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Estriol Generates Tolerogenic Dendritic Cells In Vivo That Protect against Autoimmunity

Tracey L. Papenfuss,* Nicole D. Powell,† Melanie A. McClain,‡ Ashley Bedarf,* Amber Singh,* Ingrid E. Gienapp,‡ Todd Shawler,‡ and Caroline C. Whitacre‡

Chronic inflammation contributes to numerous diseases, and regulation of inflammation is crucial for disease control and resolution. Sex hormones have potent immunoregulatory abilities. Specifically, estrogen influences immune cells and inflammation, which contributes to the sexual dimorphism of autoimmunity and protection against disease seen during pregnancy in multiple sclerosis (MS) and its animal model, experimental autoimmune encephalomyelitis (EAE). Although long thought to act primarily on T cells, recent evidence demonstrated that myeloid cells, such as dendritic cells (DCs), are essential in mediating estrogen’s protective effects. Estriol (E3), a pregnancy-specific estrogen, has therapeutic efficacy in MS and EAE, and we evaluated whether E3 could act exclusively through DCs to protect against the inflammatory autoimmune disease EAE. Levels of activation markers (CD80 and CD86) and inhibitory costimulatory markers (PD-L1, PD-L2, B7-H3, and B7-H4) were increased in E3 DCs. E3 DCs had decreased proinflammatory IL-12, IL-23, and IL-6 mRNA expression, increased immunoregulatory IL-10 and TGF-β mRNA expression, and a decreased ratio of IL-12/IL-10 protein production. Importantly, transfer of E3 DCs to mice prior to active induction of EAE protected them from developing EAE through immune deviation to a Th2 response. This protection was apparent, even in the face of in vitro and in vivo inflammatory challenge. In summary, our results showed that E3 generates tolerogenic DCs, which protect against the inflammatory autoimmunity disease EAE. Targeted generation of tolerogenic DCs with immunomodulatory therapeutics, such as E3, has potential applications in the treatment of numerous autoimmune and chronic inflammatory diseases.

*C Department of Veterinary Biosciences, College of Veterinary Medicine, The Ohio State University, Columbus, OH 43210;†Section of Oral Biology, College of Dentistry, The Ohio State University, Columbus, OH 43210; and‡Department of Molecular Virology, Immunology, and Medical Genetics, The Ohio State University, Columbus, OH 43210

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Address correspondence and reprint requests to Dr. Tracey L. Papenfuss, The Ohio State University, 185 Veterinary Medicine Academic Building, 1900 Coffey Road, Columbus, OH 43210. E-mail address: papenfuss.1@osu.edu

Abbreviations used in this article: CDS, cumulative disease score; CT, cycle threshold; DC, dendritic cell; E2, 17β estradiol; E3, estriol; EAE, experimental autoimmune encephalomyelitis; Flt3L, Flt3 ligand; IL-12, IL-23, and IL-6; ICOSL, ICOS ligand; MOG, myelin oligodendrocyte glycoprotein; MS, multiple sclerosis; Pb, placebo; Tg, transgenic; Tol-DC, tolerogenic DC; Treg, regulatory T cell.
abilities has resulted from studies exploring E2’s effects in EAE, and it was long thought that these actions were mediated primarily through direct actions on T cells. However, recent evidence suggested that myeloid cells, such as dendritic cells (DCs), are the primary cells responsible for mediating estrogen’s protective abilities in EAE (18, 19).

DCs are innate immune cells of the myeloid lineage uniquely able to drive the differentiation of naïve Th cells into Th1, Th2, Th17, and Tregs through a combination of MHC class II–TCR–peptide interactions, costimulatory molecules, and cytokine production (20, 21). Long known to promote inflammatory immune responses, DCs are increasingly being recognized for their potent regulatory capabilities in limiting inflammation, with possible therapeutic applications in the treatment of autoimmune and inflammatory diseases (22–24). Such tolerogenic DCs (Tol-DCs) are programmed by a variety of factors, including innate immune receptor signaling, cell–cell interactions, and microenvironmental cues (e.g., steroid hormones, cytokines, other soluble mediators) (20, 25–32). Steroid hormones, such as glucocorticoids and vitamin D, are potent generators of Tol-DCs. Studies showed that these Tol-DCs are arrested in an immature state, expressing low CD40, CD80, CD86, and MHC class II expression and decreased Ag-processing and -presentation capabilities (20, 25–29). Tol-DC populations are diverse and regulate immune responses through numerous potential mechanisms, including altered costimulatory molecule expression, inhibition of proinflammatory mediators (e.g., IL-12, TNF-α, NO, NF-kB), enhanced production of immunoregulatory factors (e.g., IL-10, TGF-β, IDO, arginase), or increased expansion and/or differentiation of Tregs (20, 21, 33–36).

Estrogen’s effects on DCs are less well studied than glucocorticoids or vitamin D; however, DCs, like other myeloid cells, are fully able to express estrogen receptors and respond to estrogens (37, 38). Most studies use E2 and show an anti-inflammatory effect of E2 on mature DCs and a potent effect on resting and inflammatory DC differentiation (37, 39, 40). Studies in autoimmune models demonstrated that E2 exposure can diminish Ag presentation, enhance Th2 responses, and enhance production of immunoregulatory IDO or Tregs (18, 33, 41). Although E2 influences DC–T cell interactions, the effects of other estrogens, such as E3, have not been thoroughly investigated, although E3 was shown to decrease proinflammatory NO and TNF-α in mouse microglia and enhance IL-10 production in CD68+ monocytes (20, 25–32).

