Ankylosing spondylitis patients display altered dendritic cell and T cell populations that implicate pathogenic roles for the IL-23 cytokine axis and intestinal inflammation

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Abstract
Objective. AS is a systemic inflammatory disease of the SpA family. Polymorphisms at loci including HLA-B27, IL-23R and ERAP-1 directly implicate immune mechanisms in AS pathogenesis. Previously, in an SpA model, we identified HLA-B27-mediated effects on dendritic cells that promoted disease-associated Th17 cells. Here we extend these studies to AS patients using deep immunophenotyping of candidate pathogenic cell populations. The aim of our study was to functionally characterize the immune populations mediating AS pathology.

Methods. Using 11-parameter flow cytometry, we characterized the phenotype and functions of lymphocyte and myeloid cells from peripheral blood, and the synovial phenotype of AS patients and age-matched healthy controls.

Results. Significantly fewer circulating CD1c-expressing dendritic cells were observed in AS patients, offset by an increase in CD14−CD16+ mononuclear cells. Ex vivo functional analysis revealed that this latter population induced CCR6 expression and promoted secretion of IL-1β and IL-6 when co-cultured with naive CD4+ T cells. Additionally, systemic inflammation in AS patients significantly correlated with increased proportions of activated CCR9+ CD4+ T cells.

Conclusion. CD14−CD16+ mononuclear cells may contribute to AS by promoting Th17-mediated AS disease pathology through CCR6, IL-1β and IL-6 induction.

Key words: dendritic cells, CD4+ T cells, ankylosing spondylitis, inflammation, CD14−CD16+ mononuclear cells.

Rheumatology key messages
• The AS immunological signature consists of altered myeloid (CD1c+:CD16+) and lymphocyte (CCR6+:CXCR3+) profiles.
• CD14−CD16+ mononuclear cells may support Th17-mediated AS disease pathology through CCR6, IL-1β and IL-6 induction.
• The intestinal environment may drive systemic inflammation in AS patients in a CCR9-dependent manner.

Introduction
AS is a chronic arthropathy of the axial and peripheral skeleton, characterized by inflammation and abnormal bone and entheseal remodelling. AS is recognized as the prototypic disease of a larger SpA group of conditions that share genetic, pathophysiological and clinical features. Although transformed by the introduction of...
TNF inhibitors a decade ago, current therapeutics remain inadequate for a substantial proportion of patients, offering a compelling case for improved pathophysiologi
discovery. Large genome-wide association studies have recently contributed several novel loci and hence novel candidate pathways in this regard. There is particular interest in the confirmed associations with IL-
23R, ERAP1 and STAT3 [1–3]. Taken together with the provisional clinical benefit accrued upon inhibition of the IL-17A pathway in clinical trials in AS patients [4], this suggests that novel immune-mediated pathways beyond TNF may now be formally implicated in pathogenesis.

Prior mechanistic studies have relied heavily on data generated using animal models. Experiments using transgenic rats overexpressing the human HLA-B27 and β2-microglobulin genes (B27-TG) have, for instance, implicated CD4+ T cells [5, 6], myeloid cells [7, 8] and endoplasmic reticulum (ER) stress [9] in the development of SpA. There are as yet few data informing analogous mechanisms operating in patients. Given that HLA-B27 is an MHC class I molecule, it is intriguing that disease in B27-TG rats occurs independently of CD8+ T cells [6]. Moreover CD4+ T cells expanded in AS patient peripheral blood are enriched for a population of Th17 phenotype [10–12]. Dendritic cells (DCs), which regulate T cell responses, have been strongly implicated in SpA pathogenesis in B27-TG rats [7, 8, 13]. Parallel studies in patients are, however, rare. Most have heretofore focused on cells generated from monocytes following ex vivo culture with GM-CSF and IL-4 (mo-DCs) [14]. These express reduced surface MHC II expression compared with cells from healthy individuals, but no alteration in production of IL-23 or other cytokines [15, 16]. On the other hand, in vitro studies do suggest that ER stress of DCs could lead to excessive IL-23 production [17], and as such they are intriguing candidate cells in initiating this effector pathway. In this regard, however, the relevance of data generated using cultured monocytes to DC biology is unclear [18]. We consider it critical now to determine the contributions of bone fide DCs and other myeloid lineages to AS pathogenesis.

