Peroxisome proliferator–activated receptor δ limits the expansion of pathogenic Th cells during central nervous system autoimmunity

Shannon E. Dunn,1,6 Roopa Bhat,1 Daniel S. Straus,5 Raymond A. Sobel,2 Robert Axtell,1 Amanda Johnson,1 Kim Nguyen,1 Lata Mukundan,3 Marina Moshkova,6 Jason C. Dugas,4 Ajay Chawla,3 and Lawrence Steinman1

Peroxisome proliferator–activated receptors (PPARs; PPAR-α, PPAR-δ, and PPAR-γ) comprise a family of nuclear receptors that sense fatty acid levels and translate this information into altered gene transcription. Previously, it was reported that treatment of mice with a synthetic ligand activator of PPAR-δ, GW0742, ameliorates experimental autoimmune encephalomyelitis (EAE), indicating a possible role for this nuclear receptor in the control of central nervous system (CNS) autoimmune inflammation. We show that mice deficient in PPAR-δ (PPAR-δ−/−) develop a severe inflammatory response during EAE characterized by a striking accumulation of IFN-γ+IL-17A− and IFN-γ+IL-17A+ CD4+ cells in the spinal cord. The preferential expansion of these T helper subsets in the CNS of PPAR-δ−/− mice occurred as a result of a constellation of immune system aberrations that included higher CD4+ cell proliferation, cytokine production, and T-bet expression and enhanced expression of IL-12 family cytokines by myeloid cells. We also show that the effect of PPAR-δ in inhibiting the production of IFN-γ and IL-12 family cytokines is ligand dependent and is observed in both mouse and human immune cells. Collectively, these findings suggest that PPAR-δ serves as an important molecular brake for the control of autoimmune inflammation.

Pathogenic Th1 and Th17 cells specific for myelin autoantigens are regarded as critical for both initiation and perpetuation of inflammation in multiple sclerosis and its animal model, experimental autoimmune encephalomyelitis (EAE; Diveu et al., 2008). Cytokines involved in the Th1 and Th17 axes of inflammation are detected in active lesions of multiple sclerosis patients (Windhagen et al., 1995; Lock et al., 2002; Kebir et al., 2009) and in the central nervous system (CNS) of mice with EAE (Langrish et al., 2005). Adoptive transfer studies indicate that both Th1 and Th17 cells can initiate the autoimmune cascade in this disease (Stromnes et al., 2008; Kroenke et al., 2008).

Peroxisome proliferator–activated receptors (PPARs) are members of the nuclear hormone receptor superfamily. This group of ligand-activated transcription factors regulates diverse processes including lipid and glucose homeostasis, cell proliferation and differentiation, and inflammation (Straus and Glass, 2007). PPARs are considered to be nutritional sensors, as they are bound and activated by fatty acid intermediates and can thus translate lipid levels into altered gene transcription. PPAR-γ binds oxidized and nitrosylated fatty acids as well as certain eicosanoids and prostaglandins (Schopfer et al., 2005; Shiraki et al., 2005), whereas PPAR-δ and PPAR-α have overlapping specificities for long-chain unsaturated fatty acids (Forman et al., 1997; Chawla et al., 2003; Hostetler et al., 2005). Upon binding their
In this paper, we show that PPAR-δ−/− mice exhibit enhanced Th cell expansion and cytokine production during EAE, as compared with WT mice, resulting in the enhanced accumulation of IFN-γ+ and IL-17A− and IFN-γ−/IL-17A+ CD4+ cells in the spinal cord and more severe demyelination. Moreover, we found that the PPAR-δ ligand GW0742 inhibits the production of IFN-γ and IL-12 family members in both mouse and human immune cells. Together, these findings suggest that PPAR-δ serves as an important molecular brake for the control of CNS autoimmune inflammation.

RESULTS AND DISCUSSION

PPAR-δ limits the development of CNS inflammation during EAE

To investigate the role of endogenously activated PPAR-δ in CNS autoimmunity, we compared the clinical course of EAE in WT versus PPAR-δ−/− mice. Upon immunization with myelin oligodendrocyte glycoprotein (MOG) p35-55 in CFA, WT mice developed classical relapsing-remitting EAE (Fig. 1, a and b). In contrast, PPAR-α appears to inhibit inflammation by shifting Th responses from Th1 to Th2 (Jones et al., 2003; Dunn et al., 2007; Gocke et al., 2009). How the PPAR-δ agonist GW0742 functions to attenuate EAE is not clear. The role of endogenously activated PPAR-δ in the control of CNS inflammation is also unknown.