We found that E3 exposure generates Tol-DCs with stable regulatory abilities that are resistant to inflammatory challenge. In vivo E3 exposure increased the expression levels of activation markers CD80, CD86, and MHC class II and inhibitory costimulatory markers PD-L1, PD-L2, B7-H3, and B7-H4 in DCs compared to levels seen in placebo (Pb)-treated mice. E3 Tol-DCs also expressed decreased proinflammatory IL-12, IL-23, and IL-6 mRNA levels and increased immunoregulatory IL-10 and TGF-β mRNA levels compared with Pb DCs; had a decreased ratio of IL-12/IL-10 protein production; and diminished T cell proliferation. Importantly, E3 Tol-DCs protected mice from developing EAE through immune deviation to a Th2 response, and in vitro and in vivo inflammatory stimuli failed to abrogate the protective effect of E3 Tol-DCs. Taken together, these data suggest that immunomodulatory compounds, such as E3, generate Tol-DCs that may be useful in the treatment of autoimmune or chronic inflammatory diseases.

Materials and Methods

Mice

Female C57BL/6 (H-2b) mice (4–6 wk old) were purchased from Jackson Laboratories (Bar Harbor, ME), housed four or five/cage, and maintained on a 12-h light/dark cycle. Mice were allowed to rest for 7 d prior to pellet implantation; 5–7 d later they were given Flt3L to expand DCs, and age-matched mice were used for DC transfer and EAE studies. All animals received identical commercially available chow and were housed and cared for according to the institutional guidelines in the University Laboratory Animal Resources at The Ohio State University. Myelin oligodendrocyte glycoprotein (MOG) TCR transgenic (Tg) mice were maintained in-house and screened by flow cytometry for expression of the Vα3.2/Vβ11 MOG-specific TCR.

Hormone pellet administration and in vivo DC expansion

Sixty day-release 5-mg estradiol (E3) pellets or Pb pellets containing carrier without hormone (Innovative Research of America, Sarasota, FL) were implanted s.c. between the shoulder blades; mid- to late-pregnancy levels were achieved by day 5 postimplantation, as previously reported (42). Mice were allowed to recover for 5–6 d prior to Flt3L (Flt3L) administration. Human recombinant Flt3L (hFL; kindly provided by Angen Thousand Oaks, CA) was used to expand DCs in vivo through daily administration of hFL (200 ng/ml hFL in 200 μl 0.1% mouse serum albumin) s.c. in the nape of the neck for 9 d. Control mice were given vehicle alone (200 μl 0.1% mouse serum albumin). Alternatively, DCs were expanded in vivo by administering 4 million B16 melanoma cells (expressing murine Flt3L s.c.) and mice were sacrificed 3–4 wk after administration. All comparisons of DC populations were from identically treated mice (i.e., E3 and Pb DCs from nonexpanded mice were compared or E3 or Pb DCs from Flt3L-expanded mice were compared).

CD11c+ and CD4+ cell preparation and culture

Splenic DCs from C57BL/6 female mice were positively selected and purified using CD11c magnetic bead separation (Miltenyi Biotec, Auburn, CA) and pulsed with MOG35–55 for 2 h in supplemented RPMI 1640 containing 10% FBS, 25 mM HEPES, 2 mM l-glutamine, 50 μM penicillin, 50 μg/ml streptomycin, and 5 × 10−5 M 2-ME. Purified CD11c+ DCs were washed and administered i.v. to naive recipients (1–2 × 106) 1 d prior to EAE induction or used for companion in vitro cocultures. CD11c+ DCs were phenotypically evaluated for cell surface marker expression (flow cytometry) or cytokine production (ELISPOT) and in functional CD11c+/CD4+ cocultures (proliferation assays, ELISA, ELISPOT, and flow cytometry). Spleen and lymph node CD4+ T cells were purified using CD4+ magnetic bead separation (Miltenyi Biotec, Auburn, CA) for cocultures, and DCs were plated at a ratio of 1:5 or 1:10 with 2 × 105 purified CD4+ T cells and 4 × 104 or 2 × 105 DCs, respectively. Purity of the negative and positive fractions was assessed by flow cytometry analysis, and cell populations with a purity >95% were used for experimentation. Cells were cultured in supplemented RPMI 1640 with or without the following stimuli: with medium, 10 g/ml MOG35–55, 5 μg/ml LPS, 3 μg/ml ConA, or 3 μg/ml anti-CD3 for 24–96 h. Cells were cultured in round-bottom 96-well plates (2 × 103 T cells/well) or in transwell 96-well plates (separating purified DCs from responder [2 × 105] splenocytes) for 24–96 h, with responder cell numbers remaining constant. Cell viability was assessed by 7-aminoactinomycin D staining, with subsequent evaluation by flow cytometry at 24, 48, 72, and 96 h of culture.

Proliferation assays

Cellular proliferation was evaluated by culture of cells for 48–72 h, with incorporation of [3H]thymidine for the last 18 h of culture. Cells were harvested and counted using Perkin Elmer Top Count NXT, with Packard’s Top Count NXT software. Results are expressed as the total cpm (mean...
cpm of cultures with Ag/mean cpm of cultures with medium alone) ± SEM for all animals in the group.

