.J. Freeman) and The National Multiple Sclerosis Society (L.M. Francisco). G.J. Freeman draws royalties from patents regarding PD-L1. The authors declare that they have no other conflicts of interest.

Submitted: 17 April 2009
Accepted: 12 November 2009

**REFERENCES**