In this paper, we show that PPAR-δ−/− mice exhibit enhanced Th cell expansion and cytokine production during EAE, as compared with WT mice, resulting in the enhanced accumulation of IFN-γ+ and IL-17A− and IFN-γ−/IL-17A+ CD4+ cells in the spinal cord and more severe demyelination. Moreover, we found that the PPAR-δ ligand GW0742 inhibits the production of IFN-γ and IL-12 family members in both mouse and human immune cells. Together, these findings suggest that PPAR-δ serves as an important molecular brake for the control of CNS autoimmune inflammation.
sexes (Fig. 1, a and b) but appeared more prominently in males (Table I). Thus, males were used for subsequent experiments.

To gain insight into why EAE failed to remit in PPAR-Δ⁻/⁻ mice, we sacrificed male WT and PPAR-Δ⁻/⁻ mice together at the time when WT mice displayed the first improvement of clinical signs and conducted a histological analysis of the brain and spinal cord. As compared with WT mice, PPAR-Δ⁻/⁻ mice displayed more inflammatory foci in the CNS (Fig. 1, c and d). The majority of parenchymal lesions were detected in the spinal cord in both groups; however, only in PPAR-Δ⁻/⁻ mice were there lesions in the cerebrum and cerebellum (Fig. 1 g). As compared with the WT group, PPAR-Δ⁻/⁻ mice also exhibited higher vacuolation/demyelination scores (mean ± SEM = 2.4 ± 0.4 in PPAR-Δ⁻/⁻ vs. 1.2 ± 0.2 in WT, P = 0.04). Consistent with histological findings, a higher number of CD4⁺ and CD19⁺ cells were recovered from spinal cords of PPAR-Δ⁻/⁻ versus WT mice during the acute phase of EAE (Fig. 1 f). Together, these results indicate that the more severe EAE that developed in PPAR-Δ⁻/⁻ mice was the result of immune-mediated myelin and axon damage.

**PPAR-Δ inhibits Th1 and Th17 autoimmunity in the periphery and CNS**

Because Th1 and Th17 cells can initiate and perpetuate the CNS inflammatory cascade in EAE (Diveu et al., 2008), we next examined the cytokine production by CNS-infiltrating CD4⁺ cells during this disease. Mononuclear cells were isolated from spinal cords of WT and PPAR-Δ⁻/⁻ mice at 4 d after the onset of clinical signs, and the production of IFN-γ and IL-17A by CD4⁺ cells was determined by intracellular cytokine staining. This analysis revealed the presence of a higher number (Fig. 2 a) and frequency (Fig. 2 b) of IFN-γ-producing CD4⁺ cells in spinal cords of PPAR-Δ⁻/⁻ versus WT mice. Although we did not detect a difference in the number or frequency of IL-17A-producing cells between these two groups (Fig. 2, a and b), we did observe a trend for higher amounts of CD4⁺ cells that coproduced both IFN-γ and IL-17A in PPAR-Δ⁻/⁻ spinal cords (629 ± 167 in PPAR-Δ⁻/⁻ vs. 181 ± 49 in WT, P = 0.06, n = 3 experiments; Fig. 2 c). Recent studies suggest that this Th subset is expanded by IL-23 and is highly pathogenic in the acute phase of EAE (Axell et al., 2006; Kebir et al., 2009).

Next, we investigated whether MOG p35-55–reactive Th1 cells also underwent preferential expansion in the periphery of PPAR-Δ⁻/⁻ mice during EAE. We harvested spleens from WT and PPAR-Δ⁻/⁻ mice at 8 d after immunization and examined the recall proliferation and cytokine production in response to MOG p35-55 stimulation. Consistent with spinal cord findings, we found that splenocytes from PPAR-Δ⁻/⁻ mice produced higher levels of IFN-γ, IL-17A, and TNF in response to peptide stimulation as compared with WT counterparts (Fig. 2 d). We also detected higher frequency of IFN-γ⁺ IL-17A⁺ CD4⁺ cells in PPAR-Δ⁻/⁻ versus WT splenocytes by intracellular cytokine analysis (6.6 ± 1.5 in PPAR-Δ⁻/⁻ vs. 3.0 ± 1.3 in WT, P = 0.02, n = 4 experiments). In addition, PPAR-Δ⁻/⁻ MOG p35-55–reactive splenocytes proliferated more robustly than WT cells (Fig. 2 e). In contrast, the serum titers of MOG p35-55–specific IgM or IgG were not different between WT and PPAR-Δ⁻/⁻ during EAE (Fig. 2 f). Thus, the more severe CNS inflammatory response in PPAR-Δ⁻/⁻ mice correlated more with the enhanced expansion and CNS infiltration of IFN-γ– and IL-17A–producing CD4⁺ cells.

