Role of Tonsillar Chronic Inflammation and Commensal Bacteria in the Pathogenesis of Pediatric OSA

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Immune responses at the boundary between the host and the world beyond are complex and mucosal tissue homeostasis relies on them. Obstructive sleep apnea (OSA) is a syndrome suffered by children with hypertrophied tonsils. We have previously demonstrated that these tonsils present a defective regulatory B cell (Breg) compartment. Here, we extend those findings by uncovering the crucial role of resident pro-inflammatory B and T cells in sustaining tonsillar hypertrophy and hyperplasia by producing TNFα and IL17, respectively, in ex vivo cultures. Additionally, we detected prominent levels of expression of CD1d by tonsillar stratified as well as reticular epithelium, which have not previously been reported. Furthermore, we evidenced the hypertrophy of germinal centers (GC) and the general hyperplasia of B lymphocytes within the tissue and the lumen of the crypts. Of note, such B cells resulted mainly (IgG/IgM)+ cells, with some IgA+ cells located marginally in the follicles. Finally, by combining bacterial culture from the tonsillar core and subsequent identification of the respective isolates, we determined the most prevalent species within the cohort of OSA patients. Although the isolated species are considered normal oropharyngeal commensals in children, we confirmed their capacity to breach the epithelial barrier. Our work sheds light on the pathological mechanism underlying OSA, highlighting the relevance taken by the host immune system when defining infection versus colonization, and opening alternatives of treatment.

Keywords: tonsils, oral bacteria, obstructive, sleep, apnea, mucosal immunity
INTRODUCTION

Human paired palatine tonsils are lymphoid epithelial tissues of the oral mucosa around the oropharynx. They are part of the Waldeyer’s ring of lymphoid tissue, which also comprise the adenoids and the lingual tonsils. The palatine tonsils (tonsils, from now on) are strategically located to generate mucosal immunity as they are constantly exposed to dietary and airborne antigens (Ags). Moreover, they have evolved for direct transport of foreign material from the exterior to the lymphoid cells through deep and branched crypts and the absence of Ag-degrading digestive enzymes (1). The surfaces of the human body, including the oropharynx, are colonized by several microbes, mainly bacteria, that establish a mutualistic relationship with the host. Tonsils are predominantly B-cell organs, immunologically most active between 4 and 10 years old. Some children (also some adults, but were not included in this study) present tonsillar hyperplasia and hypertrophy for yet unknown reasons. Such enlargement is the major pathophysiological sign underlying OSA, a highly prevalent disease, recognized as a major public health burden. OSA is characterized by repeated events of partial or complete upper airway obstruction during sleep that lead to disruption of normal ventilation with all the consequences implied due to hypoxemia. An increment in pro-inflammatory cytokines have been reported in blood from OSA patients (2–4). It is assumed that the B cell hyperplasia and hypertrophy that cause OSA are the result of chronic inflammation of palatine tonsils in children.

B cells contribute to immune responses during infectious, inflammatory and autoimmune diseases. In the last two decades, we have learned that B cells are able to modulate physiological and pathological processes not only by producing antibodies (Abs) and presenting Ags but also by producing cytokines. It has been shown that B cells can occur in the form of several cytokine-secreting subsets with either pro- or anti-inflammatory functions (5). Being an important reservoir of human B cells, the tonsils serve as a platform to study such subsets. Within this framework, we have recently demonstrated that OSA tonsils rendered significantly lower percentages of interleukin 10 (IL10) producing B cells (Bregs) than tonsils excised due to recurrent tonsillitis, showing that Bregs have a more complex and interesting role in tonsillar disease than was hitherto appreciated. Moreover, such defect in Breg population correlated with an increment in the proportion of germinal center (GC) cells. This correlation reveals a role for the Breg subset in controlling GC reactions, as it is by means of GC cells that tonsils are hypertrophied (6). GCs are the microanatomical sites within secondary lymphoid organs, critical for memory B and plasma cell generation. There are also many other cellular and molecular players involved in controlling GC activity. For instance, it has been long known that the pro-inflammatory cytokine tumor necrosis factor alpha (TNFα) is a required autocrine B-cell growth factor (7).

In the present paper, we found that tonsillar mononuclear cells (TMC) from OSA tonsils effectively exhibit a pro-inflammatory cytokine profile rapidly in culture. Interestingly, under certain stimulating conditions, B lymphocytes became one of the main cell populations driving TNFα levels in culture ex vivo. On the other hand, tonsillar interleukin 17 (IL17) was produced primarily by CD4+ T cells. At tissue level, we discovered CD1d expression by tonsillar stratified as well as reticular epithelium and corroborated the persistent hypertrophy of GC and the concomitant hyperplasia of B lymphocytes, prevalently IgG/IgM positive (IgG/IgM)+. We identified a number of bacterial species in the tonsillar core of patients with tonsillar hypertrophy, considered normal oropharyngeal commensals in children. However, we confirmed their presence beyond the epithelial boundaries by fluorescence in situ hybridization (FISH). Thus, our observations suggest that tonsillar hypertrophy is a multifaceted condition not associated to the presence of a particular microorganism but more likely to a failure of normal immune homeostatic mechanisms caused by the local loss of the capacity to discriminate between commensals and pathogens of the host. All in all, when such discrimination is lost, commensals become pathogens. Therefore, we support the notion that OSA in children is of infectious nature, clearly not associated to a single species.

MATERIALS AND METHODS

Isolation of cells

Primary human mononuclear cells were isolated from tonsils obtained from patients (total n=54 taking into account all experiments, the particular number of samples per experiment were detailed in the corresponding Figure legends) undergoing tonsillectomy due to OSA. TMC were prepared as follows. Briefly, tonsils were collected in phosphate buffered saline (PBS) buffer containing 50 µg ml⁻¹ amphotericin B (Richet, BA, Arg). Tissues were chopped with a scalpel in IMDM medium (see below) and passed through a 70 µm-pore-size cell strainer (Falcon, Thermo Fisher, BA, Arg). TMC were purified by density gradient centrifugation with Ficoll-Hypaque (GE Healthcare, Uppsala, Sweden). The viability of primary cells, as determined by trypan blue exclusion was greater than 99% in all preparations. Informed consent was obtained from subjects before the study. The institutional ethics committee (Clinical Hospital, School of Medicine, Buenos Aires) approved the collection and use of clinical material, conforming to the provisions of the Declaration of Helsinki (as revised in Edinburgh 2000). Informed consent was obtained from all participants and/or their legal guardian/s. FACS experiments were performed with freshly isolated cells only.