Our previous analyses revealed a systemic deficiency in a specific DC population in B27-TG rats, which in turn promoted elaboration of Th17 responses [19]. Because this cytokine axis may be associated with AS pathology, we hypothesized that myeloid populations could be similarly altered in AS patients. Accordingly, we performed deep phenotyping of blood and SF leucocyte populations clearly altered in AS patients. Accordingly, we performed we hypothesized that myeloid populations could be similar

**Materials and methods**

**Patients**

AS patients meeting the modified New York criteria [20] were recruited from the Glasgow Royal Infirmary rheumatology clinic between March 2011 and June 2013. Informed, written consent was obtained from all study participants according to the Declaration of Helsinki. Ethical approval for the study was awarded by the West of Scotland Research Ethics Service (Institute of Infection, Immunity and Inflammation Research Tissue Bank, REC: 11/S0704/7). Consenting age- and sex-matched healthy controls (HCS) were recruited under ethics approved by the College of Medical, Veterinary, and Life Sciences Ethics Committee, University of Glasgow (Project #2013007). Patient clinical features are outlined in supplementary Table S1, available at Rheumatology Online.

**Reagents**

Cells were cultured in RPMI 1640 supplemented with 10% fetal calf serum, 100 U/ml penicillin, 100 μg/ml streptomycin, 2 mM l-glutamine and 50 μM 2-mercaptoethanol (complete medium).

**Peripheral blood mononuclear cell isolation**

Isolation of peripheral blood mononuclear cells was performed over a Histopaque–1077 gradient. Following centrifugation, the peripheral blood mononuclear cell layer was harvested.

**Antibodies**

Directly conjugated or biotin-labelled antibodies targeting CD3 (UCHT-1), CD4 (OKT4), CD14 (M5E2), CD15 (W6D3), CD16 (3G8), CD19 (HIB19), CD25 (BC96), CD45RA (HI100), CD56 (MEM-188), HLA-DR (L243), TcRβ (IP26) and CXCR3 (GO25H7) were from Biolegend (San Diego, CA, USA). Antibodies targeting CCR9 (248621) and CCR10 (314305) were purchased from R&D systems (Minneapolis, MN, USA). CD1c (AD5-B7) and anti-SLAN (M-DC8, DD-1) antibodies were from Miltenyi Biotec (Auburn, CA, USA). CD11c (B-ly6), CD141 (1A4), CCR6 (11A9) and CCR4 (1G1) antibodies were purchased from BD Biosciences (Oxford, UK).

**Flow cytometry**

Following Fc receptor inhibition (eBioscience, San Diego, CA, USA), staining was performed in PBS with 2% fetal calf serum and 2 mM EDTA. Where biotin-conjugated antibodies were used, a streptavidin step was performed. Samples were acquired using LSR II (BD Biosciences) or MACSQuant (Miltenyi Biotec) flow cytometers, or purified using a FACSAria cell sorter (BD Biosciences). Data were analysed using FlowJo software (version 9.2; Tree Star, Ashland, OR, USA).

**Naive CD4+ T cell isolation**

Naive CD4+ T cells were isolated from peripheral blood mononuclear cells using a naive CD4+ T cell isolation kit.
Mixed leucocyte reaction
Myeloid populations were re-suspended in complete medium and co-cultured with CFSE-labelled allogeneic naive CD4+ T cells for 5 days. Supernatants were harvested and proliferation assessed through CFSE dilution.