**PPAR-Δ limits CD4⁺ cell expansion and effector cytokine production**

To determine whether the enhanced T cell autoimmunity in PPAR-Δ⁻/⁻ mice was a result of intrinsic differences in CD4⁺ cells, we contrasted the cytokine profiles of WT and PPAR-Δ⁻/⁻ naive (CD62L⁺) CD4⁺ cells in response to ex vivo stimulation
Figure 2. PPAR-δ−/− mice exhibit higher Th cytokine production during EAE. (a and b) Mononuclear cells were isolated from spinal cords of WT or PPAR-δ−/− mice at 4 d after the onset of clinical signs, and then they were pooled (n = 3/group) and processed for intracellular cytokine staining. The number (a) and frequency (b) of CD4+ mononuclear cells that had intracellular accumulation of IFN-γ (IFN-γ/IL-17A+ and IFN-γ/IL-17A−) or IL-17A (IL-17A+ IFN-γ− and IL-17A− IFN-γ−) are shown. (c) Representative FACs plot of IFN-γ and IL-17A staining in CD4+ gate. (d and e) Splenocytes from male age-matched WT or PPAR-δ−/− mice were harvested at 8 d after immunization and were stimulated ex vivo with MOG p35-55. Cytokines (d) were measured in culture supernatants using ELISA. Proliferation (48 h; e) was measured by [3H]-thymidine incorporation assay. Values are means + SEM of triplicate culture wells. Data in a–e are representative of three experiments. (f) Serum was collected from WT and PPAR-δ−/− mice during EAE. Titers of MOG p35-55–specific IgM and total IgG were determined by ELISA. Values are absorbance measurements of individual mice from two experiments that contained n = 5 and n = 11 mice/group. *, significantly different from WT.
Figure 3. **PPAR-δ limits cytokine production by CD4+ cells.** (a–c) Naive CD4+ cells were isolated from the spleens of WT or PPAR-δ−/− mice and stimulated with anti-CD3 and anti-CD28 under Th0-skewing (a, c, e, and g) or Th17-skewing (b, d, f, and h) conditions. After 4 d of culture, CD4+ cells were processed for intracellular cytokine staining. (a and b) Representative FACs plots of events in the CD4+ gate. (c and d) Cytoxine levels in culture supernatants after 48 h of culture under Th0-skewing (c) or Th17-skewing (d) conditions. (e–h) mRNA levels of IFN-γ and T-bet (Th0 conditions) or IL-17A and RORγt (Th17 conditions; relative to β-actin mRNAs) were determined using real-time PCR. *, significantly different from WT. Values are means ± SEM of triplicate cultures (c and d) or triplicate reactions (e–h). Data are representative of two to five experiments.

Figure 3. PPAR-δ limits cytokine production by CD4+ cells. Naive CD4+ (a) or CD19+ (b) cells were isolated from the spleens of WT or PPAR-δ−/− mice and were stimulated as indicated. Proliferation of these cells was measured using a [3H]-thymidine incorporation assay. Values are means ± SEM of triplicate cultures. (c and d) Splenocytes were harvested cells were isolated from the spinal cords of PPAR-δ−/− mice reconstituted with the PPAR-δ−/−CD4+ cells exhibited higher IFN-γ and IL-17A production as compared with those transferred with WT CD4+ cells (Fig. S1 a). Although only a small fraction (~25%) of mice in both groups succumbed to EAE, more WT B6.Rag1−/− than PPAR-δ−/−B6.Rag1−/− mice developed a relapsing-remitting course (four of four of WT vs. one of four PPAR-δ−/−B6.Rag1−/−; Fig. S1 b). Although only a small fraction (~25%) of mice in both groups succumbed to EAE, more WT B6.Rag1−/− than PPAR-δ−/−B6.Rag1−/− mice developed a relapsing-remitting course (four of four of WT vs. one of four PPAR-δ−/−B6.Rag1−/−; Fig. S1 b). These results indicate that PPAR-δ acts in the CD4+ cell compartment to limit expansion of Th1 and Th17 cells and to dampen CNS inflammation during EAE.