Cell culture

TMC were cultured in IMDM medium (Life Technologies, CA, USA) containing 10% heat-inactivated fetal calf serum, 2mM L-glutamine, 100 U/ml penicillin, 100 µg/ml streptomycin, 20 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid buffer (HEPES), 1 mM sodium pyruvate and 50 µM 2-mercaptopethanol (all from Invitrogen, CA, USA). Human IL2 (20 ng/ml; R&D Systems, MN, USA) and human IL4 (20 ng/ml; R&D Systems, MN, USA) were added immediately before experiments as supplements. When indicated, human recombinant hCD40L (250 ng/ml; R&D Systems, MN, USA) and 25 µM CpG-ODN 2006 (InvivoGen, CA, USA) were used. Cells were cultured at
1x10^6 cells/ml either in 24-well culture plates (1ml) or 48-well culture plates (0.5ml).

**Antibodies and fluorescence-activated cell sorting (FACS)**

Fluorochrome conjugated mAbs specific for human CD3 (Pacific Blue, clone UCHT1), human CD20 (FITC, clone L27 and APC H7 clone 2H7), human CD4 (PerCP, clone SK3), human IL17A (APC, clone BL168), TNFα (PE, clone IT-5H2 and BV711, clone Mab11), and respective isotype control mAbs were purchased from BD Biosciences (CA, USA) and Biolegend (CA, USA). Fixable viability dyes were used either eFluor 780 (eBioscience, CA, USA), Zombie Green (Biolegend, CA, USA) or Zombie Aqua (Biolegend, CA, USA) depending on the staining scheme. To detect intracellular cytokines in cultured cells, the latter were stimulated with PMA/Ionomycin/Brefeldin A for the last 5 hs of culture. Then, they were incubated with Cytofix/Cytoperm (BD PharMingen, CA, USA) for 20 minutes (min) in the dark and washed with Perm/Wash solution (BD PharMingen, CA, USA). Following permeabilization, the cells were stained with the respective anti-cytokine mAb. Cells were acquired using FACSaria II (BD Biosciences, CA, USA) and analyzed with FlowJo software (Treestar, OR, USA). Single stained controls were used to set compensation parameters. Fluorescence minus one and isotype-matched Ab controls were used to set analysis gates.

**Statistical analyses**

The results were analyzed using GraphPad Prism 7.0 software (Graph Pad Inc, CA, USA). The normality of variable distribution was assessed when p > 0.05. Once the hypothesis of normality was accepted, comparisons were performed using unpaired Student t test. When normality resulted rejected, a Mann-Whitney test was used. In both cases, the null hypothesis being that 2 populations do not differ.

**Bacterial cultures**

Tonsillar samples were cultured on Columbia agar containing 5% sheep blood and chocolate agar (Laboratorios Britania, Argentina) at 37°C with 5% CO2 for 24 to 48 h. Anaerobes were cultured onto Brucella agar supplemented with hemin and vitamin K under anaerobic conditions. All isolates were identified by MALDI-TOF MS. The isolates were identified by the direct colony on plate extraction method as previously described (8). Mass spectra were acquired using the MALDI-TOF MS spectrometer in a linear positive mode (Microlens, Bruker Daltonics). Mass spectra were analyzed in a m/z range of 2,000 to 20,000. The MALDI Biotyper library version 3.0 and MALDI Biotyper software version 3.1 were used for bacterial identification.

**Immunofluorescence**

Cryostat sections (5-10 µm thickness) of tonsils were fixed and stained with mouse anti-human CD1d (clone NOR3.2/13.17 Santa Cruz Biotechnology, CA, USA) followed by chicken anti-mouse IgG antibody AlexaFluor 488 (Thermo Fisher Scientific, BA, Arg.). (IgG/IgM)’ cells were detected by addition of goat anti-human IgM+IgG (H+L) F(ab’)2 (Jackson Immunoresearch, PA, USA) followed by donkey anti-goat IgG AlexaFluor 594 (Jackson Immunoresearch, PA, USA). IgA’ cells were detected by addition of goat anti-human IgA F(ab’)2 (InVivoGen, CA, USA) followed by anti-goat IgG AlexaFluor 594 (Jackson Immunoresearch, PA, USA). Cell nuclei were visualized with 4,6-diamidino-2-phenylindole staining (DAPI, Thermo Fisher Scientific, BA, Arg.). Finally, slides were rinsed with phosphate buffered saline, air dried, mounted in Vectashield (Vector Labs, CA, USA) and sealed with a glass coverslip. Samples were examined with a Nikon Eclipse Ti-E fluorescence microscope (Nikon instruments Inc, Tokyo, Japan). Corresponding isotype controls were always added (see Supplementary Information).

**Bacterial localization and immunostaining**

A solution of 0.5 mg/ml of lysozyme (Sigma Aldrich, Darmstadt, Germany) in 0.1 M Tris-HCl and 0.05 M Na2EDTA was added to the cryosections, followed by incubation at 37°C for 3 h. Fixed, permeabilized tonsillar sections were then incubated in a moist chamber for 4hs at 48°C in hybridization buffer [0.9 M NaCl, 20 mM Tris-HCl (pH 7.6), 0.01% sodium dodecyl sulfate, 30% formamide] containing either the universal probe EUB388 AlexaFluor 488 labeled probe or the negative control (nonsense) NONEUB388 AlexaFluor 546 (Thermo Fisher Scientific, BA, Arg.). Stringent washing was performed by incubating the slide in washing buffer [20 mM Tris-HCl (pH 7.6), 0.01% sodium dodecyl sulfate, 112 mM NaCl] at 48°C for 15 min in a moist chamber a number of times, and subsequently rinsed with ddH2O for 5min. Finally, probed-hybridized tonsil cryosections were subjected to immunofluorescence staining as described above to determine the distribution of bacteria within the different tissue compartments. Images were acquired with a Nikon Eclipse Ti-E microscope (Nikon Instruments Inc., Tokyo, Japan) or an Olympus FV1000/IX81 confocal microscope (Olympus Corporation, Tokyo, Japan) using an oil-immersion objective (60X; numerical aperture, 1.42).