RNA extraction
A MicroRNA kit (Qiagen, Venlo, Netherlands) was used for RNA extraction. Genomic DNA was removed using an RNase-free DNase kit (Qiagen). cDNA was generated using a Superscript First Strand Kit (Invitrogen).

Real-time quantitative PCR
Gene expression was measured using the Brilliant III Ultra Fast SYBR GREEN qRT-PCR Master Mix (Agilent Technologies, Santa Clara, CA, USA). Reactions were analysed using a 7500 Fast Real-Time PCR System machine (Applied Biosystems, Foster City, CA, USA). Primers used were TATA-binding protein (TBP): forward AGA CCT TCCTGTATCCCTTGG, reverse TAGGTGGGTCAGCTG; ATF4: forward GACCAGTTGATGACCTCG, reverse GGGAAAGGTTGGTTAAGAGGTG; PERK: forward TGCTGGTCTGGAAGACACACC, reverse TGAGTCGATCCCATGGGTAGGA; EIF2a: forward GCTCTTGACAGTCCGAAGG, reverse CATTGCCCAAGGCAAAAG; unspliced XBP-1: forward AGACAGCGCTTGGGGATGGA, reverse CCGTTGACAGGTCAGCTAG; spliced XBP-1: forward AGAGAGGCTGGGGATGGA, reverse CCGTTGACAGGTCAGCTAG; CCR9, CCR4 and CCR10 were not differentially expressed on HC and AS patient CD4+ T cell populations (Fig. 1A). Elevated levels of CCR9 and reduced CXCR3 expression were observed on activated (CD25int) CD4+ AS patient T cells (Fig. 1C and D). CXCR3 expression was also reduced on AS CD4+ memory T cells (Fig. 1D).

Cytokine detection
Cytokines IL-1β, IL-4, IL-5, IL-6, IL-10, IL-17A, GM-CSF, IFNγ and TNFα were detected using a custom-made Luminex assay (R&D systems), using the BioPlex 200 system (Bio-Rad, Hercules, CA, USA). Plates were read using an MRX microplate reader (Dynex technologies, Chantilly, VA, USA).

Results
The CD4+ T cell CCR6/CXCR3 axis is altered in AS patients
T cells belonging to the Th17 lineage have been associated with disease pathogenesis in B27-TG rats [9, 19] and AS patients [10–12]. We performed detailed immunophenotyping of the circulating CD4+ T cell profile in AS through quantification of T cell populations and their effector status. The latter aspect was determined through examination of chemokine receptor expression (commonly used surrogate markers of T helper cell polarization) [21, 22]. We first segregated four CD4+ TcRαβ+ populations based on CD25 and CD45RA expression (Fig. 1A), namely naive (CD45RA+CD25−), memory (CD45RA−CD25+), activated (CD45RA−CD25−) and non-activated (CD45RA+CD25+) phenotypes and Tregs (CD45RA−CD25+). All Tregs expressed FoxP3 (Fig. 1A). Enumeration of CD4+ T cell subsets revealed no differences between AS patients and HCs (Fig. 1B). Elevated levels of CCR6 and reduced CXCR3 expression were observed on activated (CD25int) CD4+ AS patient T cells (Fig. 1C and D). CXCR3 expression was also reduced on AS CD4+ memory T cells (Fig. 1D).

AS patient plasma contains elevated levels of Th17-associated cytokines
To understand how the balance of T helper subsets might be regulated in AS patients, we examined the plasma cytokine and growth factor profiles in our cohort. We observed a modest increase in TNFα levels in AS patient plasma (Fig. 2A). IL-1β levels were similar to those of controls (Fig. 2B). Assessment of several plasma Th-associated cytokines revealed a significant reduction in plasma IL-4 in AS patients, while IL-10 and IFNγ levels resembled those of HCs (Fig. 2C). In contrast, there was significant upregulation in the circulating levels of the Th17-associated cytokines IL-6 and IL-23p19, but no increase in IL-17A itself (Fig. 2C). Thus, the cytokine milieu in AS patients is skewed towards the Th17/IL-23 axis. Interestingly, Fli3L, a myeloid haematopoietic growth factor, was significantly upregulated in AS patient plasma compared with HCs (Fig. 2D).

