Introduction

Autoimmune diseases such as multiple sclerosis (MS), rheumatoid arthritis (RA), and Grave’s disease preferentially affect women (1) and likely involve an oligoclonal expansion of autopathogenic Th1 cells directed at tissue-specific antigens (2). In general, women have a stronger immune response than men (3), perhaps contributing not only to a more effective immune surveillance and increased longevity, but also to a propensity for developing autoimmune diseases. Gender differences in the immune response and susceptibility to autoimmune diseases usually become apparent after sexual maturity (4). Sex hormones may contribute to susceptibility or resistance by influencing the development and function of pathogenic T cells, but conceivably, these hormones could also affect regulatory T cells that modify the course of disease.

Increased levels of sex hormones produced during pregnancy have been associated with clinical remissions in MS and RA patients, whereas reduced levels postpartum lead to an increased number of exacerbations (5–7). Similarly, pregnant animals develop less severe signs of experimental autoimmune encephalomyelitis (EAE) (8, 9), and estrogen, administered at levels found during pregnancy (100-fold higher than basal levels), can suppress EAE (10, 11). Conversely, castration of female mice accelerates the onset of EAE (11). These observations suggest that whereas endogenous estrous and diestrous levels of estrogen may exert a limited inhibitory effect, higher levels of supplemental estrogen are needed to suppress development of inflammatory Th1 responses (12).

Estrogen receptors (ER) have been demonstrated on human blood mononuclear cells, splenic and thymic cells (13), as well as HLA-DR negative human peripheral and thymic T cells (14), with some further indication that, within human T-cell subtypes ER expression might be limited to CD8+ cells (15, 16). However, estrogen has been shown to directly influence the cytokine secretion profile of human CD4+ T-cell clones (17, 18), strongly implicating functional ER, and we recently demonstrated ERα message in purified CD4+ T cells from mouse spleen (19). Moreover, Benten et al. (20) described a novel estradiol signaling pathway, present in both CD4+ and CD8+ T cells, which was mediated through plasma membrane receptors rather than the classical nuclear estrogen receptors. Taken together, these findings provide strong evidence that estrogen can bind to and regulate CD4+ T cells, thus allowing the possibility that estrogen might dampen inflammation either by directly inhibiting or deviating Th1 cells or by promoting development of regulatory Th2 (21) or CD8+ (16) cells.

EAE is an inflammatory demyelinating disease of the central nervous system (CNS) that is induced by immunization of susceptible mice with myelin proteins or peptides in complete Freund’s adjuvant (CFA) (22–25). Autoreactive CD4+ Th1 cells home specifically to the CNS and initiate the accumulation of inflammatory mononuclear cells, resulting in demyelination and clin-
ical disease (26). As in MS, female mice develop more severe EAE than male mice (27, 28). We developed a new EAE treatment model using recombinant BV8S2 proteins to induce regulatory T cells in male transgenic mice on the B10.PL background expressing the β chain of the T-cell receptor (TCR) for myelin basic protein (MBP) Ac1-11 (29). T cells in unprimed mice proliferated vigorously to Ac1-11, but produced low levels of IFN-γ and IL-10, indicating that these were naïve effector precursor cells. The enhanced T-cell response to Ac1-11 encoded by the transgene induced a cognate T-cell response to the TCR BV8S2 protein expressed by the MBP-specific T cells, suggesting natural priming in vivo. Induction of EAE by immunization with MBP-Ac1-11/CFA prompted MBP-specific T cells to greatly increase production of IFN-γ, provoking a concomitant increase of cytokine secretion by TCR-specific T cells. However, the mice developed relapsing EAE, indicating that the naturally induced anti-TCR T cells were not sufficient to inhibit disease. Vaccination with BV8S2 protein in incomplete Freund’s adjuvant (IFA) before EAE induction reduced incidence and severity of disease and stimulated TCR-specific T cells to secrete elevated levels of both IFN-γ and IL-10 (29) These regulatory T cells expressed the same Tg BV8S2 chain and the same AV2S3 chain as the encephalitogenic T cells, differing only in the AV-CDR3 junctional region (30), and could inhibit activation and transfer of EAE after coculture with the MBP-Ac1-11-specific T cells (29) through release of IL-10, IL-4, and IFN-γ, but not TGF-β (31). These data demonstrate that immune regulation in this model occurs through a nondeletional cytokine-driven suppressive mechanism involving TCR-specific T cells (32). We now show that, in contrast to males, female littermate mice developed more severe EAE and were only transiently protected from EAE after vaccination with BV8S2 protein. To clarify these gender differences in susceptibility and treatment in EAE, we further evaluated the combined effects of TCR vaccination and estrogen on both pathogenic and regulatory T cells.