**RESULTS**

**Characterization of TNFα Production by B Cells From Hypertrophied Tonsils**

It is long established that B-cell–derived TNFα plays a crucial role in the development of follicular dendritic cells (FDCs) and B-cell follicles in spleen (9–11). TNFα and IL10 are cytokines with antagonistic actions. Interestingly, tonsils from OSA patients present a defective Breg compartment which correlates with higher proportion of GC B lymphocytes [BGC (6)], larger GC (12), and increased numbers of T follicular helper cells (Tfh) (13) than tonsils excised by other pathologies. Taking all into account, it would be expected that OSA tonsils exhibit a significant lymphocyte-derived TNFα compartment. We examined TNFα expression at the single cell level by FACS, upon TMC culture. A set of cultures was treated for 24 hs with IL2 and IL4 (IL2+IL4), which promote survival of all lymphocyte subsets through slight stimulation (14). Another set of cultures was supplemented for 24
hs and 48 hs with CpG and CD40L (in addition to IL2 and IL4, IL2+IL4+CD40L+CpG) to target stimulation towards B cells, aiming to assess specifically their capacity to contribute to the tonsillar pool of TNFα. Single cells were selected based on FSC-A vs FSC-H analysis (Singlets gate, Figure 1A). The cells that were already dead prior to permeabilization were excluded from the analysis through a fixable dead cell staining (Viable gate, Figure 1A and Supplementary Figure 1). In agreement with previous observations (14), elevated levels of cell death were linked to stimulation of tonsillar cultures (Figures 1A, B), dominated by a variety of terminally differentiated and highly activated B cells. In fact, we used this decay in the viability of the TMC cultures to monitor for suitable activation. In our experience, TMC culture stimulation is quite susceptible to minimal changes in experimental conditions (cellular density or FCS and activation cocktail batch, for instance). Therefore, when comparing cytokine secretion within a cohort of patients, those cultures that did not present such evolution in terms of viability, proportion of CD3+ and CD20+ populations and CD20 down-modulation (Figure 1A), were not considered. Also, CD3+ cells served as an internal control for cytokine expression (Figure 1C).

As expected, we observed that TMC from hypertrophied tonsils produced considerable amounts of TNFα when stimulated. At 24 hs post stimulation, approximately one third (29% ± SD 22%) of the cells from the IL2+IL4 stimulated cultures, expressed TNFα and that percentage reached ~40% (37%± SD 16%) in CD40L+CpG+IL2+IL4 stimulated cultures. The latter cultures were particularly affected by the surface CD20 down-modulation which takes place in response to general stimuli (15), albeit more pronounced with CD40L stimulation, as it has been extensively described previously (6, 16). Interestingly, B cells (CD20+) expressing TNFα were those presenting lower levels of CD20 (CD20 down TNFα+ cells, Figure 1A) suggesting that TNFα expression might be another functional consequence of CD20 modulation and the downstream signaling pathways triggered post internalization. These findings confirmed that the extent of CD20 down-regulation correlates with the degree of B cell activation (Figures 1A, C) (6, 15, 16). Despite the fact that lymphoid populations other than CD20+ and CD3+ cells in TMC are negligible (Figure 1A, non-cultured panel, double negative cells), their putative expansion, under the culture conditions used, needed to be tested as they would represent a putative contaminant of the CD20 down cells. Tonsillar dendritic cells (tDC) identified as CD20 CD3 CD11c+ appeared to increase their percentage within the CD20 CD20 down gate under stimuli. Even so, they represented 0.4%-0.7% of the latter, depending on the stimulus (Supplementary Figure 2A). Moreover, we could not detect any CD11b+ cells among the cultured TMC (Supplementary Figure 2B). The proportion of innate lymphoid cells (ILC) was also assayed (Supplementary Figure 3) and proved to be negligible as well (detailed in the next section).

In order to compare TNFα expression by CD20+/down population with that of CD3+ control cells within the same sample, we measured the median fluorescence intensity (MFI) for TNFα+ cells on both populations (Supplementary Figure 4). Regardless of the type and length of stimulation, average MFI for CD3+ cells resulted around twice the same parameter for CD20+/down cells (Figure 1D). We next determined in which proportion CD20+/down and CD3+ cells were contributing to the total TNFα+ population (Figure 1E). Comprehensively, at 24 hs post stimulation with CD40L+CpG+IL2+IL4, when B cell activation peaked, CD20+/down cells represented the majority of tonsillar TNFα+ cells (52.4% ± SD 13.4% CD20+/down cells vs 41.7% ± SD 12.8% CD3+ cells). On the other hand, at 48 hs post stimulation with CD40L+CpG+IL2+IL4, when tonsillar B cells showed strongly reduced viability, tonsillar TNFα+ population was dominated by CD3+ cells (30.0% ± SD 11.7% CD20+/down cells vs 63.0% ± SD 15.9% CD3+ cells). The cultures stimulated for 24 hs only with IL2+IL4 presented a tendency towards a slightly higher CD20+/down cells contribution to the TNFα+ pool than CD3+ cells (Figure 1F). Importantly, the ability of tonsillar B cells to express TNFα was proved to be independent on the presence of the T cells in the TMC sample (Supplementary Figure 5). Finally, we also tried stimulating the TMC cultures using as a surrogate antigen F(ab')2 goat antibodies specific for IgM and IgG (in addition to CD40L+IL2+IL4) (14), obtaining very similar results to those of CD40L+CpG+IL2+IL4 stimulation (data not shown).

Kim et al. (17) have previously reported that upon culture, TMC from hypertrophied tonsils rendered higher levels of TNFα in the supernatant than TMC from children with recurrent tonsillitis. These results extend such findings, indicating that being exposed to various stimuli, B cell population can largely contribute to the TNFα pool, supporting (if not promoting) tonsillar inflammation.

