Seminal Fluid Regulates Accumulation of FOXP3⁺ Regulatory T Cells in the Preimplantation Mouse Uterus Through Expanding the FOXP3⁺ Cell Pool and CCL19-Mediated Recruitment


Abstract

Regulatory T (Treg) cells facilitate maternal immune tolerance of the semiallogeneic conceptus in early pregnancy, but the origin and regulation of these cells at embryo implantation is unclear. During the preimplantation period, factors in the seminal fluid delivered at coitus cause expansion of a CD4⁺CD25⁺ putative Treg cell population in the para-aortic lymph nodes draining the uterus. Using flow cytometry, immunohistochemistry, and real-time quantitative PCR (qPCR) for the signature Treg cell transcription factor FOXP3, we confirmed the identity of the expanded lymph node population as FOXP3⁺ Treg cells and showed that this is accompanied by a comparable increase in the uterus of FOXP3⁺ Treg cells and expression of Foxp3 mRNA by Day 3.5 postcoitum. Seminal plasma was necessary for uterine Treg cell accumulation, as mating with seminal vesicle-deficient males failed to elicit an increase in uterine Treg cells. Furthermore, seminal fluid induced expression of mRNA encoding the Treg chemokine CCL19 (MIP3β), which acts through the CCR7 receptor to regulate Treg cell recruitment and retention in peripheral tissues. Glandular and luminal epithelial cells were identified as the major cellular origins of uterine CCL19, and exposure to both seminal plasma and sperm was required for maximum expression. Together, these results indicate that Treg cells accumulate in the uterus prior to embryo implantation and that seminal fluid is a key regulator of the uterine Treg cell population, operating by both increasing the pool of available Treg cells and promoting their CCL19-mediated recruitment from the circulation into the implantation site.

cytokines, immunology, pregnancy, regulatory T cells, seminal fluid, tolerance, uterus

Introduction

A state of maternal immune tolerance is needed to avoid immune attack on the developing conceptus after implantation. This is achieved through a variety of adaptations within the immune system [1]. One such adaptation is an expansion of the suppressive CD4⁺CD25⁺ regulatory T (Treg) cell subset [2–4]. Treg cells mediate suppression of the proliferation and effector function of a range of leukocyte subsets including CD4 and CD8 T cells, B cells, and NK cells, as well as influence the maturation and function of dendritic cells (DCs) and macrophages [4, 5]. The importance of Treg cells in the reproductive process is highlighted in mouse models that display inability to maintain allogeneic pregnancy in their absence [2, 6].

Treg cells comprise approximately 5%–10% of CD4⁺ T cells in rodent species and are defined by their constitutive expression of the interleukin 2 receptor-α chain (CD25) [7] and, more recently, their expression of the hallmark Treg cell marker forkhead box P3 (FOXP3) [8, 9]. The majority of Treg cells are generated within the thymus [5], and antigens within the periphery are critical in expanding thymic emigrants and inducing the de novo generation of Treg cells from naïve CD4⁺ T cells [10]. The role of fetal alloantigens in the expansion and function of Treg cells in pregnancy remains unclear [2, 3]. However, like other T cells, Treg cells require T-cell receptor (TCR) engagement and activation to fully elicit their suppressive function [11], and evidence suggests that a further consequence of TCR engagement in vivo is proliferation [12]. Therefore, it is likely that fetal antigens encountered by maternal antigen-presenting cells act to stimulate Treg cell generation, activation, and expansion.

Our recent studies suggest that delivery of male seminal fluid at coitus is involved in activating paternal antigen-reactive Treg cells at the outset of pregnancy [13]. Semen elicits an inflammation-like response and activates immune changes in many mammalian species including mice [14] and humans [15, 16]. The transmission of seminal fluid at coitus is the earliest and most frequent way in which the female immune system is exposed to paternal antigens that are later encountered on the implanting embryo. Considerable evidence supports the hypothesis that semen facilitates induction of a state of immune tolerance [17] and that this occurs in a paternal antigen-specific manner [13, 18] as a consequence of uterine DCs cross-presenting seminal fluid antigens to activate maternal T cells [19].

Previously, it has been shown that in the period between mating and embryo implantation, expansion of the CD4⁺CD25⁺ T cell population occurs in the para-aortic lymph nodes (LNs) that drain the uterus [13, 20] and that this expansion depends on female exposure to factors in the seminal plasma fraction of the male ejaculate [13]. However, the
relevance of the LN response and the significance of insemination for Treg cells in the uterus, where these cells are presumably required at implantation to exert their suppressive effector function, have not been explored. In this study, we utilized the specific Treg cell marker FOXP3 to show that there is an increase in Treg cells in the uterus prior to embryo implantation on Day 3.5 postcoitum (pc) and that Treg cell accumulation depends on female exposure to seminal plasma. We also provide evidence that seminal fluid up-regulates expression of the T-cell chemokine CCL19 in uterine glandular epithelial cells in the region of Treg cell accumulation in the endometrial stroma. We conclude that the seminal plasma acts in female tissues to induce an elevation in the Treg cell population in the uterus and that this occurs due to increased availability of circulating Treg cells as well as Treg cell migration into the uterus due to expression of the chemokine CCL19 within uterine epithelial cells.

MATERIALS AND METHODS

Mice and Surgical Treatments

C57Bl/6 (B6) and BALB/c mice were purchased from the University of Adelaide Central Animal Facility. Mice expressing a FOXP3-green fluorescent protein (GFP) fusion protein (Foxp3-GFP) mice [9], kindly provided by Dr. Alexander Rudensky (Memorial Sloan-Kettering Institute, New York, NY), were back-crossed onto a B6 background for eight generations and were then maintained by breeding homozygous pairs. All mice were housed under specific pathogen-free conditions at the University of Adelaide Medical School Animal House on a 12L:12D cycle and given food and water ad libitum. All experiments were performed in accordance with the Guiding Principles for the Care and Use of Research Animals endorsed by the Society for the Study of Reproduction, with approval from the University of Adelaide Ethics Committee.