**Evaluation of cytokines: ELISA and ELISPOT**

For cytokine protein determination, supernatants were collected from in vitro cultures of DCs 24 h after purification or from DC–T cell cocultures. OPT-EIA Sandwich ELISA kits (PharMingen, San Diego, CA) were used to determine the levels of IL-12p40, IFN-γ, TNF-α, IL-2, IL-4, and IL-10, and colorimetric changes were read with a SpectraMax Plus 384 spectrophotometer and analyzed with SoftMax Pro software (Molecular Devices, Sunnyvale, CA). The frequency of cytokine-secreting cells was determined for IL-12, IFN-γ, IL-2, IL-4, and TNF-α using ELISPOT development modules (R&D Systems, Minneapolis, MN), and samples were run in triplicate with medium, MOG35–55, LPS, ConA, or anti-CD3. Computer-assisted image analysis, using KS ELISPOT software and microscope control processor MCP4 (Carl Zeiss Vision GmbH, Thornwood, NY), was used to analyze results. Background cytokine-producing cells/million ± SEM for all animals in a group.

**Flow cytometry**

Single-cell suspensions of spleen and lymph nodes cells were evaluated for CD4, CD25, and intracellular Foxp3 expression. For intracellular staining, cells were stained first for surface markers and then fixed and permeabilized prior to intracellular Foxp3 staining. DC populations from pelleted mice were evaluated for overall CD11c expression and gated on CD11c+ (from nonpurified spleenocytes) or CD11c+ (from purified CD11c+ populations) to determine levels of CD80/B7-1, CD86/B7-2, PD-1, CD40 (BD Biosciences, San Jose, CA), PD-L1/B7-H1, PD-L2/B7-DC, B7-H3, and B7-H4 (eBiosciences, San Diego, CA). Three-color flow cytometry was performed on DC and T cell populations using FITC-, PE-, or allophycocyanin-conjugated mAbs, and the corresponding isotype controls (BD Biosciences, San Jose, CA) and cells were processed on a BD FACSCalibur flow cytometer and analyzed using Cell Quest analysis software (BD Biosciences, San Jose, CA).

**RT-PCR and real-time RT-PCR**

Purified CD11c+ DCs from mice implanted with E3 or Pb pellets were frozen at −80°C in TRIzol reagent and then used for RNA extraction. The following primers were used in RT-PCR: 5′-AAGTGCTGCTTCCCTGCTAAGGACAAGGCTG-3′ and 3′-AACCCACACCAAGGAGAAAGGAGA-5′ for IL-12p40; 5′-ACCACCGTTGACTCATTCTG-3′ and 3′-AGGGCAAGGGCTC-5′ for TGF-β; 5′-CTCTCTGAAA-3′ and 3′-TGCTCTTTGTGATGTCAGCTG-5′ for IL-12p35; 5′-CCCGAGAGAGAGTCACGAGGAGAAG-3′ and 3′-CAAGAGACCTTTTACGGCATC-5′ for IL-6; 5′-CTCTCTAAAGGAAGACGCTG-3′ and 3′-GAATCTGAAAAGCCTCTCAAGG-5′ for TNF-α; 5′-GCCCGCGCGGACTTACATPA-3′ and 3′-AACCTTGAGTGGAGGAGGAGGAG-5′ for IL-10; whereas anti-inflammatory mediator mRNA is increased in E3 DC populations. Similar to the overall CD11c+ population, mean fluorescence levels seemed to increase in DCs exposed to E3 (Fig. 1D), whereas mean fluorescence levels did not differ significantly for most markers, with the exception of slight increases in B7-H3 (Fig. 1D). E3 did not seem to influence ICOS ligand (ICOSL) or CD40, suggesting that it increases select stimulatory and inhibitory markers in DC populations, which may contribute to the tolerogenic ability of E3 Tol-DCs.

Given that specific CD8α+ and CD8α− DCs can directly impact Th1/Th2 responses, we next determined whether E3 differentially alters the levels of CD8α+ DCs and the expression of stimulatory or inhibitory markers in CD8α+ and CD8α− DC subpopulations (43). E3 consistently increased CD8α+ DCs ∼10–15% over Pb treatment (Fig. 2A), suggesting that increased CD8α+ levels may contribute to the tolerogenic abilities of transferred E3 DCs. Similar to the overall CD11c+ population, mean fluorescence levels did not differ between E3 and Pb DCs for most markers. Only in the CD8α+ CD11c+ subset were mild increases in CD80 and PD-L1 and decreases in ICOSL and MHC class II seen (Fig. 2C), suggesting that altered levels of these molecules in the CD8α+ DC subset may significantly contribute to the tolerogenic ability of E3 Tol-DCs.

Expression of proinflammatory mediator mRNA is decreased, whereas anti-inflammatory mediator mRNA is increased in E3 Tol-DCs.