**PPAR-δ inhibits expression of IL-12 family cytokines**

Both IFN-γ/IL-17A and IFN-γ/IL-17A+ CD4+ cells were detected in spinal cords of PPAR-δ−/− mice during EAE. The former Th subset is induced by IL-12, whereas the latter subset is expanded by IL-23 (Awasthi et al., 2009; McGeachy et al., 2009). Given results that the related nuclear receptor PPAR-γ negatively regulates IL-12p40 (common to IL-12 and IL-23; Storer et al., 2005), we investigated whether myeloid cells from PPAR-δ−/− mice have a higher propensity to produce this or other IL-12 family cytokines. Indeed, we found that splenic cd11b+ cells and peritoneal macrophages from PPAR-δ−/− mice produced higher levels of IL-12p40 as compared with WT counterparts (Fig. 5, a and b). A small amount of IL-12p70 (p35/p40 heterodimer) was also detected in supernatants of PPAR-δ−/− cells at levels that were higher than those observed in WT. In contrast, we did not detect a consistent difference in the production of other cytokines between these groups. Similar to ELISA findings, we also observed that the mRNA expression of the IL-12 family members IL-12p40, IL-23p19, and IL-12p35 were markedly elevated in PPAR-δ−/− versus WT splenic cd11b+ cells after LPS stimulation (Fig. 5 c). Because myeloid cells drive the expansion of Th cells in the CNS during EAE (Bailey et al., 2007),
This finding that PPAR-δ⁻/⁻ macrophages have a higher propensity to produce IL-12 family cytokines may further explain the preferential accumulation of INF-γ/IL-17A⁻ and INF-γ/IL-17A⁺ CD4⁺ cells in the CNS of PPAR-δ⁻/⁻ mice during this disease.

PPAR-δ suppresses INF-γ and IL-12 production in a ligand-dependent manner

PPAR-δ is reported to have both ligand-dependent and -independent actions in vivo (Straus and Glass, 2007). To test how the ligand dependency of PPAR-δ affects Th cell production, we stimulated WT or PPAR-δ⁻/⁻ CD4⁺ T cells with anti-CD3 and anti-CD28 in the presence or absence of the specific PPAR-δ agonist GW0742. We used up to 1 µM of this drug because preliminary experiments indicated that GW0742 has off-target effects at ≥1-µM concentrations. These initial experiments revealed that GW0742 triggered an increase in Th1 cytokine production by WT CD4⁺ cells (Fig. 6 a). These results were unexpected given our results in PPAR-δ⁻/⁻ mice; however, they were consistent with previous results of GW0742 on Th1 cytokine production during EAE (Polak et al., 2005).

Because bovine serum is a potential source of several endogenous PPAR-δ ligands, we postulated that the failure of GW0742 to dampen Th1 cytokine production was because PPAR-δ was already saturated with a serum-derived ligand. To test this possibility, we repeated the experiment using serum-free X-vivo 20 media. Under these conditions, GW0742 had the expected effect of decreasing the production of INF-γ by WT but not by PPAR-δ⁻/⁻ CD4⁺ cells (Fig. 6, b–d).

Figure 5. PPAR-δ negatively regulates expression of IL-12 family cytokines. Splenic CD11b⁺ cells (a and c) or peritoneal macrophages (b) were isolated from adult age-matched WT and PPAR-δ⁻/⁻ mice and were cultured with 10 ng/ml LPS. (a and b) Supernatants were collected at 24 h for cytokine measurement by ELISA. (c) Cells were harvested at 0, 1, 3, or 5 h after stimulation for collection of RNA for real-time PCR measurement of p40, p35, or p19 mRNA expression (relative to β-actin). Values represent mean ± SEM of triplicate cultures (a and b) or reactions (c). *, significantly different from WT. Results are representative of three experiments.