Methods

Mice. Tg mice bearing the functionally rearranged BV8S2 gene specific for MBP-Ac1-11 on the B10.PL background were kindly provided by Joan Goverman (Seattle, Washington, USA). Male Tg mice were bred with B10.PL females, and the offspring were tested for expression of the transgene by FACS analysis of blood cells stained for BV8S2 as described previously (33). Approximately half of each litter expressed the BV8S2 transgene, with approximately half of these transgenic littersmates of each sex. For some experiments, mice expressing the BV8S2 transgene were compared with littermates that did not express the transgene. The colony was housed and cared for at the Animal Resource Facility (Veterans Affairs Medical Center, Portland, Oregon, USA) according to institutional guidelines.

Antigens. N-acetylated MBP-1-11 peptide (Ac-ASQKRP-SQRSK) was synthesized using solid-phase techniques and was purified by HPLC at the Beckman Institute, Stanford University (Stanford, California, USA). Glutathione S-transferase (GST) and GST-BV8S2 proteins were expressed and purified as described previously (34). The GST-BV8S2 fusion protein contains the complete BV, BD, and BJ regions, and the first 19 residues of the BC region from the TCR of an encephalitogenic rat T-cell clone fused to the COOH-terminal end of GST. To control for the GST-BV8S2 protein, the GST protein was produced and purified using the same expression system. The GST protein was included as a control in all tissue culture experiments using the GST-BV8S2 protein.

Induction of active EAE and protection with BV8S2 protein or estrogen therapy. EAE was induced in Tg male or female mice by injecting 400 µg MBP-Ac1-11/CFA containing 200 µg Mycobacterium tuberculosis subcutaneously over four sites on the flank. For TCR protection experiments, mice were injected with 12.5 µg recombinant BV8S2 protein/IFA (experimental), or saline/IFA, or GST/IFA (sham controls), intraperitoneally, on days −7 and +3 relative to injection of the MBP-Ac1-11, according to the protocol developed by Kumar and Sercarz (35). In an alternative protocol, mice were given the initial two injections and then boosted weekly with 12.5 µg BV8S2 protein or saline given subcutaneously. Groups of male and female mice that were treated with TCR protein (Figure 1) were littersmates, although data from the male mice have been reported previously (29). For estrogen hormone therapy or combined estrogen plus TCR therapy, 3-mm pellets containing varying amounts of 17β-estradiol or estriol (Innovative Research of America, Sarasota, Florida, USA) were implanted subcutaneously on the animal’s back 7 days before induction of EAE. Control mice were sham operated but received no pellet. The estrogen pellets provide continuous controlled release of a constant level of hormone over a period of 60 days. The concentration of 17β-estradiol in pellets used in our experiments and the expected serum concentration of secreted hormone maintained in the mice are listed in Table 1, along with the established range of physiological serum hormone levels during the estrus cycle and pregnancy (36). Serum concentrations of estrogen monitored before and during the course of EAE in representative control and implanted mice consistently fell within the expected ranges.