**Expression of proinflammatory and immunoregulatory factor mRNA from E3 and Pb DCs was investigated using real time RT-PCR. E3 DCs demonstrated decreased expression of proinflammatory IL-12p40, IL-23p19, IL-6, and inducible NO synthase (NOS2) mRNA and increased expression of immunoregulatory TGF-β and IL-10 mRNA (Fig. 3). Decreased IL-12 mRNA levels are consistent with DC–T cell coculture results in Fig. 2B and, together with decreased IL-12p40, IL-23p19, and IL-6 mRNA levels, suggested that E3 may decrease the proinflammatory capabilities of DCs, which may affect the generation of Th1 and Th17 cells. Similarly, increased expression of immunoregulatory cytokines, such as TGF-β and IL-10 mRNA, may also contribute to the immunoregulatory role of E3 DC-mediated protection. No consistent differences were seen in the expression of TNF-α and immunoregulatory mediators arginase (Arg2) and IDO mRNA between E3 and Pb DCs, suggesting that these factors are not involved in E3 DC action (Fig. 3). Interestingly, slight increases in IL-12p35 expression were noted in E3 DCs (Fig. 3).**

**Results**

**In vivo E3 exposure increases DCs expressing activation and inhibitory cell surface markers**

To determine whether pregnancy levels of estrogen could distinctly affect DCs, splenic DCs from mice implanted with E3 (pregnancy levels) or Pb pellets were phenotypically evaluated. E3 exposure increased the levels of activation markers CD80, CD86, and MHC class II in DCs compared with mice implanted with Pb pellets (Fig. 1A). CD40 levels did not differ significantly, although trends of increased CD40 levels in E3 DCs were seen in some experiments (data not shown). Mean fluorescence data suggested that although E3 increased the percentage of these markers, only CD80 mean fluorescence levels seemed to increase in DCs exposed to E3 (Fig. 1C). CD86, CD40, and MHC class II mean fluorescence levels did not increase, and MHC class II levels may have decreased slightly in E3-exposed DCs. These results suggested that E3 increases the overall level of activation in the splenic DC compartment but may not upregulate the expression of activation markers on individual DCs. We next evaluated whether E3 affected the expression of inhibitory costimulatory molecules. Similar to what was seen in activation markers, levels of the inhibitory markers PD-L1, PD-L2, B7-H3, and B7-H4 were increased after E3 exposure (Fig. 1B), whereas mean fluorescence levels did not differ significantly for most markers, with the exception of slight increases in B7-H3 (Fig. 1D). E3 did not seem to influence ICOS ligand (ICOSL) or CD40, suggesting that it increases select stimulatory and inhibitory markers in DC populations, which may contribute to the tolerogenic ability of E3 Tol-DCs.

**E3 DCs have an increased IL-10/IL-12 ratio due to decreased IL-12 production and not through increased IL-10 production**

Given the described roles of IL-12 and IL-10 in modulating T cell responses and influencing EAE, E3 and Pb DCs were cultured...
in vitro with or without LPS, and IL-12 and IL-10 protein expression was measured (44). E3 DCs produced less IL-12p40 after 96 h in culture in the media control group and after stimulation with LPS (Fig. 4A). To determine whether IL-12 production by E3 DCs could be enhanced or altered by Ag exposure, the neuroantigen MOG was added to DCs and cocultures of DCs with naive Ag-specific CD4+ T cells specific to MOG. Fig. 4B demonstrates that IL-12 production on a per-cell basis was lower in E3 DC cultures compared with Pb DCs in the media control group (98 $\pm$ 614 and 174 $\pm$ 3, respectively), as well as following MOG stimulation (124 $\pm$ 64 and 192 $\pm$ 11, respectively). These results demonstrated that the decreased IL-12 production by E3 DCs remains, even when DCs are stimulated directly with LPS or indirectly by Ag-stimulated T cells. Given the importance of IL-10 in regulating immune function in select Tol-DCs, we evaluated IL-10 production following in vitro stimulation of E3 and Pb DCs. IL-10 production did not differ between E3 and Pb DCs in the media control or following LPS stimulation (Fig. 3C), but the ratio of IL-10/IL-12 was increased for E3 DCs in the media-treated mice (ratio of 1) and LPS-stimulated mice (ratio of 1.59) compared with Pb DCs (ratios of 0.5 and 0.53 for media and LPS, respectively). This increased IL-10/IL-12 ratio occurred as a result of decreased IL-12 rather than because of elevated IL-10.

T cells proliferate less when cocultured with E3 Tol-DCs

To assess whether the described E3 Tol-DC phenotype had functional consequences on T cell differentiation, E3 DCs and Pb DCs were cultured with purified naive CD4+ MOG-specific T (Th0) cells from MOG TCR Tg mice. MOG Th0 cells cultured with E3 DCs proliferated less than did those cultured with Pb DCs in the presence of media, the neuroantigen MOG, and APC-stimulating LPS, but they did not differ when T cells were specifically stimulated with anti-CD3 (Fig. 5A). These results suggested that E3 DCs have a baseline ability to inhibit T cell proliferation (i.e., media group) that is resistant to DC-activating stimuli (i.e., LPS group). Importantly, E3 DCs are also able to inhibit T cell proliferation in response to Ag (i.e., MOG group) but not in response to anti-CD3 (Fig. 5B). Additionally, soluble factors produced by DCs separated from splenocytes in a transwell system (Fig. 5B) and splenocytes cultured with E3 Tol-DC-conditioned media (data not shown) suggested that cell–cell interaction is necessary for the full spectrum of regulatory effects of E3 DCs on T cells.