Presence of Inflammatory T Cells Co-Expressing IL17 and TNFα in Hypertrophied Tonsils

It has been suggested that development of OSA is associated with peripheral Th17/Treg imbalance and is characterized by a pro-inflammatory cytokine microenvironment (2). We have previously reported on the ability of tonsillar B cells to secrete IL17 (6). However, while B cells as a population can dominate tonsillar TNFα production, we found IL17 secretion is indeed controlled by CD3+ cells at every culture condition tested, including those specifically stimulating B lymphocytes (Figures 2A, B). TMC cultures supplemented with IL2+IL4 as well as those supplemented with CD40L+CpG+IL2+IL4 rendered an IL17+ population which comprised ~90% CD3+CD4+IL17+ cells at 24 and 48 hs. We confirmed this observation by measuring cytokines on the supernatant of some of the cultures by multiplex biomarker immunoassay (Luminex). We tried 4 different conditions to specifically stimulate B cells (CpG; CD40L; anti-(IgM/IgG)+CpG; anti-(IgM/IgG)+ CD40L), all of them in addition to IL2+IL4. We also added a culture treated only with IL2+IL4, as control. IL17 concentration did not significantly varied with culture condition compared to the control, as opposed to that of TNFα, which exhibited significant increments under B cell stimulating conditions (data not shown). As expected, CD3+IL17+ cells were confirmed to be CD3+CD4+IL17+ (Th17) cells (Figure 2B). Moreover, we tried to estimate the putative contribution of cells other than Th17 and B
FIGURE 1 | TNFα expression by tonsillar mononuclear cells. (A) Freshly isolated TMC were cultured on IL2+IL4 alone or CpG+CD40L+IL2+IL4, for the time points indicated on the top of each panel. Non-cultured cells served as control (conserved at 4°C). Cells were stained for surface CD20, CD3 and intracellular TNFα. Samples were subsequently analyzed by FACS. Gating strategy is partially illustrated. Singlets were gated by plotting FSC-H vs FSC-A for each sample (upper panels). Within the singlets population, dead cells were determined by a fixable viability dye. In this case, the viable gate corresponds to the eFluor 780-cells (middle panels). Within the viable gate, lymphoid gate was determined through SSC-A vs FSC-A (not shown). Lower panels: CD20 vs CD3 contour plots. Percentages designate frequencies of the populations indicated. Dashed red lines show the decline of expression of CD20 post stimulation. Data from one experiment representative of 10 independent experiments performed with different individuals each one, are presented. Gates were manually adjusted due to the changes experienced by the cells in culture upon each treatment. (B) Line graph plotting the mean percentage ± SD of the frequencies of the viable gate pooled from the 10 independent experiments performed with stimulated TMC and detailed in (A). (C) TMC were cultured under the same conditions and stained for the same molecules as in (A). Gating strategy as in (A) (singlets-viable-lymphoid, not shown). Contour plots show TNFα and CD20 expression by TMC. Percentages designate frequencies of populations in each quadrant. CD20+/down cells appear highlighted in red, based on the gating performed on the corresponding CD20 vs CD3 graph as in (A), using the backgating option, to exclude any CD3 contamination. Quadrants were manually adjusted due to the changes experienced by the cells in culture upon each treatment. Data from one experiment representative of 10 independent experiments performed with different individuals each one, are presented. (D) Graph showing the MFI fold increment=(TNFα MFI of CD3+TNFα+/TNFα MFI of CD20+/down TNFα+) for each independent experiment at every stimulating condition. Calculations were made following FACS analysis depicted in Supplementary Figure 4. (E) Gating strategy used to estimate B and T cell specific contribution to the TNFα+ cell pool from the 10 samples analyzed as shown in (A–C). P1 denotes percentage of TNFα+ cell population determined by TNFα vs SSC-A contour profile under the stimulations described above. Panel on the right, representative contour plots for CD20+/down (red) and CD3+ (blue) populations within the P1 gate. (F) Histograms presenting the mean percentage ± SD of the CD3+ and CD20+/down/+ cell population frequencies from 10 independent experiments within the TNFα+ cell pool for each stimulating condition, calculated as shown in (E). p value was calculated through unpaired t test, normality was confirmed by the Shapiro-Wilk test.
lymphocytes, to the IL17+ cell population pool. ILC3 are characterized by their capacity of secreting IL17 when stimulated. ILC3 represented around 0.03% of the TMC lymphoid gate (P3, Supplementary Figure 3). Their negligible fraction precluded us from detecting whether the culture conditions used stimulated them to express IL17. The same applied to Natural Killer (NK) cells which collectively signified 0.2% of the TMC lymphoid gate (P3, Supplementary Figure 3) but only a sub-fraction would be able to express IL17.

Th17 cells have appeared to exhibit plasticity of function and often co-produce other pro-inflammatory cytokines, particularly at sites of inflammation. Hence, we investigated whether tonsillar Th17 also expressed other cytokines as TNFα and interferon γ (IFNγ). The pseudo-color dot plots shown in Figure 2C were