The proportion of circulating CD16+ mononuclear cells increased in AS
Data from our laboratory support a relationship between systemic myeloid dysregulation and AS/SpA; we previously demonstrated that B27-TG rats lack CD103+ CD172a+ DCs [19], while Fli3L was significantly upregulated in AS patients (Fig. 2D). Thorough examination of myeloid populations may elucidate their involvement in promoting the aberrant T cell phenotypes observed in

Statistics
Results were presented as +s.d. Data were analysed by Student’s t-test or Mann–Whitney U test, followed by a Bonferroni post-test using GraphPad prism (San Diego, CA, USA). Linear regression, Kruskal–Wallis Spearman correlation tests were performed with Dunn multiple comparisons post-tests. Values of P<0.05 were considered statistically significant.
our patient cohort. To our knowledge, no prior study has directly addressed this question \textit{ex vivo} without prior cell expansion in culture. Lineage-CD14-CD11c+ HLA-DR+ cells were subdivided into three myeloid populations based on their differential expression of CD141 and CD16 (supplementary Fig. S2A and B, available at Rheumatology Online). The CD141-CD16- population expresses CD1c (data not shown) and thus represents CD1c+ DCs. The CD141-CD16+ population [23], collectively termed CD14-CD16+ mononuclear cells, was heterogeneous for 6-Sulpho LacNAc (SLAN) (M-DC8) and thus contained CD14-CD16+ SLAN- and SLAN+ subsets (supplementary Fig. S2A, available at Rheumatology Online). No difference was observed in the proportion of total DCs (CD1c+ and CD141+ DCs) between HCs and AS patients (Fig. 3A). Unfortunately, our study was insufficiently powered, with too few HLA-B27+ HCs and HLA-B27+ AS patients to examine the influence of HLA-B27 on immunological disease parameters (Fig. 3A). Analysis of plasmacytoid DCs (HLA-DR+...
CD123+ CD304+ revealed no significant difference between HCs and AS patients (supplementary Fig. S2C, available at *Rheumatology* Online). We observed no disparities in the frequencies of CD141+ DCs in AS patients (Fig. 3B). However, analogous to that population we predicted would be deficient from our B27-TG experiments [19], analysis of clinical data (supplementary Table S1, available at *Rheumatology* Online) revealed a significant correlation between increased proportions of CD141+ DCs and greater disease severity when assessed by BASDAI (disease activity) but not by BASFI (for functional limitation see supplementary Fig. S3, available at *Rheumatology* Online).

Unexpectedly, we observed a significant proportional reduction in CD1c+ DCs in AS patients (Fig. 3B). This finding was offset by increased frequencies of total CD14+ CD16+ mononuclear cells in AS patients (P = 0.0552), despite no proportional differences in SLAN-expressing subsets (Fig. 3C). Comparison of the major blood myeloid populations revealed a significant shift from CD1c+ DCs towards CD14+ CD16+ mononuclear cells in AS patients (Fig. 3D). The unfolded protein response (UPR) induced within myeloid cells as a consequence of ER stress initiated through HLA-B27 misfolding, is thought to contribute to disease pathogenesis in B27-TG rats [7–9]. Despite the observed myeloid dysregulation (Fig. 3B and C), we failed to detect UPR induction in AS patient CD1c+ DCs and CD14+ CD16+ mononuclear cells (Fig. 3E). Upregulation of BiP expression was observed within CD14- CD16+ mononuclear cells (Fig. 3E).