Mice were assessed daily for clinical signs of EAE according to the following scale: 0, no signs; 1, limp tail;
2, moderate hind-limb weakness (waddling gait); 3, moderately severe hind-limb weakness; 4, severe hind-limb weakness; 5, paraplegia; 6, quadriplegia, moribund condition. The cumulative disease index (CDI) was determined for each mouse by summing the daily clinical scores, and the mean CDI plus or minus SEM was calculated for the control and experimental groups. The mean clinical score (MCS) was calculated for each mouse by dividing the CDI by the duration (days) of disease, and the mean plus or minus SEM was calculated for the control and experimental groups.

**Proliferation assay.** Spleens (SPL) were removed surgically, and single-cell suspensions were prepared. Proliferative responses of T cells were determined in 96-well microtiter plates by incubating 4·10^5 spleen cells plus antigen at an optimal concentration of 20 μg/well. Cultures were incubated for 72 hours at 37°C and 7% CO2, the last 18 hours in the presence of 0.5 μCi 3H-thymidine. Cells were harvested onto glass fiber filters, and thymidine uptake was determined by liquid scintillation. Mean cpm ± SEM were calculated from triplicate wells. The stimulation index (SI) was obtained by dividing cpm from antigen-stimulated wells by cpm from wells with no antigen. SI in cultures stimulated with GST alone was subtracted from the SI induced with GST-BV8S2 protein.

**Measurement of cytokine secretion.** Spleen cells were suspended at 4·10^6 cells/mL in stimulation medium with and without specific antigens. Cell culture supernatants were recovered at 72 hours and frozen at –70°C until needed for the cytokine assay. Measurement of cytokines was performed by ELISA developed in our laboratory (19), using cytokine-specific capture and detection antibodies (PharMingen, San Diego, California, USA). Capture antibodies for IFN-γ, IL-10, and TGF-β were diluted to 2 μg/mL in bicarbonate coating buffer (0.1 M NaHCO3, pH 8.2). Standard curves for each assay were generated using recombinant mouse cytokines (PharMingen), and the concentration of cytokines in the cell supernatants was determined by interpolation from the appropriate standard curve.

**Assessment of antibody responses.** Antibody reactivity to MBP-Ac1-11 peptide and GST-BV8S2 protein was determined by indirect ELISA as described previously (37). Briefly, mouse antisera from treated and control Tg mice

![Figure 1](https://example.com/figure1.png)

**Table 2**

<table>
<thead>
<tr>
<th>Group</th>
<th>Figure</th>
<th>Treatment</th>
<th>Incidence</th>
<th>Day of onset</th>
<th>CDI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Males</td>
<td>1a</td>
<td>Controls</td>
<td>13/16</td>
<td>15 ± 1</td>
<td>49 ± 10</td>
</tr>
<tr>
<td></td>
<td></td>
<td>BV8S2</td>
<td>6/16a</td>
<td>14 ± 1</td>
<td>17 ± 7a</td>
</tr>
<tr>
<td>Males</td>
<td>1b</td>
<td>Controls</td>
<td>10/10</td>
<td>13 ± 1</td>
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<td>BV8S2,B</td>
<td>1/9c</td>
<td>15</td>
<td>7 ± 7c</td>
</tr>
<tr>
<td>Females</td>
<td>1c</td>
<td>Controls</td>
<td>5/5</td>
<td>15 ± 4</td>
<td>75 ± 12</td>
</tr>
<tr>
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<td>6/6</td>
<td>13 ± 1</td>
<td>82 ± 23</td>
</tr>
<tr>
<td></td>
<td></td>
<td>GST</td>
<td>6/6</td>
<td>12 ± 1</td>
<td>36 ± 4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>BV8S2,B</td>
<td>7/7</td>
<td>19 ± 5b</td>
<td>10 ± 7b</td>
</tr>
</tbody>
</table>