Vasectomized (VAM), seminal vesicle-deficient (SVX), and vasectomized/seminal vesicle-deficient (VAM/SVX) BALB/c male mice were purchased as previously described [21]. VAM males were proven incapable of fertilization by examination of females 8 days after mating. For experimental matings, one to three 8- to 12-week-old B6 females were caged with intact, VAM, SVX, or SVX/VAS stud males. Females that mated with intact or VAM males were checked for the presence of vaginal plugs, whereas mating with SVX males and SVX/VAS males was detected by the presence of sperm or by video recorded confirmation of copitus, respectively. The morning of confirmation of mating was termed Day 0.5 pc. Mated females were removed from males and caged in groups of up to three per cage.

Flow Cytometry

The para-aortic (iliac) LN and inguinal LN were dissected from females at estrus or on Day 3.5 pc. In some experiments, peripheral blood was drawn by cardiac puncture after administration of 2% 2,2,2-trichloroethanol anesthesia (Avertin, Sigma-Aldrich), and mesenteric and cervical LNs, spleen, and uterus were also excised. Blood was diluted 1:2 in phosphate-buffered saline (PBS) containing heparin (Sigma-Aldrich), and peripheral blood mononuclear cells were collected after centrifugation over Lympholyte (Cedarlane, Hornby, Canada) according to the manufacturer’s instructions. For LNs and spleen, single-cell suspensions of lymphocytes were prepared by mechanical dispersion between glass microscope slides. Lymphocytes were suspended in a total of 1 ml of 0.1% bovine serum albumin (BSA) in PBS-0.05% sodium azide, pH 7.4 (fluorescence-activated cell sorting [FACS] buffer). For total cell quantification, a 100-µl aliquot was removed for later analysis. The remaining portion was washed and resuspended at 10^6 cells/ml. Aliquots of 10^6 cells were incubated (10 min, 4°C) with 0.5 µg of anti-Fcγ-IR antibody (FcBlock; BD Pharmingen, San Diego, CA) to block nonspecific binding. Cells were subsequently incubated (30 min, 4°C) with 0.5 µg each of PE anti-CD4 (L3T4; BD Pharmingen) and phycoerythrin (PE)-Cy7 anti-CD25 (PC61; eBioscience, San Diego, CA). In some experiments, biotinylated anti-CCR7 (eBioscience) followed by 0.2 µg of PE-Cy5 streptavidin (BD Pharmingen) were also added. Following surface staining, cells were fixed and permeabilized using the FixPerm Staining Buffer set (eBioscience) according to the manufacturer’s instructions. Permeabilized cells were incubated (30 min, 4°C) with 0.25 µg of allopurinol (Sigma) to block FOXP3 (JFK-16s; eBioscience). After being washed with 1× fix-permeabilization buffer (eBioscience), cells were resuspended in 500 µl of fix-permeabilization buffer and analyzed with a FACSCanto flow cytometer using FACSDivadiva software (both, BD Biosciences).

For quantification of CD4^+ cell numbers, 100-µl aliquots of cells were incubated (10 min, 4°C) with 10 µl of FACS buffer containing 0.5 µg of anti-Fcγ-IR antibody, followed by incubation (30 min, 4°C) with 10 µl of FACS buffer containing 0.5 µg of PE anti-CD4. Subsequently, 20 µl of CountBright absolute counting beads (Molecular Probes, Invitrogen, Carlsbad, CA) was added with 860 µl of FACS buffer. The absolute numbers of CD4^+ FOXP3^+ cells and CD4^+ CD25^+ cells per LN were then calculated using the FACScanto Flow Cytometer and FACSDiva software.

To evaluate CCL19 expression in uterine cells, the entire uterus was dissected and trimmed of fat and mesentery, split lengthwise, and digested in 2.5 µg/ml DNase and 1 mg/ml of collagenase A (Sigma) for 1 h at 37°C with agitation, as previously described [22]. The cell suspension was filtered and 2.5 ml of 0.1% bovine serum albumin (BSA) in PBS-0.05% sodium azide, pH 7.4 was added with 860 µl of FACS buffer. The absolute numbers of CD4^+ FOXP3^+ cells and CD4^+ CD25^+ cells per LN were then calculated using the FACScanto Flow Cytometer and FACSDiva software.

Quantitative Real-Time PCR

Uterine tissue was excised from estrous or Day 3.5 pc mice mated to intact, SVX, VAM, or SVX/VAS stud males, and RNA was extracted and reverse transcribed as previously described [23] and then resuspended in 100 µl of RNase-free water and stored at −20°C until required. Quantitative real-time PCR (qPCR) reactions were performed in duplicate in a final volume of 20 µl containing 10 µl of SYBR Green Master Mix (Applied Biosystems), 5 µl of cDNA template or GenBank accession numbers, and 3 µl of cDNA template or water (as non-template negative control). Real-time qPCR conditions were 10 min at 95°C, followed by 40 cycles of 95°C for 15 sec, and 60°C for 45 sec, using an ABI Prism 7000 sequence detection system (Applied BioSystems). After DNA amplification, PCR products were subjected to melt analysis to confirm the identity of PCR products.

TABLE 1. Real-time PCR primer sequences and Genbank accession numbers.