Myeloid populations can be identified in AS patient SF

A presumed pathophysiological destination of circulating cells is the articular compartment. As tissue is not readily available from the spine, we analysed the immune cell milieu of four AS patient SF samples. Analysis of lineage-CD11c+ MHC II+ cells revealed four myeloid populations: CD14+ CD16+ monocytes, CD14+ CD16+ monocytes and two DC populations (CD141+ and CD1c+; Fig. 4A). In contrast to peripheral blood (Fig. 3C and supplementary Fig. S2B, available at *Rheumatology* Online), the CD14+ CD16- SLAN+ and SLAN- mononuclear cell subsets were absent from AS SF (Fig. 4A). High levels of CCR4 and CCR6 expression were observed on SF CD3+ CD4+ T cells (Fig. 4C), and most CD4+ CD25hi cells co-expressed both chemokine receptors (Fig. 4C). To identify inflammatory mediators driving joint pathology, the myeloid milieu from matched AS patient SF and blood samples was analysed. IL-10, IL-17A, TNFα and Flt3L were not differentially expressed between plasma and SF (Fig. 4D).
In contrast, IL-6 concentrations were dramatically increased within AS SF (Fig. 4E).

**DC-T cell interactions promote Th17-like immune responses**

Cognate interactions between antigen-presenting cells, principally DCs, and T cells are central to the regulation of immune responses. To investigate blood myeloid cell function, we assessed their ability to stimulate proliferation of naive allogeneic CD4+ T cells. HC and AS patient myeloid cells induced similar CD4+ T cell proliferation at all DC:T cell ratios (Fig. 5A). DCs have a greater capacity to induce naive T cell proliferation than blood monocytes (supplementary Fig. S4, available at Rheumatology Online). Supernatant analysis from these co-cultures revealed no difference in IL-1β and IL-6 production between HCs and AS patients (Fig. 5B). CD14+CD16+ mononuclear cell–CD4+ T cell co-cultures consistently secreted the highest levels of IL-1β and IL-6 (Fig. 5B). The Th-associated cytokines IFNγ and IL-17A were also examined; however, no significant differences were observed (data not shown). Overall, interactions between CD14+CD16+ mononuclear cells and T cells promote secretion of the Th17-associated cytokines IL-1β and IL-6.
CCR6 is often used as a surrogate marker for Th17 cells [22], although not all CCR6+ cells secrete IL-17A (data not shown). Given that published data implicate Th17 cell involvement in AS pathogenesis [10–12], and the elevated CCR6 expression on AS patient CD4+ T cells observed by us, we explored whether these phenomena could arise from altered DC–T cell interactions. We performed mixed leucocyte reactions and assessed CCR6 expression on proliferating CD4+ T cells (Fig. 5C). Blood DCs and CD14−CD16+ mononuclear cells from AS patients and HCs all induced CCR6 on interacting T cells, to similar levels (Fig. 5D).

### Disease pathogenesis linked to CCR6+ and CCR9+ CD4+ T cells

Finally, to further explore our data relative to clinical phenotypes, we evaluated the relationships between immunological and clinical parameters. We observed that increased frequencies of CCR6+ Tregs (CD25hi) were significantly associated with elevated disease severity,
assessed by both BASMI and BASFI (Fig. 6A). Although similar relationships were observed between increasing frequencies of CCR6 + activated T cells (CD25+) and BASMI and BASFI, these did not reach significance (Fig. 6A). Furthermore, we observed significant correlations between increased proportions of CCR9 + CD4+ activated T cells and elevated ESR and CRP scores (Fig. 6B).

Discussion

The genetic link between HLA-B27 and AS was identified more than 40 years ago, yet the mechanisms underpinning this association remain elusive. Studies of B27-TG rats indicate involvement of myeloid populations and CD4+ T cells (specifically the Th17 lineage) in SpA pathophysiology [5-9, 19, 24]. Parallel human studies have highlighted involvement of the Th17/IL-23 pathway in AS [1, 2, 12, 25]. Despite this, evidence for IL-17-secreting T cells in inflamed, extra-articular tissues is limited [26-28]. Furthermore, examination of the critical myeloid populations has only been performed indirectly, using in vitro-generated mo-DCs [15, 16]. These cultures do not recapitulate the diverse populations found in vivo. We have performed the first detailed phenotypic and functional analyses of freshly isolated myeloid populations in AS. We have not, however, performed these analyses using samples from patients with other inflammatory diseases. Such comparisons have previously been shown to be valuable [29, 30]. Further studies are therefore required to establish whether our immunological observations are unique to AS pathophysiology. Additionally, the majority of
our patients presented with long-standing disease; if samples were available it would be fascinating to repeat these studies using patients in the earliest stages of disease.