CDI is sum of daily scores; 12.5 μg BV8S2/IFA on days –7 and +3. *P < 0.05. **P < 0.01. ***P < 0.001. B, weekly boosts with BV8S2/IFA.
Effects of estrogen on EAE. The effects of sex hormones, including 17β-estradiol and estril, were evaluated on the clinical course of EAE by hormone depletion or addition experiments. As is shown in Figure 2a, female Tg mice unable to produce detectable levels of estrogen (<1 pg/mL) or other sex hormones after ovariectomy developed significantly more severe EAE than sham-ovariectomized females (CDI = 81 vs. 56; P < 0.001). These data demonstrate that even basal levels of ovarian factors, possibly including estrogen, provide some regulation of EAE. On the other hand, treatment of sham- or nonovariectomized females with 17β-estradiol pellets produced a dose-dependent inhibition of EAE in both Tg females (Figure 2a) and B10.PL littermate females (Figure 2b). Notably, added estrogens had a less pronounced effect on the Tg versus non-Tg females. In B10.PL females, essentially complete inhibition of EAE was produced with 15-mg pellets secreting pregnancy levels of 17β-estradiol (9,000–10,000 pg/mL serum) over a 60-day period or with 2.5-mg pellets (1,500–2,000 pg/mL serum), and substantial inhibition was produced over a wide range of estrogen concentrations from estrus (0.36-mg pellets secreting 150–200 pg/mL serum) to diestrus levels (0.10-mg pellets secreting 25–50 pg/mL serum, Figure 2b). By comparison, in Tg females, pregnancy levels of hormone (15-mg pellets) produced a marked but incomplete inhibition of EAE, with estrus levels (0.36-mg pellets) producing only moderate inhibition (Figure 2a). Estril, which is elevated only during pregnancy, had an inhibitory effect on EAE in B10.PL females equivalent to 17β-estradiol (Figure 2b). The partial resistance to estrogen therapy in Tg females may be due to the higher native frequency of Ac1-11-specific T cells afforded by the transgene (29), which with EAE were incubated in antigen-coated wells, and bound antibody was detected spectrophotometrically with peroxidase-labeled rabbit anti-mouse antibody and o-phenylene-diamine as a substrate. Differences between groups were determined using Student’s t test.

Ovariectomy. The ovaries were removed by making two bilateral incisions (5 mm) halfway between the base of the tail and the middle of the back, followed by small incisions (2.5 mm) through the peritoneal wall. The ovaries were pulled through the incisions by grasping the periovarian fat, the blood vessels were ligated, and the ovaries were removed. The incision was closed by surgical skin clips. The animals were allowed to recover for at least 1 week before initiation of experiments.

Androgen and estrogen detection. Mice were bled by cardiac puncture and the blood was allowed to clot at 4°C overnight. The samples were centrifuged, and the sera were collected and stored at –80°C until hormone analy-

Figure 2
Inhibitory effects on EAE of 17β-estradiol and estril on intact and ovariectomized female mice. Hormone pellets releasing controlled levels of estrogen (see Table 1 for the range of serum concentrations maintained by various pellet sizes) were surgically implanted 7 days before induction of EAE with MBP-Ac1-11 peptide/CFA + pertussigen. (a) Intact and ovariectomized Tg females, as indicated. (b) Intact (nonovariectomized) B10.PL female littermates. Note increased severity of EAE in ovariectomized versus intact Tg females, in intact Tg versus B10.PL females, and dose-dependent inhibition of EAE in both Tg and B10.PL females, with reduced sensitivity in Tg females. A total of 30 sham-operated and 81 experimental B10.PL female mice, and 36 sham-operated and 56 experimental Tg female mice were used in a total of nine combined experiments, with total group sizes ranging from five to 33 mice. *Significant difference between control and experimental (P < 0.05). Ovx, ovariectomized.
likely accounts for an increased severity of EAE (CDI = 56 in Tg vs. 39 in non-Tg females; \( P < 0.001 \)). Consistent with this notion, estrogen treatment of ovariectomized female Tg mice reduced the severity of EAE to about the same level as sham-ovariectomized Tg mice treated with estrogen (Figure 2a). However, the inability of estrogen to fully inhibit EAE in Tg females at very high levels approximating pregnancy suggests that a portion of the encephalitogenic cascade is estrogen insensitive.