Figure 6. GW0742 suppresses INF-γ and IL-12 family cytokines in a PPAR-δ-dependent manner. CD4⁺ cells (a–e) or peritoneal macrophages (f) were stimulated in the presence of 100 nM GW0742 or vehicle of equal volume. Cells were cultured in complete RPMI (a) or X-vivo 20 serum-free media (b–f). CD4⁺ cells were stimulated with anti-CD3 under Th0-skewing (a–d) or Th17-skewing (e) conditions, whereas peritoneal macrophages were stimulated with LPS. Cytokine production was measured after 24 or 48 h of culture by ELISA and intracellular staining was conducted after 4 d of culture as in Fig. 3. Values represent means ± SEM of triplicate cultures. *, significantly different from WT. †, significantly different from vehicle counterpart. Results are representative of three experiments.
Similarly, GW0742 suppressed the production of IL-12p40 in a PPAR-δ-specific fashion by macrophages (Fig. 6 f). Interestingly, treatment with GW0742 did not lower the production of IL-17A by WT T cells as detected by ELISA (Fig. 6 e), indicating that the enhanced production of IL-17A by PPAR-δ⁻/⁻ CD4⁺ cells was likely related to general effects of PPAR-δ on T cell expansion.

Transcriptional regulation of IFN-γ is complex, as it is dictated by the interaction of functional genomic elements acting as far as 50 kb from this gene (Aune et al., 2009). Comparatively, the LPS-induced transcription of IL-12p40 is more straightforward, as it involves the binding of well-characterized transcription factors (i.e., NF-κB) to genomic elements in the proximal promoter of this gene (Murphy et al., 1995). We thus pursued studies of the IL-12p40 promoter to gain insight into the mechanism of ligand inhibition of cytokine production. We observed that GW0742 repressed IL-12p40 promoter activity in RAW264.7 cells that were transfected with an IL-12p40/luciferase reporter construct (Fig. S2). This ligand effect was lost and PPAR-δ dependent (Fig. S2). This result indicates that PPAR-δ works via a mechanism of transcriptional repression to regulate IL-12p40 gene expression.

PPAR-δ inhibits the production of IFN-γ and IL-12 family cytokines by PBMCs

Given that GW0742 is a ligand agonist of both mouse and human PPAR-δ, we investigated whether this drug modulates cytokine production by human immune cells. We obtained PBMCs from three male healthy donors and then cultured these cells in serum-free media containing either 100 nM GW0742 or vehicle control in the presence of either anti-CD3 and anti-CD28 or LPS. Similar to findings in the mouse system, we found that 100 nM GW0742 selectively reduced the production of IFN-γ by PBMCs in response to anti-CD3 and anti-CD28, and it blunted the LPS induction of IL-12p40 and IL-23 (Table II). These effects were observed in three of three donors.

Concluding remarks

This study provides new evidence that PPAR-δ is an important negative regulator of CNS autoimmune inflammation. We found that mice deficient in this nuclear receptor exhibit a severe course of EAE characterized by a preferential accumulation of IFN-γ⁺IL-17A⁻ and IFN-γ⁺IL-17A⁺ CD4⁺ subsets in the CNS. This appeared to occur as a result of several immune aberrations that included a hyperresponsive CD4⁺ T cell population and higher IL-12 family cytokine production by myeloid cells. We also found that the PPAR-δ ligand GW0742 inhibits the production of IFN-γ and IL-12p40 by both mouse and human immune cells, indicating that these immune regulatory effects of PPAR-δ are ligand dependent and conserved between mouse and man. Taken in this light, our observation of immune phenotypic differences between WT and PPAR-δ⁻/⁻ mice, in the absence of treatment with synthetic ligands, suggests that PPAR-δ likely binds an endogenous ligand to mediate its anti-inflammatory effects in vivo. Future studies will investigate the nature of these ligands as well as the molecular mechanism of inhibition of IFN-γ and IL-12 family cytokines by PPAR-δ.