<table>
<thead>
<tr>
<th>Target mRNA*</th>
<th>Forward primer</th>
<th>Reverse primer</th>
<th>GenBank accession no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Foxp3 (1)</td>
<td>5′-AGG AGA AAG CTT GGA CTT ATG A-3′</td>
<td>3′-GTG GCT ACT ATG CAG GA-5′</td>
<td>NM_054019</td>
</tr>
<tr>
<td>Foxp3 (2)</td>
<td>5′-TGCC TCT CAG AGA CTA CTT GGA-3′</td>
<td>3′-TTT TGT GAC TAC GAA TTG-5′</td>
<td>NM_054019</td>
</tr>
<tr>
<td>Ccl19</td>
<td>5′-GGG CTC CAC ACC GGA ATG C-3′</td>
<td>3′-CGG AAG GTC TTT AGC ATT-5′</td>
<td>NM_011888.2</td>
</tr>
<tr>
<td>Ccl22</td>
<td>5′-CAC CCC TCC CCA ATA CTT G-3′</td>
<td>3′-CCT GGT AGC GGA CCA ATG-5′</td>
<td>NM_009137.2</td>
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<tr>
<td>Ccr4</td>
<td>5′-CCG AAC AAG GTC GTP CTT TCC-3′</td>
<td>3′-TTT CTT AGC GTG ATG GGC-5′</td>
<td>NM_00916.2</td>
</tr>
<tr>
<td>Ccr5</td>
<td>5′-GCC CCT GCC CCG CCA ATA CT-3′</td>
<td>3′-TCA GCC TCT CCA GCT TTT GGC-5′</td>
<td>NM_009117.5</td>
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<tr>
<td>Ccr7</td>
<td>5′-GAC ACC CAG GAA AAT GCT-3′</td>
<td>3′-CGG TGG TAT TCT CCG CCA A-5′</td>
<td>NM_007719.2</td>
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<td>Iga</td>
<td>5′-TGG TTC TCT TCA TGA GAG CCG GGG GC-3′</td>
<td>3′-AAG TCG TCC TGC TGC TGC GGC TTG-5′</td>
<td>NM_008399.2</td>
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<tr>
<td>Actb</td>
<td>5′-CTG GGG CCG CCC TAG GCA CCA-3′</td>
<td>3′-ACA AGC AGC TCA TTC TGA-5′</td>
<td>NM_007393.3</td>
</tr>
</tbody>
</table>

* Gene symbols and synonyms are according to nomenclature specified by Mouse Genome Informatics at http://www.informatics.jax.org/ [22].
amplicon size. Assay optimization and validation experiments were performed using cDNA from murine placental tissue to define the amplification efficiency of each primer pair. Cycle threshold (Ct) values were determined for serial dilutions of cDNA, and the linearity of detection was confirmed to have a correlation coefficient of >0.99 over the detection range when plotted as Ct versus log of RNA concentration. Primer sets for target genes were acceptable if their amplification efficiency varied from that of the Actb housekeeper primers by <10%. All RNA samples were reverse transcribed in a single batch and assayed with each individual primer set in the same PCR run. The Actb mRNA expression was shown not to vary among treatment groups. Messenger RNA abundance data were normalized to Actb mRNA expression using delta Ct (ΔCt = C_{target} - C_{actb}) values and expressed in arbitrary mRNA units as a percentage of the mean value of the estrous control group and calibrated so that the mean of the control group was equal to 1. PCR data were then log transformed to ensure normal distribution before statistical analysis.

**Immunohistochemistry**

To detect FOXP3 in paraffin-embedded uterine tissue, 7-μm-thick longitudinal mid-sagittal sections were dewaxed in xylene and rehydrated. Sections were subjected to antigen retrieval by incubation in citrate buffer (10 mM sodium citrate, pH 6.0) in an autoclave (121°C, 15 psi) for 10 min. Nonspecific antibody binding was blocked by incubation (1 h, room temperature) with 10% normal rabbit serum (Sigma-Aldrich) in 1% PBS-0.025% Tween-20 (Sigma-Aldrich), and endogenous peroxidase activity was blocked by incubation for 10 min in 3% hydrogen peroxide (Sigma-Aldrich). Sections were then incubated with anti-FOXP3 (FKJ-16s) primary antibody (1:400 dilution) overnight at 4°C, followed by incubation with biotinylated goat anti-rat (Dako, Glostrup, Denmark) secondary antibody (1 h, room temperature). Sections were treated with streptavidin-conjugated horseradish peroxidase (Vector Laboratories, Burlingame, CA) according to the manufacturer’s instructions, and detection was performed using a diaminobenzidine peroxidase substrate kit (Vector Laboratories). Images were captured using a NanoZoomer 1.0 unit (Hamamatsu, Shizuoka, Japan) at a zoom equivalent of a 20× objective lens. FOXP3+ cell density was quantified as the number of FOXP3-positive cells per mm² area of tissue (both endometrial and myometrial compartments in a total of 12 sections per mouse), which was calculated by tracing the section outline using NDP-view software (Hamamatsu) and manual counting of individual FOXP3+ cells.

To colocalize CCL19 and FOXP3, the uterus was excised on Day 3.5 pc after mating with intact, SVX, VAS, or SVX/VAS BALB/c males. A) The proportion of CD4+CD25+ cells in para-aortic LN expressing FOXP3, as detected by reactivity with FKJ-16s antibody, is shown. B) Expression of CD4 and CD25 in para-aortic LN FOXP3+ cells, as detected by reactivity with FKJ-16s antibody, is shown. C) Expression of FOXP3, as detected by reactivity with FKJ-16s of GFP positivity in para-aortic LN cells from Foxp3Gfp mice, for expression of either FOXP3 marker is shown. Data in A–C are representative of n ≥ 4 mice on Day 3.5 pc, and similar staining patterns were evident in n ≥ 4 estrous mice. 

D) Absolute numbers of CD4+FOXP3+ cells in the para-aortic LN are shown. E) Absolute numbers of CD4+FOXP3+ in the inguinal LNs are shown. F) Absolute numbers of CD4+CD25+ cells in the para-aortic LNs are shown. G) Absolute numbers of CD4+CD25+ in the inguinal LNs are shown. Data are means ± SEM, and different letters denote statistical significance among groups. Groups in D–G comprise n = 10–15 mice per group.
FIG. 2. The effect of seminal fluid exposure on FOXP3\(^+\) cell density and Foxp3 mRNA expression in the uterus. Uterine tissue was recovered from B6 virgin estrous mice (EST) or B6 mice on Day 3.5 pc after mating with intact, SVX, VAS, or SVX/VAS BALB/c males. Representative photomicrographs of FOXP3\(^+\) cells detected by IHC with FJK-16s antibody in uterine tissue at estrus (A) or on Day 3.5 pc after mating with intact males (B, C), and a negative control tissue with primary antibody omitted (D) are shown. A–D, bar = 100 μm. LE, luminal epithelium; Gl, endometrial glands; My, myometrium. Small arrows are FOXP3\(^+\) cells.

Nikon C1-Z confocal microscope and EZ-C1 version 3.2 software (Tokyo, Japan).