**FIGURE 2** | Tonsillar Th17 cells co-express TNFα and IL17. (A) Freshly isolated TMC were cultured on CpG+CD40L+IL2+IL4 for 24 hs. Samples were subsequently analyzed by FACS. Gating strategy is partially illustrated. Singlets were gated by plotting FSC-A vs FSC-H for each sample (not shown). Within the singlets population, dead cells were determined by a fixable viability dye (see Methods). Within the viable gate, lymphoid gate was determined through SSC-A vs FSC-A (not shown). Cells were stained for surface CD20, CD3 and intracellular IL17. Pseudo-color plots show CD3 and CD20 expression by TMC upon stimulation, percentages designate frequencies of populations in each gate (left panel). Upper right panel: dot plot shows IL17 and CD20 expression by TMC, CD20+IL17+ population is highlighted in black and its frequency within CD20+ population is indicated as percentage. Lower right panel: dot plot shows IL17 and CD3 expression by TMC, CD3+IL17+ population is highlighted in black and its frequency within CD3+ population is indicated as percentage. Data from one experiment representative of 10 independent experiments performed with different individuals each one, are presented. (B) Gating strategy used to estimate the populations contributing to the IL17+ cell subset upon stimulating strategy as in (A). Singlets were gated by plotting FSC-A vs FSC-H for each sample (not shown). Within the singlets population, dead cells were determined by a fixable viability dye (see Methods). Within the viable gate, lymphoid gate was determined through SSC-A vs FSC-A (not shown). P1 denotes IL17+ cell population determined by IL17 vs SSC-A contour profile. Panel on the right, representative contour plots for CD20+ and CD4+ populations within the P1 gate. Percentages designate frequencies of the populations indicated relative to P1 gate. Data from one experiment representative of 6 independent experiments performed with different individuals each one, are presented. (C) Freshly isolated TMC were cultured on IL2+IL4 alone or CpG+CD40L+IL2+IL4, for the time points indicated on the top of each panel. Singlets were gated by plotting FSC-A vs FSC-H for each sample (not shown). Within the singlets population, dead cells were determined by a fixable viability dye (see Methods). Within the viable gate, lymphoid gate was determined through SSC-A vs FSC-A (not shown). Cells were stained for surface CD20, CD3, intracellular TNFα and IL17 and finally analyzed by FACS. Upper panels: pseudo-color plots show IL17 vs CD3 expression by TMC. They correspond to events within singlets, viable, lymphoid and CD3+ gates. Percentages designate frequencies of IL17+ cells within CD3+. Lower panels: pseudo-color plots show TNFα and IL17 expression by TMC, gated as in the upper panels. Percentages designate frequencies of populations in each quadrant, relative to the CD3+ gate. Data from one experiment representative of 12 independent experiments performed with different individuals each one, are presented. (D) Histograms presenting the mean percentage ± SD of the Th17 TNFα+ cell population frequencies from 12 independent experiments for each stimulating condition, calculated as shown in (C), top right quadrant, p calculated through a Mann-Whitney test as the distribution of data in the middle column was not normally distributed.
gated on CD3+ cells and show CD3 vs IL17 (upper panel) and IL17 vs TNFα (lower panel) staining. We detected co-expression of TNFα in a significant fraction of the Th17 cell population upon all culture conditions tested. At 24 hs post stimulation, nearly half of the Th17 population (44% ± SD 14%) from the IL2+IL4 stimulated cultures co-expressed TNFα. This percentage significantly increased when TMC were cultured with CD40L+CpG+IL2+IL4 for 24 hs (59% ± SD 14.5%) as well as for 48 hs (60%± SD 18%) indicating a positive correlation between higher frequencies of TNFα and IL17 and TNFα co-expression (Figure 2D). Low levels of expression of IFNγ observed upon the stimulating conditions tested precluded us from definitive conclusions regarding co-expression of IFNγ and IL17.

We concluded that tonsillar IL17 was produced primarily by Th17, which largely coproduced TNFα. Interestingly, several studies have linked the Th17 pathway with formation of GCs in mice spleens and ectopic B cell follicles at sites of inflammation (18–20). On the whole, TMC from hypertrophied tonsils promptly exhibited a pro-inflammatory cytokine profile in culture.

The Complex Histological Milieu of Tonsils and the Potential B Cell Contribution to Their Tissue Specific Inflammation

Like all mucosae, tonsillar immune actions (anti- and pro-inflammatory) must be tightly regulated, to balance the protection against virulent germs and the tolerance to harmless flora and innocuous Ags entering with air and food. Collectively, the results described above and others we have previously published (6, 14) are in agreement with the notion that tonsillar hypertrophy is a result of an imbalance between regulatory and effector immune functions which in turn lead to a local chronic inflammation. To this point, we have examined isolated cells. However, in vivo, immune processes occur in a complex tissue context which has to be comprehended. In order to recognize the histologic compartments within tonsils, we performed immune-fluorescence staining on cryosections from excised tonsils. The palatine tonsils are secondary lymphoid organs belonging to the mucosal associated lymphoid tissue (MALT), with unique histological characteristics. Unlike the lymph nodes, tonsils do not have afferent lymphatics, but present deep and branched crypts. Ag sample occurs through their distinctive tissue architecture and this is why tonsils are called lympho-epithelial organs. The tonsillar epithelium provides protection as well as serving to transport foreign material from the lumen to the lymphoid compartment. To evidence this mucosal surface, we tested the tonsillar epithelial cells’ reactivity to antibodies against CD1d, since it has been reported its expression in other mucosal epithelial cells, like intestinal (21), epidermal keratinocytes (22) and vaginal epithelium (23). We confirmed CD1d expression in tonsillar epithelium. Anatomically, the surface epithelium of the palatine tonsils is an extension of the stratified squamous epithelium of the oral mucosa (Figure 3A, Supplementary Figures 6, 7). Importantly, in the crypts, the stratified epithelium becomes narrower and laxly textured [reticulated epithelium as termed by Olahe in 1978 (24)] and highly infiltrated with lymphocytes (Figures 3B–D). We observed

CD1d expression in reticulated epithelium as well as in stratified squamous epithelium. Nevertheless, the level of that expression did not appear uniform at all levels of the multilayered squamous epithelium. The basal cells, which maintained an orientation perpendicular to the basement membrane, presented negligible immunoreactivity to anti CD1d. Conversely, CD1d was expressed strongly in the epithelial cells of the supra-basal and intermediate layers, significantly declining its expression in the apical layers. A similar pattern of CD1d staining is observed in the vaginal stratified epithelium (23). Expression was restrained mostly to the cell membrane in all cases (Figure 3). This epithelium was supported by a layer of connective tissue (Figure 3A) that separates it from the lymphoid component, and resulted negative for CD1d, with the exception of sporadic infiltrating lymphoid cells. Figures 3B, C show a region of crypts lined by reticulated epithelium. In fact, the crypt in Figure 3C, is lined by reticulated epithelium on one side and stratified squamous epithelium on the other. Two characteristics of the former are clear from the figures, and these are the desquamation of the upper cell layers and the absence or disruption of the basal layer of the epithelium. In this case, there is no boundary between the epithelium and the underlying lymphoid tissue. The cells in the intermediate layer were distorted and often separated from one another by the invading lymphoid cells, yet again CD1d exhibited high expression in this layer. The reticular epithelium trailed the curving shapes of the underlying follicles. To our knowledge, this is the first report on CD1d expression by tonsillar epithelium.