MATERIALS AND METHODS

Mice and EAE induction. Mice that have targeted disruption of the last exon of PPAR-β/δ gene containing the ligand-binding and RXR heterodimerization domains have been described previously (Peters et al., 2000). Homozygote mice express a truncated unstable PPAR-δ transcript, resulting in no detectable PPAR-δ protein (Peters et al., 2000). These mice, herein referred to as PPAR-δ⁻/⁻ mice, were originally on an outbred (129 × B6) background and were crossed six generations to C57BL/6. PPAR-δ⁻/⁻ mice, in the absence of treatment with synthetic ligands, suggests that PPAR-δ⁻/⁻ mice could not be generated past the sixth generation. Thus, WT and mutant colonies were established from the sixth generation. Genotyping of PPAR-δ⁻/⁻ mice and WT counterparts was performed by amplifying a portion of the ligand-binding domain of PPAR-δ or the neo cassette insert of the mutant allele from tail DNA (cyling conditions: 1 min at 94°C; 1 min at 60°C, and 1 min at 72°C). Primer sequences were: PPAR-δ forward, 5’-CAGGATGTCCTTCCACAGAGACAG-3’; and neo cassette, 5’-GCAATCATCATCTGGG-3’. EAE was induced in PPAR-δ⁻/⁻ mice (8–12 wk of age) and age-matched WT counterparts via subcutaneous immunization with 100 μg MOG p35-55 in an emulsion with CFA (Dunn et al., 2007). Mice were also

| Stimulus | Donor 1 | Donor 2 | Donor 3 |
|----------|---------|---------|---------|
|          | Vehicle | GW0742  | Vehicle | GW0742  | Vehicle | GW0742  |
| α-CD3 and α-CD28 |         |         |         |         |         |         |
| IL-2     | 122 (7) | 160 (8)³ | 178 (5) | 200 (11) | 92 (11) | 93 (4)  |
| IL-17A   | 21 (2)  | 25 (1)  | 14 (1)  | 16 (5)   | 16 (1)  | 14 (1)  |
| TNF      | 140 (12)| 113 (13)| 491 (17)| 400 (88) | 1,019 (50)| 1,178 (50)|
| IFN-γ    | 1,231 (213)| 716 (33)³ | 4,205 (129)| 3,125 (271)³ | 3,087 (111)| 2,547 (97)³ |
| LPS      |         |         |         |         |         |         |
| IL-12p40 | 7,160 (152) | 4,541 (162)³ | 5,497 (264) | 3,203 (328)³ | 8,881 (367) | 5,562 (318)³ |
| IL-23    | 3,581 (54) | 2,507 (109)³ | 2,262 (41) | 1,450 (50)³ | 3,775 (136)| 2,508 (124)³ |
| IL-12p70 | 322 (38) | 275 (36) | 1,486 (145) | 991 (85)³ | 1,195 (187) | 1,335 (157) |

Values are means ± SEM of cytokine levels (pg/ml) in triplicate cultures.
³Significant difference from vehicle control cultures using a Student’s t test (P = 0.05, two-tailed).

Table II. Cytokine production by human PBMCs treated with 100 nM GW0742

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injected intravenously with 50 ng Bordetella pertussis toxin (BD) at the time of, and 2 d after, immunization. Mice (n = 6–10/group) were examined daily for clinical signs of EAE and were scored on a five point scale: 0, no clinical disease; 1, limp tail; 2, hindlimb weakness; 3, complete hindlimb paralysis; 4, hindlimb paralysis plus some forelimb paralysis; and 5, moribund or dead. We found that clinical phenotype of PPAR-δ−/− mice was pronounced in males. Thus, we conducted all subsequent experiments in male mice. Animal protocols were approved by the Division of Comparative Medicine at Stanford University and animals were maintained in accordance with the guidelines of the National Institutes of Health.

MOG p35–55–specific Th responses. At 8 d after immunization (in EAE experiments), spleens were recovered from nonsick immunized mice (n = 3 mice/group) and were dissociated into a single cell suspension. Red blood cells were lysed and splenocytes were cultured in 96-well plates (0.5 × 10^5 cells/well) with 0–25 µg/ml MOG p35–55 in complete RPMI (RPMI 1640 supplemented with 2 mM L-glutamine, 1 mM sodium pyruvate, 0.1 mM nonessential amino acids, 100 U/ml penicillin, 0.1 mg/ml streptomycin, 0.5 µM 2-mercaptoethanol, and 10% fetal calf serum). Culture supernatants were harvested at 24–72 h after stimulation for cytokine measurement by ELISA (BD, R&D Systems, and eBioscience kits).

Rag transfer experiments. CD4+ cells were isolated from spleens of male PPAR-δ−/− mice (n = 10 mice/group), pooled together, and 7–10 × 10^6 of these cells were injected intravenously into B6.Rag1−/− recipients (The Jackson Laboratory). B6.Rag1−/− transferred with WT CD4+ cells served as controls. EAE was induced in these mice at 48 h after transfer as described previously (Dunn et al., 2007). Approximately 25% of mice in each group developed clinical signs. Nonsick mice were used to investigate the role for T cell–expressed PPAR-γ in the regulation of MOG p35–55 splenocyte responses.