Significantly, our analyses support reports of a systemic imbalance in circulating CD4+ T cell subsets in AS, favouring cells of the Th17 lineage. We identified this imbalance using CCR6 and CXCR3 as surrogate markers of...

Fig. 6 Immunological parameter correlations with disease progression and inflammation in AS

(A) Correlations between AS patient (n = 14) CCR6-expressing CD4+CD25hi T cells (Tregs; Top) and CCR6+CD25int T cells (activated; bottom) with BASMI (left) and BASFI (right) clinical scores. (B) ESR and CRP correlations with AS patient (n = 16) CCR9+CD4+CD25int (activated) cell frequencies. Each dot represents one individual. Analyses were performed and assessed using linear regression and the Kruskal–Wallis Spearman correlative statistical tests (r), using the Dunn multiple comparisons post-test. *P < 0.05, **P < 0.01 and ***P < 0.001.
Th17 and Th1 differentiation, respectively [21, 22]. The elevations we observe in circulating levels of the Th17-associated cytokines IL-6 and IL-23 may promote induction and survival of these Th17-like cells in vivo [31, 32]. IL-23 has also been shown to directly drive SpA-associated pathology in animal models, and has been associated with human disease [26, 33, 34]. Our data support these observations, implicating involvement of CCR6/Th17-associated CD4+ lymphocytes and the IL-23 cytokine axis in AS pathogenesis [9, 19, 25].

We previously identified a systemic deficiency in the CD103+ CD172a+ tolerogenic DC population in B27-TG rats [19]. We hypothesized that the analogous human subset, CD141+ DCs, would be absent or depleted in AS patients. However, we observed no differences in the frequencies of circulating CD141+ DCs between AS patients and HCs. In fact, elevated proportions of circulating CD141+ DCs were weakly associated with increased disease severity. Thus, unlike in B27-TG rats, a loss of this DC subset does not appear to drive human disease. Hence, several immunological processes driving human AS appear to differ from those driving SpA-like symptoms in B27-TG rats. In contrast, we observed a significant reduction in the frequency of the largest blood DC population, CD1c+ DCs, in AS patients. It should be noted that there was a 15-year difference between the average ages of the HC and AS patient cohorts. We consider it unlikely that this disparity caused the observed differences between our HC and AS patient populations because our analyses did not detect any influence of donor age on the number or frequency of any DC population in our cohorts (data not shown). In myeloid cells of B27-TG rats, the UPR, driven by misfolded HLA-B27 molecules, induces release of pro-inflammatory cytokines and cell death [9, 35, 36]. However, we could detect no evidence for UPR activation in circulating CD1c+ DCs and CD14− CD16+ mononuclear cells from AS patients. Our results thus support previous observations using more heterogeneous cell preparations from blood or intestine [37–41] and implicate involvement of alternative disease-associated immunopathogenic mechanisms. If HLA-B27-induced UPR induction is indeed the force driving AS, future investigations will need to carefully examine whether any tissue-resident cell populations show evidence of ER stress or UPR activation.