E3 Tol-DC recipient mice develop less severe EAE

To assess the biological relevance of the E3 DC regulatory capabilities, we evaluated the protective ability of E3 DCs in the inflammatory autoimmune disease EAE. Regardless of the mechanism, simple hypoproduction of proinflammatory mediators (e.g., IL-12) by E3 Tol-DCs is likely insufficient to explain the prevention of EAE. Specifically, mice receiving E3 Tol-DCs have a full complement of resident DCs presumably capable of generating IL-12 and IL-23 necessary to promote Th1 and/or Th17 responses responsible for EAE clinical disease. Mice receiving E3 DCs developed less severe disease than did Pb DC recipients, had a decreased cumulative disease score (CDS) and decreased clinical severity, and failed to relapse or develop significant chronic EAE compared with Pb DC recipients (Fig. 6, Table I). The protective effect of E3 DCs was not influenced by exposure to LPS in vitro or to CFA and pertussis toxin adjuvants in vivo (Fig. 7, Table II). These results suggested that E3 DCs are resistant to in vitro and
in vivo inflammatory signals and that the regulatory abilities of these E3 DCs are maintained in vivo.

E3 DC recipients promote immune deviation without generating Tregs

Active induction of EAE generates Th1 and/or Th17 cells that drive development of disease, whereas Th2 and Treg responses are protective. To assess whether E3 DCs protected mice through altering the Th1/Th2/Th17/Treg balance, peripheral immune organs were evaluated at day 10 postimmunization. Lymph node cells from immunized mice showed decreased Ag-specific proliferation to the immunizing neuroantigen MOG in E3 DC recipients (Fig. 8A). Proliferation did not differ when lymph node cells were stimulated with anti-CD3 or LPS (Fig. 8A). Because Tregs did not seem to be involved in the E3 DC-mediated protection against EAE, cytokine profiles were evaluated in E3 and Pb DC recipients. Lymph node cells from immunized mice receiving E3 DCs prior to immunization produced less IFN-γ and increased IL-4 following MOG stimulation compared with Pb DC recipients (Fig. 9A). Although IL-17 levels were slightly higher in the Pb DC recipients, the levels did not reach statistical significance (Fig. 9B). Taken together, these data suggested that E3 Tol-DCs alone were able to protect mice from developing EAE.

FIGURE 3. E3 DCs express decreased proinflammatory and increased immunoregulatory mRNA levels. E3 and Pb DCs were purified with CD11c magnetic beads and evaluated by real-time RT-PCR to determine the expression of proinflammatory and immunoregulatory mediators. Adjusted average ΔCT values are shown for proinflammatory IL-12p40, IL-23p19, IL-12p35, IL-6, TNF-α, and immunoregulatory TGF-β and IL-10. E3 and Pb DCs were obtained from pooled splenocytes of mice implanted with E3 or Pb pellets. *p < 0.05 of triplicates. Individual p values were determined based on triplicates of individual experiments and reported when a consistent difference (trend or p < 0.05) was seen in at least three separate experiments (at least six mice per E3 or Pb treatment).
and do so through Ag-specific immune deviation to a Th2 response. Additionally, this protection is resistant to inflammatory challenge in vitro (i.e., LPS) or in vivo (i.e., Th1/Th17-promoting adjuvants).

Discussion
Chronic inflammation is recognized as a significant contributor to a wide array of human diseases, including autoimmunity, cardiovascular disease, asthma, allergies, and cancer. The ability to regulate such inflammation is crucial for control and resolution of autoimmune and chronic inflammatory diseases. Although Tregs are known to have potent regulatory abilities, increasing evidence demonstrates that myeloid cells can dramatically regulate innate and adaptive immune responses and can be used therapeutically to regulate inflammation. Tol-DCs are one such regulatory myeloid cell population, and there is much interest in immunotherapy with these cells (20, 21). A primary difficulty in Tol-DC immunotherapy is the inadequate understanding of these cells and the factors that program them to become tolerogenic (32). Given the importance of myeloid cells in estrogen-mediated protection against autoimmunity, we asked whether E3 could generate Tol-DCs capable of protecting against the inflammatory autoimmune disease EAE. We found that E3 exposure altered the subset composition, levels of activation, and inhibitory markers in DCs and generated functional Tol-DCs. Importantly, transferred E3 Tol-DCs alone could protect mice from EAE, and the protective abilities of E3 Tol-DCs seem to be resistant to in vitro and in vivo inflammatory challenge. Other estrogens, such as E2, were shown to mediate protection through actions on myeloid cells, such as DCs, but safety concerns limit the therapeutic application of E2 (3, 6, 9, 12, 13, 45). This study demonstrated that E3, an estrogen with efficacy...

FIGURE 4. E3 DCs produce less IL-12 and similar IL-10 compared with Pb DCs. A, Purified CD11c+ E3 or Pb DCs were cocultured with purified CD4+ naive MOG-specific T cells for 96 h, and IL-12p40 and IL-10 protein were evaluated in culture supernatant by ELISA. B, Cell-specific IL-12 production (ELISPOT) was lower in E3 DC cultures compared with Pb DC cultures in the media control group and following MOG stimulation. C, IL-10 cytokine production between E3 and Pb DCs show decreased IL-12 production in E3 DC groups. Data are representative of at least three separate experiments. Bar graphs show mean ± SEM and are representative of three separate experiments (at least six mice per E3 or Pb treatment). *p < 0.05.