To visualize the lymphoid compartment, we stained for (IgM/IgG)+ B cells (Figure 3). Our results confirmed previous reports (25). (IgM/IgG)+ B cells (and plasma cells, distinguishable by their size) appeared located not only in follicles, GC and their mantle zone, but also uniformly scattered throughout the tonsillar tissues (Figures 3A–F), generally associated in clusters when infiltrating reticular epithelial tissue (Figure 3D). IgA+ B/plasma cells on the other hand, were mainly found outside the germinal centers, especially in the epithelium of crypts and rarely appear in the GCs (Figure 3F). Also, it can be observed the protrusion of lymphocytes and plasma cells from the sites of desquamation to the lumen (Figures 3C, D). IgG and IgM can be observed in its soluble form, accumulated on the apical surface of the epithelial layer (Figures 3A, D). It has been previously shown that nasopharyngeal epithelium widely expresses FcRn, which mediates IgG transport across the epithelial cell layer (26). On the other hand, it does not express pIgR, hence the transport of dimeric IgA is unfeasible through this kind of epithelia (27), making IgG the predominant protective isotype in tonsils. The histological findings we described above are in agreement with those reports. Moreover, FcRn and CD1d belong to the same family of proteins, the non-classical MHC class I molecules and both have been shown to display a restriction in their expression to specific tissues, namely epithelial cells from other mucosal sites (28). Finally, follicular hyperplasia was present in all samples as well as intensively active germinal centers (Figure 4). Such hyperplasia is attributable to proliferation of BGC, as we and others have previously shown (6, 12, 29). In light of the results shown in previous sections, such misguided
B cell response can potentially fuel the local inflammatory microenvironment, extending the impairment of immune homeostatic control.

**Microbiological Aspects of Hypertrophied Tonsils**

The most evident potential source of a local persistent immune response would be a local persistent infection. However, as opposed to recurrent tonsillitis, tonsillar hypertrophy has been long considered of non-infectious etiology. Actually, hypertrophied tonsils serve as non-infected control for tonsils affected by recurrent tonsillitis in a number of publications, including relevant and recent ones in which such control should be crucial (30). It has been generally considered that bacterial cultures from hypertrophied tonsils largely reflect oropharyngeal colonization. It is actually quite complex to distinguish infection versus colonization, provided that tonsils, being part of the mucosal immune system, are constantly exposed to the environment.
In order to try to discriminate between commensals and pathogens we started by naming the bacterial species that we could retrieve alive from the core of the samples upon cauterization of the surface, trying to identify abundant, relentless and viable populations from the clinical tissue. We cultured the core tonsillar tissue of 44 children undergoing tonsillectomy due to OSA, whose ages ranged from 2 to 15 years old (Table 1). Within our cohort of patients, the frequency of tonsillectomy varied with the age, peaking at 5-6 years old (Figure 5A). All samples rendered viable bacterial cultures except for one. At phylum level, the tonsil cultures were dominated by Firmicutes (predominantly from the genera Streptococcus and Staphylococcus in that order), Proteobacteria (mostly from the genera Neisseria and Haemophilus), Bacteroidetes (principally genera Prevotella), Actinobacteria and Fusobacterium (Tables 2 and 3) in agreement with recent data obtained by next generation sequence (NGS) (31).

Considering that certain microorganisms like Streptococcus pyogenes (S. pyogenes), Streptococcus pneumoniae (S. pneumoniae), Moraxella catarrhalis (M. catarrhalis), Haemophilus influenzae (H. influenzae) and Staphylococcus aureus (S. aureus) had been found either causing ear, nose and tonsil (ENT) pathology or as harmless local flora, both situations in competent hosts, we wanted to investigate their prevalence in our patients. We did not detect M. catarrhalis within our cohort of patients. We isolated S. pneumoniae from two patients. On the other hand, S. aureus, isolated from 45% of patients, accounted for the most frequent potentially pathogenic bacteria among the tonsils we analyzed, closely followed by H. influenzae (present in 27% of the patients) and S. pyogenes (on 23% of the samples). Bacterial isolates potentially pathogenic from the excised samples are shown in Figure 5B. In Argentina, vaccination against H. influenzae type b, S. pneumoniae, Neisseria meningitidis (N. meningitidis) and Corynebacterium diphtheria, which have been long known as commensal species in tonsils, is mandatory. We did not detect any patient carrying Corynebacterium diphtheriae but we found Neisseria meningitidis in the tonsils of a single patient and we did not serotype any isolate of H. influenzae.

In most cases we obtained co-isolates of different bacteria. Only 4 patients rendered cultures of single species, 3 of those

| Age group (years) | N° patients (%) | N° bacterial isolates | Co-isolated bacteria |
|------------------|----------------|-----------------------|---------------------|
| 2 - 5            | 18 (41)        | 58                    | 3.2 ± 1.5           |
| 6 - 9            | 18 (41)        | 52                    | 2.9 ± 1.3           |
| 10 - 13          | 7 (16)         | 17                    | 2.4 ± 1.0           |
| 14 - 15          | 1 (2)          | 6                     | 3.0 ± 1.4           |

Gender

| Gender | N° patients (%) | N° bacterial isolates | Co-isolated bacteria |
|--------|----------------|-----------------------|---------------------|
| Males  | 27 (61)        | 80                    | 3.0 ± 1.6           |
| Females| 17 (39)        | 53                    | 3.1 ± 1.3           |

The values represent the number of patients in each group who donated samples after undergoing surgery due to obstructive sleep apnea (OSA). n=44 patients were analyzed, n=1 of them did not render viable bacterial cultures.

The percentages were calculated relative to the total number of patients (n=44).

Total number of bacterial isolates.

Mean ±SD numbers of species co-infecting each individual in this cohort of patients per category of gender and age.
were *Streptococcus pyogenes* group A. Co-isolates ranged from 2 to 7 species. Details of the mean of species co-isolated from each child and all isolated organisms from the samples are listed in Tables 1–3. As we isolated a plethora of viable bacteria considered commensal and potential pathogenic as well, we were still unable to discriminate whether the tonsillar hypertrophy is of infectious nature.

The basic distinction between a pathogen and a commensal is that the first one displays aggressive tools for invasion. Usually this represents that such pathogen is able to penetrate the epithelial barriers. Therefore, we set out to detect bacterial presence in tonsillar biopsies by FISH, which allowed us to investigate its distribution and organization in situ. We used a fluorescent universal eubacterial (EUB338) probe followed by immune-fluorescence staining on cryosections from excised tonsils. As shown in Figures 5C, D, we detected bacterial aggregates not only associated with the surface of the epithelium but also within the lymphoid compartment, having breached the reticular epithelium. Therefore, while we cannot ascertain that the microorganisms detected in situ as well as through culture are the initiators of the ongoing inflammatory response, we did evidence that the chronification of the process must be certainly related to bacterial spreading beyond the normal boundaries. Also, such invasion is not associated to a single species as it can be observed in different patients harboring different colonizers, which have become pathogens, given the host’ requirement of the surgery to restore homeostasis.