**Combined TCR and estrogen therapy.** Because TCR and estrogen therapy were both partially effective for preventing EAE in Tg females, we sought to compare directly the effects of single versus combined therapy. Consistent with the results shown above, Tg females vaccinated with BV8S2 protein (with weekly boosting) had delayed onset but eventually developed severe EAE, whereas mice treated with estrus levels of 17\( \beta \)-estradiol (0.36-mg pellets secreting 150–200 pg/mL serum over 60 days) had normal onset but generally less severe disease (Figure 3, Table 3). However, combined treatment with both TCR protein and estrus levels of 17\( \beta \)-estradiol produced almost complete protection against EAE (Figure 3), with only three of 16 mice developing very mild disease (Table 3). A similar degree of protection was provided in ovariectomized Tg females treated with BV8S2 protein and estrogen (Table 3), demonstrating that the enhanced therapeutic effect was dependent on 17\( \beta \)-estradiol, rather than a combination of estrogen and other gonadally produced sex hormones.

**Estrogen skew response to Ac1-11 and potentiates response to BV8S2 protein.** To investigate the mechanism(s) involved in individual and combined therapies, we evaluated proliferation and cytokine responses of immune T cells from naive and treated mice. As is shown in Figure 4a, BV8S2 naive Tg males and females had equivalent proliferation responses to Ac1-11 peptide, purified protein derivative (PPD), and ConA, but naive Tg females had a strikingly reduced reactivity to the BV8S2 protein. This finding suggests that Tg females have a diminished native capacity to regulate an encephalitogenic response.

During development of EAE, splenic T-cell responses to Ac1-11 peptide were characterized by moderate proliferation and production of TGF-\( \beta \) and essentially absent secretion of IL-10 (Figure 4b). Secretion of IFN-\( \gamma \) in response to Ac1-11 peptide was modest, reflecting preferential migration of inflammatory T cells to draining lymph nodes and the CNS as observed previously (29). Treatment with either BV8S2 protein or estrogen alone reduced proliferation and marginally affected cytokine responses to Ac1-11. However, combined treatment with both BV8S2 protein and estrogen markedly reduced proliferation and dramatically enhanced production of IL-10 and TGF-\( \beta \), but not IFN-\( \gamma \) in response to Ac1-11 peptide (Figure 4b). In contrast, splenic proliferation and IL-10 responses to BV8S2 protein were enhanced by both treatments individually and further potentiated with combination therapy, with no significant effects of treatment on IFN-\( \gamma \) and TGF-\( \beta \) secretion (Figure 4c). Additionally, combination therapy reduced circulating levels of Ac1-11–specific IgG2a antibody associated with Th1 response, with no effect on IgG1 response (Figure 4d).

**Discussion**

The results presented above demonstrate, we believe for the first time, the powerful effects of combined TCR plus estrogen therapy on EAE in Tg female mice that are only partially responsive to either treatment alone. Concomitant administration of TCR protein and supplemental estrogen not only potentiated IL-10 production by regulatory TCR-reactive T cells, but also induced robust secretion of IL-10 and TGF-\( \beta \) by MBP-Ac1-