FIGURE 5. T cells proliferate less in the presence of E3 DCs, and cell–cell contact is required for this effect. Purified CD11c+ E3 DCs or Pb DCs were cultured directly with CD4+ TCR transgenic T cells specific for MOG (A) or separated from MOG Tg T cells by transwell culture inserts for 96 h in the presence of MOG, anti-CD3, and LPS (B). Cellular proliferation was determined by the amount of incorporated [3H]thymidine and read as cpm. Data are representative of five or three separate experiments, respectively. Bar graphs show mean ± SEM. *p < 0.05 for standard proliferation assays from at least three separate experiments (12 mice per E3 or Pb treatment) and two separate experiments (5 mice per E3 or Pb treatment) for transwell experiments.

FIGURE 6. E3 DC recipients have reduced autoimmunity. Clinical scores of mice actively immunized 1 d after receiving 8–10 million E3 or Pb DCs (i.v.) pulsed in vitro with MOG35–55. Data are representative of seven separate experiments of at least five mice per group.
in MS patients and an increased safety profile compared with E2, programmed DCs to become tolerogenic and that E3 Tol-DCs protected mice against inflammatory autoimmune disease (3, 6, 9, 11, 12).

Tol-DCs are diverse and promote Th2 and other regulatory responses through a variety of mechanisms, including actions through specific DC subsets, soluble factors, and cell–cell contact. Our results showed that E3 increased the levels of activation and inhibitory molecules in the overall splenic DC compartment and specifically increased the percentage of CD8α+ DCs (Figs. 1, 2). E3 did not affect the levels of plasmacytoid DCs, which were consistently 3–5% of our purified population (data not shown).

Numerous studies suggested that DC subsets impact tolerogenic versus regulatory immune responses, with CD8α+ potently producing IL-12 and promoting Th1 responses (46–50). Thus, it was surprising that E3 increased the CD8α+ subset, yet decreased IL-12 production was seen. Although increased CD8α+ levels may result from Flt3L’s described propensity to increase CD8α+ subsets, the reason for overall decreased IL-12 production, even in the face of increased CD8α+ DCs, is not known. It is possible that E3 decreases the functional ability of CD8α+ DCs to produce IL-12 or promote Th2 responses, allowing the effects of CD8α− DCs to be more pronounced in the E3 DCs. Indeed, it was reported that CD8α− DCs are more tolerogenic (43, 46). However, Pb DCs have increased CD8α− levels, so increased CD8α− DC levels alone do not promote protective responses. Altered expression of surface markers in CD8α− DCs (e.g., increased CD80 and PD-L2) or alterations in IL-10 and IL-12 expression in either subset may contribute to E3 Tol-DC action. Given IL-12’s important role in determining the Th1/Th2 balance in vivo, decreased IL-12 production by DCs, particularly CD8α+ DCs, may contribute to the

![Image](http://www.jimmunol.org/)

**FIGURE 7.** E3 DC regulatory function is not affected by inflammatory challenge, and protection from autoimmunity is maintained. Actively immunized mice received 1–2 million purified E3 or Pb DCs exposed to MOG35–55, with or without LPS, in vitro 24 h prior to EAE induction. Delayed onset and protection from clinical EAE was evident in E3 (± LPS) DC recipients. Data are representative of two independent experiments with five mice per group.

<table>
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<th>DC Treatment</th>
<th>Incidence (%)</th>
<th>Onset (d)</th>
<th>Peak (d)</th>
<th>CDS</th>
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<td>1.2 ± 0.7*</td>
<td>1.3 ± 0.8*</td>
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<tr>
<td>Pb DC</td>
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DC recipients have reduced autoimmunity; they had decreased peak severity, clinical severity, and resolution of clinical signs. Data are representative of seven separate experiments of at least five mice per group.

**Table I.** Incidence, day of disease onset, peak day of clinical disease, and CDS of E3 and Pb DC recipients.

**Table II.** Peak day of clinical disease and CDS of mice receiving E3 DCs and Pb DCs with or without LPS exposure.

<table>
<thead>
<tr>
<th>DC Treatment</th>
<th>CDS Peak (d)</th>
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<td>E3</td>
<td>5.2 ± 5.2*</td>
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<tr>
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<tr>
<td>Pb</td>
<td>23.6 ± 7.7</td>
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<tr>
<td>Pb + LPS</td>
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Protection from autoimmunity was maintained, even with inflammatory challenge. Recipients had decreased peak severity, clinical severity, and resolution of clinical signs. Data are representative of three separate experiments, with a total of 10 mice per group. All groups were compared with Pb DC recipients.

**FIGURE 8.** E3 DC recipients proliferate less to immunizing Ag, and Tregs do not mediate the protection. The proliferative ability of lymph node cells in E3 DC recipient mice was evaluated. A, Lymph node cells from E3 DC recipient mice proliferated less to MOG than did Pb DC recipients. The decreased proliferative ability was not through increased numbers of Tregs. E3 DC recipients are not protected by increased numbers of CD8α+ and CD8α− subsets and their cytokine profiles to mediated E3 DC protection are not known but are the subject of ongoing studies.