Accompanying the reduced frequency of blood CD1c+ DCs was an increase in the heterogeneous CD14− CD16+ mononuclear cell population. This population contains both patrolling CD16+ blood monocytes [23] and MDC8/SLAN+ DCs [42, 43]. To understand the functional consequences of these changes, we assessed the ability of these myeloid populations to activate naïve T cells. On a per-cell basis, no differences were observed between AS patient and HC myeloid cells; CD14− CD16+ mononuclear cells promoted secretion of the Th17-associated cytokines IL-1β and IL-6 and were capable of inducing CCR6 expression on interacting T cells. Consequently, the observed shift towards the CD14− CD16+ myeloid population in AS patients may contribute to the induction of the pathology-associated Th17 response. Of note, SpA patient CD1c+ DCs express elevated levels of CD1d, which was shown to have immunoregulatory properties [44]. Our observation, that there is a reduction in the frequency of circulating CD1c+ DCs in AS patients, contributes to the evidence suggesting that myeloid cells may be involved with disease progression. To test this connection, further studies are required to address the role of DCs in disease sites of ongoing inflammation in AS.

Analyses of the more readily available cells in blood may not accurately reflect immunological processes occurring in disease-affected tissues. Examination of tissue populations can therefore be helpful in elucidating pathogenic mechanisms. SF samples from peripheral joints, although not necessarily representative of AS spinal or musculoskeletal pathophysiology, remain an important resource for characterizing potential tissue effector populations. We identified CD1c+ DCs in AS SF, as previously observed in RA SF [45]. Additionally, we identified a high frequency of CD141+ DCs in AS SF. These SF-resident DCs may perpetuate disease by driving local T cell responses. However, because these results were generated using very limited numbers of samples, further phenotypic and functional analyses must be performed to better understand the role of the immune populations residing within the SF. Of note, CD14− CD16+ mononuclear cells were completely absent from AS SF, indicating that they do not act locally to drive peripheral joint disease in AS. CD14− CD16+ SLAN+ cells have, however, been identified in inflamed extra-articular tissues, including tonsil and psoriatic skin [46, 47]. Together with our results, these data suggest that extra-articular CD14− CD16+ cells in lymphoid or other peripheral tissues may contribute to AS pathogenesis.

To understand how immunological changes may contribute to disease development, we must consider how these differences relate to patient clinical characteristics. In our cohort, it was clear that CCR6+ Tregs were more frequent in patients with the highest disease scores. From this correlation we infer that in AS patients, CCR6-expressing Tregs may be induced in an attempt to suppress pathology. Further investigation is clearly required to accurately assess whether this population has any effect on disease progression.

Interestingly, our data indicate a relationship between intestinal immune responses and systemic inflammation in AS; elevated ESR and CRP values correlate with higher proportions of circulating CCR9-expressing activated T cells. CCR9 is induced on T cells activated in intestinal lymphoid tissues [48]. Thus, this correlation indicates that systemic inflammation in AS is associated with intestinal activation of T cells. This is potentially extremely important, given the absence of inflammation from B27-TG rats housed under germ-free conditions, and the fact that ~50% of AS patients show evidence of subclinical intestinal inflammation [49, 50]. Together, these results indicate that antigen-presenting cells of intestinal origin may play a central role in driving systemic inflammation in AS.
patients. Given this association between systemic inflammation and the intestinal immune response, it is important to understand the impact of peripheral disease manifestations, especially signs of IBD, on disease and immunological parameters. Unfortunately, our study was not sufficiently powered (containing too few patients with IBD) to address this directly.

In summary, we have performed extensive immunophenotyping of AS patients and have identified disease-associated alterations in circulating T lymphocytes and myeloid populations. Specifically, elevated proportions of CD14+CD16+ mononuclear cells, capable of inducing CCR6-expressing T cells and IL-1β and IL-6 production, may contribute to the Th17 immune responses that are associated with AS. Furthermore, intestinal immune responses appear to contribute to systemic inflammation in AS patients, implicating loss of intestinal homeostasis as an important factor driving disease. Together, these observations enhance our understanding of the pathogenesis of AS, and support the idea that aberrant T cell responses and intestinal inflammation contribute to the pathogenesis of this disease.

Supplementary data
Supplementary data are available at Rheumatology Online.

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