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**Table 3**

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Incidence</th>
<th>Onset</th>
<th>Peak</th>
<th>Mortality</th>
<th>Average CDI (10–30)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>60/64</td>
<td>11.8 ± 1.1</td>
<td>4.7 ± 0.1</td>
<td>12/49</td>
<td>56.4 ± 7.0</td>
</tr>
<tr>
<td>BV8S2</td>
<td>45/49</td>
<td>17.4 ± 2.2^^</td>
<td>3.9 ± 1.1^^</td>
<td>1/49</td>
<td>32.0 ± 13.2^^</td>
</tr>
<tr>
<td>17( \beta )-estradiol</td>
<td>25/33</td>
<td>15.1 ± 2.1^^</td>
<td>3.6 ± 0.7^^</td>
<td>1/33</td>
<td>37.3 ± 6.2^^</td>
</tr>
<tr>
<td>BV8S2 + 17( \beta )-estradiol</td>
<td>3/16</td>
<td>18.8 ± 8.0^^</td>
<td>0.5 ± 0.4^^</td>
<td>1/16</td>
<td>3.1 ± 4.0^</td>
</tr>
<tr>
<td>Ovx</td>
<td>27/27</td>
<td>10.6 ± 0.9</td>
<td>5.3 ± 0.7</td>
<td>7/27</td>
<td>81.2 ± 17.3^^</td>
</tr>
<tr>
<td>BV8S2</td>
<td>23/26</td>
<td>18.0 ± 1.9</td>
<td>3.9 ± 1.3</td>
<td>4/26</td>
<td>40.1 ± 20.9^^</td>
</tr>
<tr>
<td>17( \beta )-estradiol</td>
<td>11/15</td>
<td>13.6 ± 1.5</td>
<td>3.5 ± 0.8</td>
<td>4/15</td>
<td>40.0 ± 22.1^^</td>
</tr>
<tr>
<td>BV8S2 + 17( \beta )-estradiol</td>
<td>3/16</td>
<td>20.8 ± 6.0</td>
<td>0.9 ± 0.9</td>
<td>0/16</td>
<td>7.4 ± 6.7^^^</td>
</tr>
</tbody>
</table>

Data are combined from a total of 10 separate experiments. \(^*\)Significant difference between control and experimental \(( P < 0.0001 \)). \(^\text{a}\)Significant difference between Ovx control and Ovx experimental \(( P < 0.0001 \)). \(^\text{b}\)Significant difference between Ovx and non-Ovx control \(( P < 0.001 \)). Ovx, ovariectomized.
11–specific T cells. This demonstration that supplemental estrogen given at estrus levels can enhance the efficacy of TCR-based therapy prompts us to speculate that estrogen might also augment other immunoregulatory strategies for human autoimmune diseases.

It is apparent from these studies that the underlying mechanisms involved in TCR-based regulation and estrogen therapy are highly complementary. Vaccination with TCR protein preferentially induces Th2-like cells that inhibit Th1-cell activation and transfer of EAE by secretion of regulatory cytokines, including IL-10 and IL-4, but not TGF-β (29, 31). Estrogen given at estrus levels also enhanced IL-10 production by TCR-reactive T cells, even in the absence of additional TCR protein. In addition, estrogen alone inhibited proliferation of Ac1-11–specific T cells, but had little effect on cytokine production. However, the combination of both approaches had an additive effect on BV8S2 reactive cells that promoted the highest levels of IL-10, as well as a synergistic effect on MBP-Ac1-11–specific T cells that triggered production of both IL-10 and TGF-β, but not IFN-γ. IL-10 has been shown to be a critical cytokine for regulating EAE in mice (39), and an important mechanism underlying our observations is that combined TCR plus estrogen therapy resulted in optimal IL-10 production by both TCR- and MBP-specific T cells.

It seems probable that the supplemental estrogen acted directly on both MBP-Ac1-11– and BV8S2-reactive T cells to modify their function. In addition to our recent observation that ERα is expressed on CD4+ splenocytes from mice (19), Benten et al. (20) demonstrated that estradiol can bind to the plasma membrane of all CD4+ (as well as all CD8+) T cells through receptors that are not the classical genomic ERs. Binding of estradiol stimulated Ca2+ influx through Ca2+ channels and release of Ca2+ from the endoplasmic reticulum, resulting in impaired signaling through the TCR. Thus, estrogen could conceivably act through either classical or nonclassical pathways to functionally modulate activation and cytokine profiles of both pathogenic and regulatory CD4+ T cells. The estrogen-enhanced production of anti-inflammatory lymphokines by the TCR-reactive T cells might have provided the milieu needed to cause immune deviation of MBP-Ac1-11–specific T cells and antibodies, as has been suggested previously (40–42). Autopathogenic T-cell specificities that have been skewed to produce Th2 cytokines, including IL-4, IL-10, and TGF-β, have been associated with a nonencephalitogenic phenotype (43), except under one specific but unusual circumstance (44). Moreover, the local secretion of these inhibitory lymphokines by deviated or genetically transduced pathogenic T cells could protect against EAE by suppression of bystander pathogenic T cells (45–48).