Statistics and Data Analysis

Normally distributed data were analyzed using one-way ANOVA and Tukey post hoc test using GraphPad Prism 5 software for Microsoft Windows (GraphPad software Inc., San Diego, CA; Microsoft, Seattle, WA). When data sets were shown by the Kolmogorov-Smirnov test to be not normally distributed, nonparametric Kruskal-Wallis and Mann-Whitney U-tests were performed using SPSS version 13.0 software for Windows (SPSS, Inc., Chicago, IL). Individual datum points were excluded as outliers if they were greater than 2 standard deviations from the mean. Relationships between mRNA transcript abundance were analyzed by bivariate correlation using Pearson correlation coefficient R. Data are displayed as means ± standard errors of the means (SEM). Differences in groups were considered significant if P values were <0.05.

RESULTS

Effect of Seminal Fluid on CD4\(^+\)FOXP3\(^+\) Cell Number in the Uterine-Draining Lymph Nodes

Recently we reported that hyperplasia in the para-aortic LN draining the uterus after mating combined with an elevation in the proportion of CD4\(^+\) cells expressing CD25 resulted in a greater than 2-fold increase in the number of CD4\(^+\)CD25\(^+\) cells [13]. We reported that 90% of the CD4\(^+\)CD25\(^+\) cells in para-aortic LN express FOXP3 [13], and other studies have shown that these CD4\(^+\)CD25\(^+\) cells exert immune-suppressive activity in vitro [2, 3], implying they are Treg cells; however, the CD4\(^+\)CD25\(^+\) subset may also contain newly activated T cells other than Treg cells, so the extent to which Treg cells comprise the expanded cell population has not been formally defined.

In this study, an anti-FOXP3 antibody and single tube cell quantification method were used to more precisely define para-aortic LN Treg cells as CD4\(^+\)FOXP3\(^+\) cells. Initially, flow cytometry analysis was performed with cells stained for surface CD4 and CD25 and then fixed, permeabilized, and stained with anti-mouse FOXP3 antibody FJK-16s. Of the total CD4\(^+\)CD25\(^+\) cell subset, approximately 80% of cells were colabeled with FJK-16s (Fig. 1A). Of the total FJK-16s reactive population, 86% of cells expressed both CD4 and CD25, 8% of cells were CD4\(^-\)CD25\(^+\), 4% of cells were CD4\(^+\)CD25\(^-\), and 2% of cells were CD4\(^-\)CD25\(^-\) (Fig. 1B). This staining distribution was similar in all estrous and mated mice evaluated (n ≥ 4 per group). To confirm that the antibody FJK-16s specifically labeled FOXP3\(^+\) cells, para-aortic LN cells from Foxp3\(^{GFP}\) mice were fixed and permeabilized and then labeled with FJK-16s. Of the cells that expressed the FOXP3-GFP fusion protein, 99% of cells were labeled by FJK-16s, whereas conversely, 95% of cells labeled by FJK-16s expressed the FOXP3-GFP fusion protein. Of the cells positive for either FOXP3-GFP expression or labeled by FJK-16s, 93.2% of cells were positive for both markers (Fig. 1C). These data demonstrate that positivity for FJK-16s staining is a reasonable marker of FOXP3 expression in para-aortic LN cells and show that despite substantial overlap between CD4\(^+\)CD25\(^+\) cells and CD4\(^+\)FOXP3\(^+\) cells, there is not complete identity between these two populations.

We then assessed the consequence of seminal fluid exposure at mating for the CD4\(^+\)FOXP3\(^+\) and CD4\(^-\)CD25\(^+\) cell populations in the para-aortic LN on Day 3.5 pc and, for comparison, included cells from the inguinal LN, which does not drain the uterus. Compared to virgin estrous animals, there was a 2.2-fold increase in the absolute number of CD4\(^+\)FOXP3\(^+\) cells in the para-aortic LNs of B6 females mated to intact BALB/c males (P < 0.011) (Fig. 1D). This was not seen in the inguinal LNs, where no significant differences in CD4\(^+\)FOXP3\(^+\) cell numbers were observed at Day 3.5 pc between intact-mated mice and virgin mice (Fig. 1E).

To assess the relative contributions of sperm and seminal plasma fractions of the ejaculate in expanding the CD4\(^+\)FOXP3\(^+\) cell population, we mated females with VAS,
SVX, or SVX/VAS BALB/c males. In the para-aortic LNs, expansion of the CD4<sup>+</sup>FOXP3<sup>+</sup> cell population was dependent on the seminal plasma but not on the sperm fraction of the ejaculate, as mating with seminal plasma-deficient (SVX or SVX/VAS) males resulted in no increase in the number of CD4<sup>+</sup>FOXP3<sup>+</sup> cells compared to those in virgin estrous animals (Fig. 1D). Conversely, mating with VAS studs resulted in an increase in CD4<sup>+</sup>FOXP3<sup>+</sup> cells (P 0.01) that was similar to that in intact-mated mice. In the distal inguinal LNs, there were no significant differences between any of the mated groups compared to virgin estrous mice (Fig. 1E). This implicates local events in the uterine tissue, rather than systemic endocrine events associated with mating, in driving expansion of the para-aortic LN CD4<sup>+</sup>FOXP3<sup>+</sup> cell population.

Among CD4<sup>+</sup>CD25<sup>+</sup> cells, the magnitude of change for each mating group relative to that of the estrous mice was similar or slightly greater than that seen in CD4<sup>+</sup>FOXP3<sup>+</sup> cells (Fig. 1, D and E). This is consistent with the considerable but not complete overlap between the two populations and is likely accounted for by the presence of activated T cells other than FOXP3<sup>+</sup> cells in the CD4<sup>+</sup>CD25<sup>+</sup> pool.