MOG p35–55–specific IgG and IgM responses. Mice were bled at day 26 of disease. MOG p35–55–specific antibodies were detected in the sera of mice using an ELISA assay. In brief, plates were coated overnight at 4°C with 10 µg/ml MOG p35–55 peptide and then blocked with 3% fetal calf serum before serial probing with sera (1:100 diluted, 2 h), followed by a 1-h incubation with horseradish peroxidase–conjugated goat anti–mouse IgG or IgM (SouthernBiotech). Plates were washed three times with 0.5% PBS containing Tewen 20 between each step. Horseradish peroxidase conjugates were detected using tetramethylbenzidine substrate. Absorbances were measured at 450 nm and background levels were subtracted.

Histopathology. Brains and spinal cords were dissected from male mice (n = 5/group), fixed in 10% formalin in PBS, and embedded in a single paraffin block. 6–10-µm-thick sections were stained with hematoxylin and eosin and luxol fast blue. The number of meningeal and parenchymal inflammatory foci was counted in brain and spinal cord sections by a trained neuropathologist who was blinded to the clinical and treatment status of the animal. A demyelination/vacuolation score was also assigned to each specimen that ranged in severity from 0 (absence of demyelination/vacuolation) to 4 (severe demyelination/vacuolation).

Isolation of CNS mononuclear cells. CNS mononuclear cells were isolated from mice (n = 3 mice/group) at 5 d after the onset of EAE. Mice were anesthetized (80–100 mg/kg ketamine hydrochloride: 10 mg/kg xylazine) and perfused with PBS containing 5 U/ml heparin. Spinal cords were pooled together (n = 3), scissor minced into Hank’s buffered saline, and then digested for 45 min at 37°C with 300 U/ml Clostridial collagenase type IV (Roche) and 1 mg/ml DNase (Sigma-Aldrich). After digestion, the spinal cord tissue was dissociated through a 70-µm sieve. This filtrate was centrifuged and the pellet containing the mononuclear cells was resuspended in 30% Percoll (GE Healthcare). This suspension was underlaid with 70% Percoll and centrifuged at 1,800 rpm for 10 min. Mononuclear cells were collected from the 30%/70% interface, washed twice, and suspended in complete RPMI.

FACS staining of cell subsets in the spleen and CNS. FACs staining of splenocyte and CNS mononuclear cells was done using BD antibodies and protocols unless otherwise noted. Antibodies directed against the following mouse cell surface antigens were used: CD4, CD8, CD19, CD11b, CD11c, CD44, CD45, CD62L, and F4/80 antigen (AbD Serotec). Intracellular staining of mouse FoxP3 was performed using a kit from ebioscience. For intracellular cytokine staining, cells were stimulated for 4 h with 10 ng/ml PMA and 750 nM monomycin in the presence of GolgStop to cause intracellular accumulation of cytokines. Cells were then stained with anti–CD4 and fixable aqua live/dead dye (Invitrogen) before fixation, permeabilization, and intracellular staining for mouse anti–IFN-γ and IL-17A. Data were acquired using a FALzer II cell sorter (BD) or LSRII analyzer and the frequency of CD45+/CD4+ cells expressing IFN-γ and IL-17 was analyzed using FlowJo software (Tree Star, Inc.).

Culture of isolated myeloid and Th cell populations. CD11b+ or CD19+ cells were isolated by positive selection and CD4+ cells by negative selection using reagents (Miltenyi Biotec). The naive CD4+ fraction was further fractionated using CD62L+ magnetic beads provided in the naive mouse CD4+ isolation kit (Miltenyi Biotec). Percutaneous macrophages were harvested from the peritoneum of mice at 3 d after i.p. injection with thioglycollate. These cells were washed, plated in flat-bottomed plates (2 × 10^5/well), and allowed to rest overnight (with vehicle or GW0742) before stimulation with LPS. All cells were resuspended in complete RPMI or in X-vivo 20 serum-free media (Lonza; serum-free media also contained 2 mM L-glutamine, 1 mM sodium pyruvate, 0.1 mM nonessential amino acids, 100 U/ml penicillin, 0.1 mg/ml streptomycin, and 0.5 µM 2-mercaptoethanol). T cells were cultured with 0.1–0.5 µg/ml of plate-bound anti-CD3 (clone 145-2C11; BD) and α–CD28 (clone 37.51; BD) under Th0–skewing conditions (no added cytokines) or Th17–skewing conditions (20 ng/ml of mouse recombinant IL-6, 3 ng/ml of human recombinant TGF-β, and 10 µg/ml of anti–IFN-γ (XMG1.2). Percutaneous macrophages or splenic cd11b+ cells were cultured in flat-bottomed 96-well plates with 1–100 ng/ml LPS. Cytokines were measured in the supernatants of cultured cells at 24–72 h using commercial ELISA kits.