**DISCUSSION**

Tonsils offer a useful model to study the interactions between the human mucosal immune system and the microbes that populate them at the epithelial level. Clinical material is not scarce, as tonsillectomy remains one of the most frequent pediatric surgeries carried out worldwide as a result of diagnosis with OSA. They rank among the secondary lymphatic organs, so every single developmental B cell subset is represented. In this study, initially we show that B cells as a population are able to produce TNFα at similar levels than those reached by T cell population, depending on the stimulating conditions. We have previously demonstrated B cell ability to rapidly produce other pro-inflammatory cytokines as IL6, IL8 and IL17 (6). Taken together, these results led us to conclude that B cell hyperplasia boost the pathological inflammatory processes and tissue alterations characteristic of OSA tonsils. Furthermore, we
TABLE 2 | Taxonomic breakdown of isolates from the core of hypertrophied tonsils at phylum level.

| Phylum       | N° species | N° bacterial isolates* (%)b |
|--------------|------------|-----------------------------|
| Firmicutes   | 16         | 65 (49)                     |
| Proteobacteria| 13         | 39 (29)                     |
| Bacteroidetes| 7          | 15 (11)                     |
| Actinobacteria| 5          | 8 (6)                       |
| Fusobacterium| 2          | 6 (5)                       |
| Total        | 43         | 133 (100)                   |

*aThe values represent the number of bacterial isolates from tonsils represented at phylum level.
*bThe percentages were calculated relative to the total number of isolates identified from the patient’s samples.

TABLE 3 | Taxonomic breakdown of viable bacterial isolates from the core of hypertrophied tonsils at genus and species level.

| Aerobic bacterial isolates                             | N° patients* (%)b |
|--------------------------------------------------------|-------------------|
| Actinomyces odontolyticus                               | 1 (2)             |
| Actinomyces graevenitii                                 | 1 (2)             |
| Corynebacterium argentoratense                          | 1 (2)             |
| Escherichia coli                                        | 1 (2)             |
| Gemella haemolytica                                     | 1 (2)             |
| Haemophilus influenza                                   | 12 (27)           |
| Haemophilus parainfluenza                               | 1 (2)             |
| Neisseria elongata                                      | 3 (7)             |
| Neisseria flavalescens                                  | 3 (7)             |
| Neisseria mucacae                                       | 1 (2)             |
| Neisseria meningitans                                   | 1 (2)             |
| Neisseria perflava                                      | 1 (2)             |
| Neisseria sicca                                         | 1 (2)             |
| Neisseria spp.                                          | 2 (5)             |
| Neisseria subflava                                      | 10 (23)           |
| Rothia dentocariosa                                     | 1 (2)             |
| Rothia mucalagnosa                                      | 4 (9)             |
| Staphylococcus aureus                                   | 20 (45)           |
| Staphylococcus intermedius                              | 1 (2)             |
| Streptococcus anginosus                                 | 4 (9)             |
| Streptococcus constellatus                              | 1 (2)             |
| Streptococcus dysgalactiae                              | 2 (5)             |
| Streptococcus intermedius                               | 1 (2)             |
| Streptococcus mitis                                     | 3 (7)             |
| Streptococcus parasanginis                              | 1 (2)             |
| Streptococcus pneumoniae                                | 2 (5)             |
| Streptococcus pyogenes                                  | 10 (23)           |
| Streptococcus salivarius                                | 9 (20)            |
| Streptococcus viridans                                  | 6 (14)            |
| Aggregatibacter aphrophilus                             | 2 (5)             |
| Aggregatibacter segnis                                  | 1 (2)             |

| Anaerobic bacterial isolates                            | N° patients* (%)b |
|--------------------------------------------------------|-------------------|
| Capnocytophaga sp.                                      | 1 (2)             |
| Fusobacterium necrophorum                               | 4 (9)             |
| Fusobacterium nucleatum                                 | 2 (5)             |
| Prevotella intermedia                                   | 5 (11)            |
| Prevotella oris                                          | 3 (7)             |
| Prevotella melaninogena                                  | 1 (2)             |
| Prevotella nanocensis                                   | 3 (7)             |
| Prevotella nigrescens                                   | 1 (2)             |
| Prevotella spp.                                          | 1 (2)             |
| Solobacterium moorei                                     | 1 (2)             |
| Veillonella atypica                                      | 2 (5)             |
| Veillonella parvula                                      | 1 (2)             |

*aThe values represent the number of patients that rendered the isolated designated in the column of the left.
*bThe percentages were calculated relative to the total number of patients (n=44).

detected a relevant proportion of CD3+ cells secreting IL17, even though we did not use culture conditions capable of specifically stimulating T cells. It has been previously reported that Th17/Th1 ratios measured in PBMC is positively related to the severity of OSA and serum levels of C-reactive protein (2). Interestingly, there are a number of human diseases in which B cells contribute to the pathology through the induction of a pro-inflammatory cytokine profile and thereby supporting T cell–mediated inflammation (18–20, 32, 33).

While others have reported on the expression of CD1d in different mucosal epithelial cells (21–23), we are the first to report its expression by tonsillar stratified as well as reticular epithelium. CD1d is a non-polymorphic MHC class I-like antigen-presenting molecule, restricted to lipid Ag presentation to invariant natural killer T cells (iNKT) cells. CD1d-binding lipid Ags from tumors as well as commensal and pathogenic microorganisms have been well characterized by many authors including ourselves (34–37). Tissue specific functions of iNKT cells were recently reviewed in (38). Of note, CD1d-mediated Ag presentation by epithelial/parenchimal cells can shape the outcome of immune responses by regulating iNKT cells function. This has been demonstrated for the case of hepatocytes, which through CD1d-lipid presentation induce iNKT cell tolerance in the liver, protecting from hepatic inflammation. Whereas iNKT cells were confirmed to be present in tonsils (39), it remains unknown their specific function at this tissue. Provided we found abundant epithelial expression of CD1d, the diversity of the tonsillar microbiota and the constant contact with environmental Ags, we suggest iNKT cells might be functionally as relevant at this site as they are in the gut (40).