**Effect of Seminal Fluid on FOXP3<sup>+</sup> Cell Number and Foxp3 mRNA Expression in the Peri-Implantation Uterus**

Next, we investigated whether the increase in Treg cells in the para-aortic LNs was accompanied by an elevation in FOXP3<sup>+</sup> cells in the uterus on Day 3.5 pc. To address this, the density of FOXP3<sup>+</sup> cells in Day 3.5 pc uterine tissue recovered from B6 mice mated to either intact, SVX, VAS, or SVX/VAS males. In the para-aortic LNs, expansion of the CD4<sup>+</sup>FOXP3<sup>+</sup> cell population was dependent on the seminal plasma but not on the sperm fraction of the ejaculate, as mating with seminal plasma-deficient (SVX or SVX/VAS) males resulted in no increase in the number of CD4<sup>+</sup>FOXP3<sup>+</sup> cells compared to those in virgin estrous animals (Fig. 1D). Conversely, mating with VAS studs resulted in an increase in CD4<sup>+</sup>FOXP3<sup>+</sup> cells (P 0.01) that was similar to that in intact-mated mice. In the distal inguinal LNs, there were no significant differences between any of the mated groups compared to virgin estrous mice (Fig. 1E). This implicates local events in the uterine tissue, rather than systemic endocrine events associated with mating, in driving expansion of the para-aortic LN CD4<sup>+</sup>FOXP3<sup>+</sup> cell population.

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BALB/c males was analyzed using immunohistochemistry (IHC) and compared to that in virgin estrous mice. In tissue from estrous mice, FOXP3⁺ cells were sparsely distributed individually or in clusters throughout the endometrial tissue, with fewer cells in the myometrial tissue (Fig. 2A). Tissue from Day 3.5 pc mice mated with intact males contained more FOXP3⁺ cells, which were frequently clustered in the endometrial stroma, predominantly in the deeper regions and rarely located in the superficial stroma subjacent to the luminal epithelium or in the myometrium (Fig. 2A).

Quantification showed that in uterine tissue from mice mated with intact males, the density of cells expressing FOXP3 was 2.1-fold increased (P < 0.05) compared with that in estrous mice (Fig. 2E). Tissue from females mated with VAS males displayed a similar pattern of location (not shown) and comparable increase in density of FOXP3⁺ cells (2.1-fold; P < 0.05), whereas SVX- and SVX/VAS-mated mice showed no change in FOXP3⁺ cell number compared with estrous mice (Fig. 2E).

Real-time qPCR analysis showed a significant increase in the relative amount of Foxp3 mRNA in the uterine tissue of mice mated with intact males compared to that in virgin estrous mice (5.6-fold; P < 0.001). Smaller increases were seen in uterine tissue from females mated with VAS and SVX males (2.1-fold and 2.9-fold, respectively; both, P < 0.01), whereas there was no significant increase seen when females were mated with SVX/VAS males. Similar results were obtained for each of the two different primer sets for Foxp3 (data not

**FIG. 4.** Correlations between mRNAs encoding chemokines and chemokine receptors and Foxp3 mRNA in Day 3.5 pc uterine tissue are shown. Uterine tissue was recovered from B6 mice on Day 3.5 pc after mating with intact, SVX, or VAS BALB/c males, and mRNAs were measured by real-time qPCR as shown in Figures 2 and 3. Linear regression analysis was performed to assess the correlation between abundance of Foxp3 mRNA and abundance of (A) Ccl4, (B) Ccl5, (C) Ccl19, (D) Ccl22, (E) Ccr4, (F) Ccr5, (G) Ccr7, and (H) Itgae mRNAs in uterine tissue. Symbols represent the expression of Foxp3 mRNA (x axis) against the chemokine or chemokine receptor transcript of interest (y axis) for the same tissue sample. Circles, triangles, and squares represent data from the INT, SVX, and VAS groups, respectively. The relationship between transcript abundance was analyzed by bivariate correlation using Pearson correlation coefficient R. The coefficient of determination is indicated as the $r^2$ value, and P values are shown.
so data reported herein are from mean values for the two primer sets.

**Effect of Seminal Plasma on Uterine Expression of mRNAs Encoding Chemokines and Chemokine Receptors**

To investigate the possible role of chemokines in uterine accumulation of FOXP3+ cells in the preimplantation period, we then assessed expression of mRNAs encoding a range of chemokines and chemokine receptors in uterine tissue collected at estrous and at Day 3.5 pc after mating with intact, VAS, SVX, or VAS/SVX males. Chemokines shown previously to be involved in Treg cell migration in tissue sites other than the uterus were chosen for analysis. These included the Ccl4, Ccl5, Ccl19, and Ccl22 chemokine ligands and the Ccr4, Ccr5, and Ccr7 chemokine receptors. In addition, we measured the Itgae mRNA encoding the key Treg cell-retaining integrin CD103 [24].

The increase in Ccl19 expression was dependent on exposure to both seminal plasma and sperm, as expression was not increased when females were mated with VAS or SVX males (Fig. 3C). Conversely, increased Ccr5 expression was shown), so data reported herein are from mean values for the two primer sets.

**FIG. 5.** Immunolocalization and flow cytometric analysis of CCL19 expression in the uterus on Day 3.5 pc are shown. CCL19 protein was localized by IHC using anti-CCL19 antibody in sections of uterine tissue recovered on Day 3.5 pc from Foxp3GFP mice mated with intact males, enabling colocalization of CCL19 and FOXP3+ cells. Representative photomicrographs of CCL19 in uterine tissue (A, B, C) and a negative control tissue with irrelevant primary antibody (D) are shown. Bar = 50 μm for A and D; bar = 100 μm for B and C. LE, luminal epithelium; GI, endometrial glands. Large arrows are FOXP3+ cells (green staining) and CCL19 immunoreactivity is evident in luminal epithelium and glandular epithelium (red staining). CCL19+ cells were analyzed for coexpression of the epithelial cell-reactive lectin DBA or leukocyte markers CD45, EMR1 (F4/80), and ITGAM (CD11B) in uterine cell suspensions recovered from B6 females on Day 3.5 pc after mating to intact, VAS, or SVX males (E–K). Representative FACS plots show that the majority of CCL19+ cells recovered from females mated with intact (E), VAS (F), or SVX males (G) also reacted with DBA, with the number in each corner indicating the percentage of cells within that quadrant. The DBA/CCL19+ population was analyzed for CD45+ expression (H), and the CCL19+CD45+ population was analyzed for the macrophage markers ITGAM and EMR1 (I). The percentage of DBA+ epithelial cells expressing CCL19 (J) and their intensity of CCL19 expression (K) were quantified by flow cytometry. Data are the means ± SEM from n = 4–6 mice per group. Data were analyzed by Kruskal-Wallis and Mann-Whitney U-tests, and different letters indicate statistical significance between groups (P < 0.05).
related to Foxp3 mRNAs encoding migratory regulators might be causally related to both seminal plasma and sperm (Fig. 3, D and G).