It is well known that IL-12 and IL-23 potently drive pathogenic T cell differentiation in EAE (i.e., Th1 and Th17, respectively), whereas IL-10 and TGF-β can dramatically regulate inflammation and EAE (51). Although E3 DCs produce less IL-12 protein, our results also showed decreased mRNA levels of proinflammatory IL-12, IL-23, IL-6, and NO, suggesting that E3 decreases the expression of numerous proinflammatory mediators. Impaired NF-κB signaling may be responsible for decreased proinflammatory mediators in E3-exposed DCs. E2 is known to influence genomic and nongenomic actions of NF-κB and other signaling pathways, and studies showed impaired or altered NF-κB signaling in E2-exposed CNS macrophages or splenic CD11c+ cells (52–54). Although E3 was shown to inhibit NF-κB in T cells, its actions on DCs are not known and were beyond the scope of this study; however, it is likely that E3 alters NF-κB, similar to E2 (55).
Such alterations may contribute to the hypoproduction of pro-inflammatory mediators, such as IL-12 seen in our studies and reported by other investigators in human monocyte-derived and mouse splenic DCs following estrogen exposure (18, 41, 55). Alternatively, TGF-β is known to reduce the stability of IL-12p40 mRNA and may contribute to the resultant decrease in IL-12 protein (56, 57). Regardless of the mechanism, simple hypoproduction of proinflammatory mediators (e.g., IL-12) by E3 Tol-DCs is likely insufficient to explain the prevention of EAE. Specifically, mice receiving E3 Tol-DCs have a full complement of resident DCs presumably capable of generating IL-12 and IL-23 necessary to promote Th1 and/or Th17 responses responsible for EAE clinical disease. Thus, active immunoregulatory mechanisms likely contribute to E3 Tol-DC-mediated protection.

TGF-β and IL-10 are potent immunoregulatory factors used by Tol-DC populations. IL-10, in particular, was shown to be used by other Tol-DCs (20, 58) and is upregulated in DCs exposed to E2 and in splenic immune cells exposed to E3 (18, 23, 41, 59). Our data showed that E3 Tol-DCs had increased expression of IL-10 and TGF-β mRNA (Fig. 3), suggesting that both of these immunoregulatory factors may contribute to E3 Tol-DC actions. Surprisingly, IL-10 protein levels increased after in vitro culture, but this is most likely attributed to suboptimal IL-10 production with LPS stimulation. Interestingly, a recent study suggested that CD40 signaling is important in inducing IL-12 and IL-23 in E3 Tol-DCs, so the absence of alterations in CD40 may explain why IL-10 protein is not increased in E3 Tol-DCs (60). Alternatively, similar overall IL-10 levels in E3 and Pb DCs suggested that CD80α-DCs in the E3 Tol-DCs may produce proportionately more IL-10 than similar CD80α-DCs in the Pb DC group, given that a lower percentage of E3 DCs are CD80α- than are Pb DCs. It is also possible that distinct increases in IL-10 production are not essential for regulation by E3 Tol-DCs but that an increased IL-10/IL-12 ratio is more important in determining the resultant immune response (47, 61).

Other immunoregulatory cytokines, such as TGF-β, may contribute to E3 Tol-DCs’ regulatory action. TGF-β is a complex cytokine that influences the Treg/Th17 balance, is an important contributor to oral tolerance and the generation of Tregs and Th3 cells, and is known to control autoimmunity through actions on DCs (62–64). The increased mRNA levels of TGF-β seen in E3 Tol-DCs suggest that TGF-β contributes to regulatory actions of these cells. Given the complexity of TGF-β regulation (i.e., post-transcriptional and post-translational modifications and latent forms), assessment of biological activity by ELISA is reportedly imprecise and were beyond the scope of this study, but our data are intriguing, and further studies are needed to evaluate the contribution of TGF-β in E3 Tol-DC regulatory actions. Although other soluble mediators, such as IDO and arginase, were reported to play important roles for other Tol-DCs or regulatory myeloid cells, we found that they did not seem to be acting in E3 Tol-DCs based on similar mRNA levels seen between E3 or Pb DCs (33, 34, 65, 66). Rather, Fig. 5B suggests that cell–cell contact mechanisms are a more important mechanism by which E3 Tol-DCs regulate T cells.

Activation and inhibitory markers are important cell contact-mediated mechanisms by which DCs regulate adaptive immune responses. Decreased expression of activation markers, such as CD80, CD86, and MHC class II, are classically associated with dexamethasone or vitamin D Tol-DCs and mediate tolerance through their arrest in an immature state. Our data suggested that E3 acts in a distinct manner to generate Tol-DCs and that E3 Tol-DCs are not immature, based on increased expression of CD80, CD86, and MHC class II. In fact, E3 actually seems to increase the overall activation state of the splenic DC compartment, as evidenced by increased levels in CD80, CD86, and MHC class II expression in E3 versus Pb DCs. Although levels of activation markers are increased, so too are numerous inhibitory markers, such as PD-L1, PD-L2, B7-H3, and B7-H4. The inhibitory abilities of PD-L1, PD-L2, B7-H3, B7-H4, and ICOSL are well known, and the interactions of these molecules with their ligands on T cells may directly contribute to E3 Tol-DC regulatory actions (67–72). Data from transwell studies (Fig. 5) and E3 DC-conditioned supernatant (data not shown) strongly suggest that cell–cell interactions in E3 Tol-DCs directly contribute to suppression of proliferation and may be a primary mechanism by which E3 Tol-DCs protect against development of EAE. Studies showing enhanced PD-1 expression on APCs after E2 exposure suggest that the interactions between PD-1 and its ligands are one potential means by which estrogens influence DC function (16). Although these studies did not evaluate PD-L1 or PD-L2 on E2-exposed APCs, we found upregulation of these ligands on Tol-DCs, highlighting the potential role of PD-1/PD-L interactions in E3 Tol-DC regulation. Less is known about the B7-H3 and B7-H4 mechanistic actions, but B7-H3 and B7-H4 can dramatically influence EAE through a variety of potential mechanisms, including inhibition of anti-CD3-induced T cell proliferation, altering cytokine production, influencing cell cycle arrest, and activating NFAT, NF-κB, and AP-1 (67, 69–77). The fact that PD-L2 and B7-H3 expression on E3 DCs or DC subsets suggest that these particular inhibitory molecules (potentially in combination with increased CD80) may play a more prominent role than others. Thus, E3 Tol-DCs may be actively regulating immune responses through alternative activation, expression of multiple inhibitory molecules, or a combination of these. At present, the relative balance of costimulatory molecules is thought to dramatically