Proliferation. Proliferation rate of T cells and B cells was measured as described previously (Dunn et al., 2007).

Real-time PCR. RNA was extracted from naive CD4+ T cells immediately after isolation or after 24 h of culture using Absolutely RNA kits (Agilent Technologies). Real-time PCR was conducted using the standard curve method, using SYBR Green reagents (Qiagen) and the following specific primer sets: IL-12p40 forward, 5′-TCCCCATTCTCTACTTCTCCCTCCT-3′; IL-12p40 reverse, 5′-GGAGGCGCATTCTTCTGTTTACA-3′; IL-12p35 forward, 5′-CACGCTAACCTCTCTTATTGGT-3′; IL-12p35 reverse, 5′-CAGCAGTCAGCAGAATAGTGT-3′; IL-23p19 forward, 5′-AATCTCTGTCATGTTAGCCCGTGG-3′; IL-23p19 reverse, 5′-GATTCTATATGTCCTCCGCCTGGT-3′; T-bet forward, 5′-AATGCGCTGTAATTGGT-3′; T-bet reverse, 5′-CCTGCCAGTTGATTCGCTCCG-3′; RORγT forward, 5′-AGATGCTTCTFACAGCAG-3′; RORγT reverse, 5′-GGCTTGACCAACGACA-3′; β-actin forward, 5′-GTGCTCTTATGCTTCTGTGT-3′; and β-actin reverse 5′-ACACGGCAATTTGCATTGGT-3′. The luciferase reporter DM-703, in which luciferase transcription was transfected using Lipofectamine 2000 (Invitrogen) and were cultured in commercial ELISA kits.

IL-12p40 promoter analysis. The RAW264.7 mouse macrophage cell line was maintained in RPMI1640 medium supplemented with 10% fetal bovine serum, 100 U/ml penicillin, and 100 µg/ml streptomycin. Cells were transfected using Lipofectamine 2000 (Invitrogen) and were cultured in 6-well plates in low-serum (0.5% FBS) media containing penicillin and streptomycin. The luciferase reporter DM-703, in which luciferase transcription is directed by a mouse IL-12p40 promoter fragment extending to −703, was obtained from K.M. Murphy (Washington University School of Medicine, Medicine at Stanford University and animals were maintained in accordance with the guidelines of the National Institutes of Health.
Culture of human PBMCs. PBMCs were isolated from peripheral blood of three healthy donors using Ficoll gradient. These cells were resuspended in X-vivo 15 media containing 2 mM l-glutamine, 1 mM sodium pyruvate, 0.1 mM nonessential amino acids, 100 U/ml penicillin, 0.1 mg/ml streptomycin, and 0.5 µM 2-mercaptoethanol. Either 100 nM of GW0742 or an equal volume of vehicle was added to cells, and they were plated (2 × 10^4 well) in flat-bottomed plates. After 2 h of vehicle or ligand treatment, Dynabead human T cell expander (1 Dynabead for every 10 T cells; Invitrogen) or 10 ng/ml LPS was added to cells. Cytokines were measured in culture supernatants at 48 or 72 h using IFN-γ, IL-2, TNF, and IL-17A ELISA kits (eBioscience) and an IL-12p40 kit (BD).

Statistical analysis. Data are presented as means ± SEM. When data were parametric (kurtosis and skewness < 2), a Student’s t test (two-tailed) was used to detect between-group differences. When data were nonparametric, ranks were compared among groups using a Mann-Whitney U test. A value of P ≤ 0.05 was considered significant.

Online supplemental material. Fig. S1 depicts the results from EAE studies using Foxp3gfp reporter mice transferred with WT or PPAR-β/δ CD4+ cells. Fig. S2 depicts the results from the IL-12p40 luciferase reporter experiment. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20091663/DC1.

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