Similarly there was no effect of sperm or seminal plasma Ccl22 expression of both fold, was also seen in tissue from mice mated with SVX males (1.8-fold, 0.05) and VAS males (2.2-fold, P 0.001) (Fig. 3F). Additionally, levels of Ccr4 and particularly Ilgae expression were both significantly correlated with Foxp3 expression (P < 0.001, r² = 0.38; and P < 0.001, r² = 0.63, respectively), and there was a trend toward a correlation between Ccr7 and Foxp3 expression (P = 0.068; r² = 0.13) (Fig. 4G).

Cellular Origin of Chemokine CCL19 in Uterine Tissue

Because uterine CCL19 was shown to be induced in early pregnancy and regulated by seminal fluid, we sought to investigate the cellular origin of CCL19 in uterine tissue and its spatial relationship with uterine FOXP3⁺ cells during the preimplantation period. IHC was utilized to detect CCL19 in fresh frozen uterine tissue from Foxp3GFP mice collected on Day 3.5 pc after mating with intact wild-type males, to allow colocalization of CCL19 and GFP-expressing FOXP3⁺ cells. CCL19 staining was evident at the apical and basal surface and to a lesser extent in the cytoplasm of most glandular and luminal epithelial cells in all (n = 6) uterine tissues examined, although there was spatial variation within individual tissues in the intensity of staining. FOXP3⁺ cells were present in the endometrial stromal tissue and were frequently located adjacent to CCL19⁺ glandular epithelial cells and to a lesser extent adjacent to luminal epithelial cells (Fig. 5, A–C).

We further investigated the identity of cells expressing CCL19 by using flow cytometry. Using the lectin DBA as a marker of epithelial cells, we confirmed that CCL19 is expressed by >80% of epithelial cells recovered from B6 females on Day 3.5 pc after mating with intact BALB/c males (Fig. 5D). A similar percent positivity for CCL19 expression was seen in DBA⁺ cells recovered from mice mated with VAS males (Fig. 5, E and I). In contrast, fewer DBA⁺ epithelial cells were positive for CCL19 on Day 3.5 pc after mating with SVX males (Fig. 5, F and I). The mean fluorescent intensity of CCL19 in DBA⁺ cells was higher in females mated with intact males than in females mated with VAS or SVX males (Fig. 5K). These results indicate that seminal plasma increases both the proportion of epithelial cells expressing CCL19 and the intensity of CCL19 expression.

Approximately 25% of the uterine cells expressing CCL19 were DBA negative. Flow cytometry confirmed that many of the DBA⁻ CCL19⁺ cells were CD45⁺ leukocytes (Fig. 5G), mainly ITGAM⁺EMR1⁺ DCs or ITGAM⁺EMR1⁺ macrophages (Fig. 5H).

Expression of the CCL19 Receptor CCR7 in Treg Cells

To determine whether Treg cells express the CCL19 receptor CCR7 in early pregnancy, B6 females were mated with intact BALB/c males, and, on Day 3.5 pc, LNs, spleen, uterus, and blood were collected for flow cytometry. Expression of CCR7 was clearly evident in CD4⁺CD25⁺FOXP3⁺ Treg cells collected from para-aortic and peripheral LNs, blood, and spleen, but of the small numbers of Treg cells detectable in the uterus, few expressed the CCR7 receptor (Fig. 6A). Within the para-aortic, mesenteric, cervical, and inguinal LNs and blood, the majority of Treg cells (50%–70%) were positive for CCR7; however, significantly fewer uterine Treg cells expressed CCR7 (mean of independent of seminal fluid exposure, and elevated expression was also seen in tissue from mice mated with SVX males (1.8-fold, P < 0.05) and VAS males (2.2-fold, P < 0.001) (Fig. 3F). Similarly there was no effect of sperm or seminal plasma deficiency on Ccl19 expression (Fig. 3B). In contrast, expression of both Ccl22 and Ccr7 mRNAs at Day 3.5 pc declined to below levels at estrus, unless females were exposed to both seminal plasma and sperm (Fig. 3, D and G).

To further examine whether changes in the expression of mRNAs encoding migratory regulators might be causally related to Foxp3 mRNA levels in uterine tissue from mated mice, we also analyzed correlations between Foxp3 mRNA and the chemokine and chemokine receptor mRNAs in uterine tissue from all groups of mated mice. This analysis revealed that the Ccl19 chemokine was significantly correlated with Foxp3 expression (P = 0.002; r² = 0.32) (Fig. 4C), while Ccr5 expression was not (P = 0.59; r² = 0.01) (Fig. 4F). Additionally, levels of Ccr4 and particularly Ilgae expression were both significantly correlated with Foxp3 expression (P < 0.001, r² = 0.38; and P < 0.001, r² = 0.63, respectively), and there was a trend toward a correlation between Ccr7 and Foxp3 expression (P = 0.068; r² = 0.13) (Fig. 4G).

FIG. 6. Expression of CCR7 in Treg cells in early pregnancy is shown. B6 females were mated to intact BALB/c males, and the percentage of CD3⁻CD4⁻FOXP3⁺ Treg cells expressing CCR7 was determined by flow cytometry in the para-aortic LN (PALN), mesenteric LN (MES), cervical LN (CLN), inguinal LN (ING), spleen (SPL), blood (BLD), and uterus (UTE). A) Representative plots of staining in para-aortic LN, blood, spleen, and uterus are shown, with the number in each corner indicating the percentage of cells within that quadrant. B) Data are the mean percentages ± SEM of CCR7⁺ cells from n = 5–7 mice per group. Data were analyzed by Kruskal-Wallis and Mann-Whitney U-tests, and different letters indicate statistical significance between groups (P < 0.05).
DISCUSSION

Prevention of deleterious immune responses against fetal antigens is critical for pregnancy success. Despite reduced antigenicity of conceptus tissue, maternal T cells still recognize and respond to fetal antigens [19, 25, 26], potentially as part of a biological "quality control" mechanism to assure conceptus health [27]. CD4\(^+\) CD25\(^+\) Treg cells have been identified as a T-cell subset that actively mediates maternal tolerance, and these cells are essential in the pre- and peri-implantation periods for establishing allogeneic pregnancy [2, 6], with insufficient Treg cells leading to implantation failure or miscarriage [6]. In this study we confirmed that exposure of the female reproductive tract to semen, specifically the seminal plasma component of the ejaculate, leads to a localized expansion of the FOXP3\(^+\) Treg cell population in the uterus-draining para-aortic LNs, and we extended this finding to show an associated increase in FOXP3\(^+\) Treg cells in the uterus at the time of embryo implantation. Additionally, we identified a role for seminal plasma-induced expression of the chemokine CCL19 in regulating uterine Treg cell accumulation prior to embryo implantation.