![FIGURE 9](http://www.jimmunol.org/)

E3 DC recipients are protected from EAE by immune deviation to a Th2 response. Lymph node cells obtained 10 d after EAE induction were evaluated for production of IL-4, IFN-γ, and IL-17. A. Cell-specific cytokine secretion of lymph node cells by ELISPOT demonstrated increased IL-4 and decreased IFN-γ production in E3 DC recipients. B. Overall IL-17 production from culture supernatants detected by ELISA showed no differences between E3 and Pb DC recipients. Data are representative of three independent experiments with five mice per group. Bar graphs show mean ± SEM. *p < 0.05.
influence the resultant immune responses, but a thorough understanding of the complex interplay and hierarchical signaling of stimulatory versus inhibitory molecules is lacking (78). Interestingly, the effects of E3 do not universally affect all costimulatory molecules because CD40 and ICOSL levels did not differ between E3 and Pb DCs, suggesting a specific effect of E3 on Tol-DC phenotype.

Our findings show that exposure of DCs to E3 programs DCs to become Tol-DCs and that E3 Tol-DCs protect through immune deviation away from pathogenic Th1/Th17 cells to a protective Th2 response. These findings mirror the Th2 response (i.e., increased IL-4 and IL-5 and decreased IFN-γ) reported in MS patients treated with E3 (10, 11). Although the mechanisms by which E3 Tol-DCs promote Th2 responses are unknown, it is surprising that neither IL-17 nor Tregs were altered, particularly given the differential effects seen in IL-23, IL-6, and TGF-β levels between E3 and Pb DCs. These cytokines play important roles in regulating the generation and balance of Th17 and Tregs, and more mechanistic studies are required to understand the relative contributions of these cytokines and other factors (e.g., retinoic acid) in affecting the generation of Th17 and CD4⁺/CD25⁺Foxp3⁺ Tregs following E3 Tol-DC exposure (16, 79–81). The absence of CD4⁺/CD25⁺ Foxp3⁺ Tregs in our study was particularly surprising given that other investigators found increased Tregs upon E2 exposure (3). It is possible that, although Treg numbers do not differ, Treg functionality may differ between E3 and Pb DC recipients or that Tregs were generated at times not evaluated in this study. Alternatively, E3 may preferentially enhance other Foxp3⁺ regulatory cells that mediate protection, such as CD4⁺/Foxp3⁺ regulatory cells (82, 83). Whether our results represent distinct actions of E3 versus E2 or whether Tregs or other regulatory cells were simply not detected at the time points evaluated remains to be determined.

Importantly, differences in estrogen, as well as model systems, may dramatically impact results. Specifically, our study evaluated in vivo effects of E3 on Flt3L DCs, whereas the majority of other studies evaluated E2 effects (typically in vitro) on GM-CSF DCs (16, 18, 84). It is known that GM-CSF generates inflammatory DCs whereas Flt3L generates steady-state (homeostatic) DCs (85). The additional impact of estrogen on both DC populations and regulatory responses (e.g., Th2, Tregs, and other regulatory cells) may likely differ depending on the relative inflammatory or resting state (40, 86). Estrogens (E2 and E3) have known effects on bone marrow DC differentiation, and our data showed that the composition of DC subsets can be altered by estrogen exposure. Thus, the very nature of DC populations may differ following estrogen exposure, which would further contribute to DC-mediated regulation of inflammation (40, 87).

In summary, we showed that the protective effects of E3 can be mediated through DCs alone and that E3 generates Tol-DCs that protect mice from developing EAE through immune deviation to a Th2 response. E3 Tol-DCs are resistant to in vitro and in vivo inflammatory stimuli, and cell–cell contact seems to be important in E3 Tol-DC actions. The precise mechanisms by which E3 alters DC actions are not known, but they may be similar to some E2-described actions, such as NF-xB inhibition, decreased production of inflammatory cytokines, inhibition through cell–cell interactions, and altered cellular migration, viability, and/or turnover (55, 56, 88, 89). E3 programs DCs to become tolerogenic, and E3 Tol-DCs protect against autoimmunity, even in the face of inflammatory challenge. The ability to program DCs to induce Th2 or tolerogenic responses has enormous therapeutic applications, and targeted generation of stable Tol-DCs to regulate inflammation is a promising therapy for the treatment of autoimmune and numerous chronic inflammatory diseases.

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Disclosures

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