Another non-classical MHC class I molecule which has been shown to be expressed by tonsillar epithelial cells is the FcRn. Tonsillar epithelium has been classified as a type II mucosal surface (27). In fact, such definition would fit for the stratified squamous epithelium. The specialized epithelium that covers the tonsillar crypts is only found in that location and is called either reticular or lymphoepithelium. Neither the squamous nor the reticular epithelium express pIgR, but they do express FcRn (41). Hence, the major post switch protective Ig isotype translocated to the lumen from the tonsillar crypts is IgG. Of note, IgM and IgG are the best isotypes to activate complement. This capacity makes them particularly efficient but also more injurious to the host due to the inflammation associated with such effector immune function. Through anti (IgM/IgG) staining we evidenced the hypothropic GCs, origin of the IgG locally produced (Figures 4, 5). We also stained for IgA and confirmed that IgA+ B cells were mainly found outside the germinal centers. Of note, it has been recently discovered in murine intestine a network of cellular interactions that drives IgM-to-IgA class switch recombination in extrafollicular B cells, compatible with the pattern we observed (42).

When studying the pathogenesis of OSA caused by hypertrophic tonsils, host immune response is only one side of the equation. The other side of it, is the microbiota populating the tonsillar mucosal surface. A diverse pool of bacteria comprising both commensal and pathogenic organisms have been isolated from OSA tonsils. In this study we worked with a...
cohort of patients whose ages ranged between 2 and 15 years old. In agreement with previous reports from patients elsewhere (43), the frequency of tonsillectomies within that age limit peaked at 5-6 years old, which is coincident with the beginning of the transition from deciduous to permanent dentition for most children. Interestingly, it has been shown that such process implies the most striking change in the salivary microbiome of all ages, when considering children from birth until 18 years old (44). Choi et al. (45) recently demonstrated that the microbiome profiles of saliva and tonsils are largely similar both, in terms of diversity and composition, in children operated due to OSA. Given the location of tonsils at the back of the oral cavity, it was somehow expected the close relation between the two microbial groups. We could therefore speculate that the important changes in the bacterial species detected when starting to acquire permanent teeth might exert extra pressure on the tonsils of some children, being the draining immune sites first handling oral Ags. Again, probably a number of factors are responsible for the loss of homeostasis, including those of the microflora but also an altered host cell immune function.

In relation to the specific bacterial species identified by culture in our samples, they are much in agreement with other studies performed either by culture or, more recently, by employing 16S rRNA gene sequencing (Tables 2, 3) (31, 45–48). In our experiments, Staphylococcus aureus (45%), Haemophilus influenzae (27%), Neisseria subflava (23%) and Streptococcus pyogenes (23%) were the most prevalent bacterial species in the core of hypertrophied tonsillar tissue within our cohort of patients. They were followed by different species of the genus Streptococcus and Neisseria. Finally, among the anaerobic prevalent isolates, we found species of Prevotella, Veillonella and Fusobacterium. As we have pointed out, all these microorganisms have been isolated from around the world, from tonsils excised due to OSA and also because of recurrent tonsillitis. Clearly, there are some differences depending on the vaccination schemes operating in different countries, which might impact in differences in the prevalence within tonsils’ crypts of S. Pneumoniae or N. meningitidis, for example. However, in general, there is consensus that the bacterial species we have found are normal oropharyngeal commensals in children. A point of particular interest to discuss is whether this means that such commensals can never be injurious. Noticeable, complications in the host immune system would imply the redefinition of most commensals as pathogens. On the other hand, a change in the model of life of these bacterial communities would have the same result. In this context, we identified bacterial aggregates in situ. Bacteria can exist either in planktonic or biofilm states. Planktonic cells are freely motile entities persisting in a liquid environment. In contrast, most of the bacterial species we named above (Haemophilus, Suturelycoccus, Streptococcus) have long been known to have the capacity of forming biofilms. In biofilms, bacteria exist as sessile aggregates encased in a self-produced complex polysaccharide matrix attached to a surface (49). The establishment of biofilms is acknowledged as a virulence factor as it allows bacteria to better resist host immune responses. While we could detect extensive aggregates attached to the surface, we did not intend to particularly identify biofilms, as we did not have samples from normal tonsils to ascertain whether biofilm presence is exclusive of pathology or not. In any case, we detect bacterial penetration through the epithelial layer to the lymphoid compartment.

The obstacle entailed by the lack of healthy pediatric tonsils to use as control group or any other comparator sample constrains the conclusions of our study to some extent. The findings described here are limited to OSA tonsils. We do not have grounds either to rule out or to extrapolate them to any other experimental model or physiological situations.

To conclude, the data presented here shed light on the contribution of the B cell compartment to the inflammatory response in OSA tonsils and underscore the importance of the interplay between the host immune system and the commensal microorganisms able to switch from asymptomatic colonization to invasive disease.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

ETHICS STATEMENT

The institutional ethics committee (Clinical Hospital, School of Medicine, Buenos Aires) approved the collection, and use of clinical material conformed to the provisions of the Declaration of Helsinki (as revised in Edinburgh 2000). Informed consent was obtained from all participants and/or their legal guardian/s. Written informed consent to participate in this study was provided by the participants’ legal guardian/next of kin.

AUTHOR CONTRIBUTIONS

LS and JR processed all the samples used in this study, performed most of the experiments and analyzed the data. RR performed a number of experiments. LB and PB performed cell sorting and advised on design of FACS experiments. GB and NS performed cryosections and H&E stainings. LS, JR, PF and BP critically reviewed and edited the manuscript. MA and BP provided number of experiments. LB and PB performed cell sorting and advised on design of FACS experiments. GB and NS performed cryosections and H&E stainings. LS, JR, PF and BP critically reviewed and edited the manuscript. MA and BP provided samples. CB, CV and FM performed the microbiologic culture and identifications. GB and NS performed cryosections and H&E stainings. LS, JR, PF and BP critically reviewed and edited the manuscript. MA and BP provided samples. CB, CV and FM performed the microbiologic culture and identifications. EA supervised and designed research, analyzed the data and wrote the manuscript. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2021.648064/full#supplementary-material
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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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