Exposure of the female reproductive tract to seminal fluid at mating can promote paternal antigen-specific hyporesponsiveness, ultimately inducing a state of transient active immune tolerance in the mother [13, 18, 28]. At least in part, this mechanism is mediated by Treg cells, which exert their actions at the time of embryo implantation [6, 13, 29]. TGFB and other factors in male seminal plasma, acting together with cytokine-related fluctuations in ovarian steroid hormones, have been implicated in eliciting increases in paternal antigen-reactive Treg cells in local LNs during the peri-conceptual period [13, 30].

Previous studies that analyzed Treg cells over gestation have generally relied on identifying Treg cells based on the coexpression of CD4 and CD25 [2, 3, 13, 31, 32]. However, CD25 expression on CD4\(^+\) cells is not an exclusive marker for Treg cells and also identifies activated CD4\(^+\) T cells [33], which could reasonably be increased in response to the antigenic stimulus of seminal fluid. Evidence of induction of other activation markers such as CD69 following insemination [14] brings into question whether elevations in the CD25\(^+\) population simply reflects an increase in activated T cells and limits the certainty of conclusions based on this identification strategy. Therefore, the hallmark Treg cell marker FOXP3 [8, 9] offers a superior marker for identifying Treg cells among the CD4\(^+\) T cell subset.

A population of CD4\(^+\) FOXP3\(^+\) cells was identified that predominantly express CD25, with more than 80% of the CD4\(^+\)CD25\(^+\) cell population also expressing FOXP3. However, a distinct population of CD4\(^+\)CD25\(^+\) cells did not express FOXP3, suggesting that at least some of these cells are activated T cells other than Treg cells. As well, there was an additional population of CD4\(^+\) FOXP3\(^+\) cells that did not express CD25. Critically, these FOXP3\(^+\)CD4\(^+\)CD25\(^+\) cells have also been shown to have suppressive function [34], and thus, we assert that definitive Treg cells in reproductive tissues are most precisely identified as CD4\(^+\) FOXP3\(^+\) cells. When cells were quantified according to the alternative classification phenotype of CD4\(^+\)CD25\(^+\) cells, the two data sets were comparable. This indicates that conclusions drawn in earlier work with Treg cells in early pregnancy, where the Treg cells are defined as CD4\(^+\)CD25\(^+\) cells, are probably valid.

In agreement with previous observations [13], on Day 3.5 pc, the total number of CD4\(^+\)CD25\(^+\) cells in the uterus-draining para-aortic LNs was greater than 2-fold increased following mating to intact males. A similar result was shown when Treg cells were defined as CD4\(^+\)FOXP3\(^+\) cells. Also consistent with earlier findings [13] was the observation that expansion of the Treg cell population following mating occurred only in the para-aortic LNs, with the inguinal LNs and spleen (data not shown) showing no significant increase in total Treg cells. Expansion was largely attributable to introduction of seminal plasma at the time of mating, confirming a critical role in FOXP3\(^+\) Treg cell expansion for the immunoreactive moieties within seminal plasma.

Treg cell numbers are higher during early pregnancy after allogeneic mating than after syngeneic mating [3]. Male antigen in the acellular fraction of the ejaculate [35] presented in the context of immunity-deviating factors such as TGFB [36] is likely to drive the postmating expansion in Treg cells in an antigen-specific manner. Experiments with mice expressing transgenic TCRs have shown that maternal antigen-presenting cells cross-present antigens from the ejaculate in the para-aortic LNs [19] and that seminal fluid-induced CSF2 (GM-CSF) facilitates this [22]. Observations that prevention of abortion in the CBA/J × DBA mating combination can be achieved if Treg cells are adoptively transferred from pregnant females, but not virgin females, are consistent with this hypothesis. Further support comes from the demonstration that paternal strain-specific tolerance of tumor cells is conferred by mating with intact but not SVX males, even in the absence of an ensuing pregnancy [13].

The largely contact-dependent mechanism of action of Treg cells dictates that their spatial distribution is critical to their suppressive function [37, 38]. For an optimal suppressive environment that would allow tolerance of the developing embryo, Treg cells would be required at the point of first contact between the conceptus and immune effector cells at implantation. Our observation of greater than 5-fold increase in Foxp3 expression and 2-fold increase in FOXP3\(^+\) cells in the endometrial stroma immediately prior to trophoblast cell penetration of the epithelial surface at implantation seems consistent with such a role. Seminal plasma appeared to be particularly important for achieving this increase in uterine Treg cells. This is thought to reflect the critical role for TGFB originating in the seminal vesicles for inducing Treg cells, an interpretation supported by experiments showing that exogenous TGFB delivered vaginally at mating can increase Treg cell numbers and alleviate abortion in CBA × DBA-mated mice [30]. A role for sperm cannot be discounted as Foxp3 mRNA expression was diminished after mating with vasectomized males, although IHC data did not support this. It is not clear why there was a difference in requirement for sperm in the RNA and protein analyses, but differences in per-cell expression of Foxp3 RNA depending on seminal fluid composition could account for this.