It is clear that basal levels of ovarian hormones exert a regulatory influence on EAE, since ovariectomized Tg females with no detectable estrogen or progesterone developed significantly more severe disease than sham-operated females. Estrogen treatment profoundly inhibited EAE in B10.PL females, but had a less pronounced effect in Tg females, perhaps owing to the higher frequency of MBP-reactive T cells afforded by the Ac1-11–reactive TCR β-chain transgene. Interestingly, treat-
ment of ovariectomized Tg females with 17β-estradiol produced a comparable degree of EAE inhibition as in sham-operated Tg females (Figure 2), indicating that this or closely related estrogen analogues contributed to natural regulation in intact females. However, in both cases, inhibition of EAE was incomplete, connoting a limited estrogen effect and an estrogen-insensitive component of the encephalitogenic cascade in Tg females.

Similarly, treatment with BV882 protein was ultimately ineffective at controlling EAE in Tg females, although this therapy did cause a significant delay in disease onset. The eventual onset of full symptoms of EAE in TCR-treated female Tg mice likewise indicated incomplete regulation in the absence of supplemental estrogen. The essentially complete protection against EAE observed with combined therapy was obviously estrogen dependent and clearly involved highly compatible dynamic interactions. Based on the well-described effects of estrogen for forming ligand-activated nuclear transcription factors that function as upstream promoters of affected genes (49), we would speculate that supplemental estrogen could act directly through the classical genomic ER to upregulate IL-10 production by TCR-reactive T cells. The enhanced levels of IL-10 might then more efficiently skew responses of the MBP-Ac1-11–reactive T cells to produce enhanced levels of IL-10 might then more efficiently skew responses of the MBP-Ac1-11–reactive T cells to produce enhanced levels of IL-10 production by TCR-reactive T cells. The enhanced levels of IL-10 might then more efficiently skew responses of the MBP-Ac1-11–reactive T cells to produce additional IL-10 and TGF-β. Estrogen, estroli, and 17β-estradiol at pregnancy levels have all been shown to enhance IL-10 as well as IFN-γ, but not IL-4 or TGF-β, secretion by CD4+ proteolipid protein–specific (PLP-specific) T cells from MS and control patients (17, 18). In our study, 17β-estradiol alone did not significantly alter IL-10 or TGF-β production by MBP-AbI-11–specific T cells when given at relatively low estrus level doses in the absence of TCR protein. The possible promoter effect of estrogen on IL-10 production by both TCR and MBP-reactive T cells is currently under investigation.

The profound protective effect of combined TCR vaccination and estrogen therapy observed in the current study has important implications for treatment of human autoimmune diseases. Many of these conditions preferentially affect women and probably involve tissue antigen–specific inflammatory T cells that can be targeted by TCR vaccines. TCR-based immunotherapy has shown a possible clinical benefit in patients with MS (50) and RA (51), but the treatment affects only about half the patients, and clinical changes are, for the most part, moderate. Clinical trials using pregnancy levels of estradiol to treat females with MS are in progress (R. Voskuhl, personal communication). However, supplemental estrogen or estradiol alone may also not fully regulate pathogenic T cells in MS, even if given at pregnancy doses, because of enhanced resistance to regulation (52). Our data suggest that a much lower estrus dose of 17β-estradiol could have both a potentiating effect on regulatory TCR-reactive T cells and a synergistic effect on MBP-reactive T cells for producing anti-inflammatory cytokines when used in combination with TCR protein vaccination. Similarly, lower doses of estrogen may be highly compatible and complementary with other forms of immune therapy that are aimed at inducing immune deviation, including altered peptide ligands (53) and low-dose oral tolerance (54).

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