CD4\(^+\) FOXP3\(^+\) cells generated in the para-aortic LNs would expand the circulating pool available to home into the uterus. The para-aortic LN are likely to be a primary site of Treg cell production and control, as the absolute numbers of Treg cells in this tissue are approximately 10-fold greater than in the entire uterus on Day 3.5 pc. In this and previous studies [13], the para-aortic LN contained approximately 3 × 10^5 Treg cells on Day 3.5 pc. Based on the observed density of 7 FOXP3\(^+\) cells per mm^2 in uterine sections, we estimate the entire uterus contains approximately 4 × 10^4 Treg cells on Day 3.5 pc. Previous studies have shown that radiolabeled
para-aortic LN cells are recruited via the blood into the uterus in early pregnancy [14].

Like other T cells, the migration and trafficking of Treg cells into peripheral tissues is largely dependent on the local expression of chemokines and the presence of the cognate chemokine receptor on the surface of the Treg cell [39–41]. In the postmating period, there are coordinated changes in the level of several chemokines synthesized in the uterus, and this occurs in concert with the influx of a range of leukocytes including neutrophils, macrophages and DCs as well as T cells [42, 43].

In both the cycling nonpregnant uterus and the mid-gestation pregnant uterus (gestational Day 10.5), changes in chemokine mRNAs have been shown to occur in parallel with increased Foxp3 mRNA [44]. To investigate which chemokines were involved in Treg cell recruitment in the peri-implantation period, we assessed uterine expression of mRNAs encoding a range of chemokine ligands and receptors and the integrin CD103, which are implicated in recruiting Treg cells [24, 41, 44–46]. Of the migratory molecules analyzed, CCL19 (also known as the macrophage inhibitory protein 3β [MIP3β]) displayed a pattern of regulation most consistent with a role in Treg cell migration. Uterine Ccl19 mRNA was elevated during the preimplantation period and was responsive to both the sperm and seminal plasma components of the ejaculate. A functional contribution to uterine Treg cell recruitment is suggested by the positive correlation between Ccl19 and Foxp3 mRNA levels. Flow cytometry and immunohistochemistry confirmed the presence of CCL19, produced mainly by uterine and glandular epithelial cells, while macrophages and DCs also contributed to CCL19 production in the early pregnant uterus.

CCL19-facilitated T-cell migration is known to occur primarily as a result of its interaction with the receptor CCR7 [46]. The trend in correlation between Ccr7 and Foxp3 mRNA expression in the Day 3.5 pc uterus suggests this chemokine receptor is linked to Treg cell recruitment. CCL19 has been shown to induce migration of CD62L+ cells, with CD62L− Treg cells displaying both preferential migration toward CCL19 and enhanced suppressive capabilities [47]. The Cdh21 mRNA was not assessed in this study, but this potentially important subset of Treg cells warrants further investigation.

Under noninflammatory conditions, CCL19 is expressed almost exclusively in secondary lymphoid organs [48, 49]. CCL19 expression is also documented in peripheral tissues including the intestinal tract [50] and pancreas [51] and is critical for T-cell recruitment across the blood–brain barrier and into the central nervous system [52, 53]. Interestingly CCL19 is reported in the human uterus, where it is produced by luminal and glandular epithelial cells, as well as leukocytes in the endometrial stroma, with elevated expression in the late secretory phase, suggesting a role in controlling lymphocyte populations associated with implantation [54]. To date, a Ccl19 null mutant mouse has not been reported, but ectopic transgenic synthesis of CCL19 in the pancreas causes formation of lymphoid aggregates [55], raising the possibility of CCL19 involvement in formation of the organized lymphoid structures present in the endometrium [56].

The small percentage of Treg cells expressing CCR7 in the uterus is consistent with observations that activated T cells that migrate into other peripheral tissues show a reduced CCR7 expression [57]. This downregulation in CCR7 expression may be critical in ensuring that Treg cells are retained within the uterus once recruited. This notion fits with reports that null mutation in CCR7 expression disrupts T cell emigration and causes their aberrant accumulation in the skin [58]. Disruption of CCR7-mediated transit of Treg cells through peripheral tissues is associated with a range of autoimmune disorders, but the extent to which signaling by CCL19 and the alternate CCR7 ligand CCL21 contribute is not resolved [46].

Previous studies have implicated CCR5 expression by Treg cells and uterine expression of the CCR5 ligand CCL4 as promoting selective migration into the gravid uterus, although no deficiency in uterine Treg cells in Ccr5 null mutant mice implies redundancy with other Treg cell chemokine receptors [44, 59]. We showed no change in uterine Ccl4 expression in early pregnancy; however, elevated Ccr5 expression was evident, irrespective of seminal plasma exposure and so potentially attributable to the effects of ovarian steroid hormones. There was no correlation between Foxp3 and Ccr5 mRNAs, which is unsurprising as several leukocytes other than Treg cells also express CCR5, including macrophages, DCs, and granulocytes. Significant correlations between Foxp3 mRNA and Ccr4 and Il6g mRNA suggest these molecules might be involved in the migration or entrapment of Treg cells in the peri-implantation period, consistent with Treg cell regulation in other tissues [24, 60]. Other chemokines participate in Treg cell recruitment, and substantial redundancy in their actions is likely. After implantation, CXCL12 (SDF-1) expression in trophoblast is thought to attract Treg cells via their expression of CXCR4 receptors [61]. Further studies to investigate these molecules in the peri-implantation uterus will be informative.

Cumulatively, these results provide evidence for two mechanisms controlled by seminal fluid that act in synergy to achieve the increase in uterine FOXP3+ Treg cells required for embryo implantation. First, seminal plasma elicits an increase in generation of Treg cells within the uterus-draining para-aortic LNs, resulting in greater availability of Treg cells for recruitment into the uterus. Second, seminal fluid-induced up-regulation of CCL19 in glandular epithelial cells would selectively promote migration of Treg cells from the circulation into the endometrium. A third mechanism might involve proliferation of recruited Treg cells in situ, and a recent report using the proliferation marker Ki67 suggests that local proliferation does indeed contribute to the uterine FOXP3+ cell population [6].

In summary, this study provides new insights into the role of seminal fluid in facilitating pregnancy through regulating FOXP3+ Treg cell accumulation in the implantation site. Given the critical role that Treg cells play in establishing pregnancy [4], a better understanding of the biological processes generating immune tolerance at embryo implantation will ultimately contribute to developing new diagnostics and treatments for implantation failure, miscarriage and other pathologies of pregnancy where an immune etiology is implicated.

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