Th2 mediated regulation in RA and the spondyloarthropathies

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Can atopy help to clarify the role of Th2 mediated regulation in these diseases?

Amongst the heterogeneity of human immune responses T helper (Th) lymphocyte subsets have been shown to have an important role.1 Of these different subsets, Th1 cells mediate cellular immunity, including cytotoxicity and delayed-type hypersensitivity responses through the specific production of interferon γ (IFNγ) and interleukin (IL) 2. Th2 cells, characterised by IL4, IL5, and IL13 production, favour humoral immunity and down regulate Th1 mediated cellular immunity. Th2 responses are associated with IL4/IL13 mediated IgE production and IL5 mediated eosinophilia. Th1 activity in its turn inhibits these responses and results in effective immune responses against several infectious agents such as bacteria and viruses. Also, in several autoimmune diseases Th1 cells contribute to the induction and persistence of inflammation and inflammation-induced tissue damage.

Numerous studies have shown that Th1-induced immunity is inhibited by suppressive Th cells other than IL4+ Th2 cells. These suppressive cells are also distinguished by their particular cytokine secretion and/or function: transforming growth factor β (TGFβ), Th3 cells, IL10+ T regulatory 1 (Tr1) cells, and CD4+CD25+ anergic/suppressive cells.2,3 Although all these subsets may contribute to suppression of Th1 activity, the balance between Th1 and Th2 cells has been shown to strongly influence many inflammatory responses. Owing to the mutually antagonising abilities of Th1 and Th2 cells in many experimental animal and human in vitro studies, the Th1/Th2 balance in rheumatoid arthritis (RA) has been extensively studied.

“Balance between Th1 and Th2 cells strongly influences inflammatory responses”

Th1 predominance in RA and the impact of atopy-induced Th2 responses

In RA synovial tissue, synovial fluid, and serum, analysis of IFNγ and IL4 production to indicate Th1 and Th2 activity, showed that Th1 activity was clearly predominant and Th2 activity was absent compared with control subjects (table 1). 

The Th1/Th2 imbalance in RA joints is associated with high numbers of activated macrophages, leading to an aggressive form of arthritis with rapidly occurring joint destruction.4 Based on the pivotal role of the Th1 predominance in RA it has been suggested that patients with RA will benefit from Th2 activity. Until now, treatments aimed at enhancing or mimicking Th2 activity—for example, through IL4 or IL10, have not provided evidence for this hypothesis.5 However, the naturally occurring mutual antagonism of atopy and RA supports this hypothesis and indicates the role of Th1/Th2 balance in RA.

“RA is associated with a 40–50% reduction in atopic disorders”

The prevalence of RA, in which Th1 predominates, was found to have a favourable impact on several atopic disorders known to be associated with a clear Th2 predominance (table 2). In five European studies RA was associated on average with a 40–50% reduction in the prevalence of atopic disorders.8,10 In one study which evaluated a limited number of patients with RA (n=40) atopy assessment by a health assessment questionnaire did not show a decreased prevalence.12 However, when allergen skin prick tests were performed to confirm atopy, a decrease of 45% in patients with positive tests was found among patients with RA compared with healthy controls.12 Similarly, in another study where hay fever was confirmed by this test, a 50% reduction in patients with RA compared with non-RA controls was seen.22 Furthermore, this was associated with a reduction of serum IgE levels and eosinophilia, further indicating suppression Th2 responses.

In addition to the inhibition of Th2 mediated immunity by Th1 cells, in hay fever/patients with RA reduced Th1 activity (IFNγ production) was seen compared with patients with RA with no hay fever (table 2). This Th1 reduction was associated with reduced disease activity as measured by reduced acute phase responses and joint scores for inflammation and joint destruction. In agreement with this, Rudwaleit et al reports in this issue of the Annals that atopic patients whose atopy started at age 20 or earlier had a reduced disease severity.13

In support of the mutual antagonism of Th1 mediated autoimmune diseases and atopy other studies have shown that both patients with multiple sclerosis and type 1 diabetes (Th1 driven diseases) had decreased prevalence and fewer symptoms of IgE mediated allergic diseases (table 2).14,15 Furthermore, an inverse

Table 1  Cytokine profiles in rheumatoid arthritis and the spondyloarthropathies

<table>
<thead>
<tr>
<th>Cytokine analysis in biological fluids and tissue</th>
<th>IFNγ</th>
<th>IL4</th>
<th>IL10</th>
<th>TNFα</th>
<th>IL1β</th>
<th>IL6</th>
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<tr>
<td>Serum*</td>
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<td>RA</td>
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<td>ReA</td>
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<td>Synovial fluid*</td>
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<td>RA</td>
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<td>AS</td>
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<td>ReA</td>
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<tr>
<td>Synovial tissue†</td>
<td>+</td>
<td>0</td>
<td>++</td>
<td>++</td>
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<td>+++</td>
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<tr>
<td>RA</td>
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<tr>
<td>T cell cytokine analysis‡</td>
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<tr>
<td>Peripheral blood*</td>
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<tr>
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</tbody>
</table>

Assessed by ELISA* or immunohistochemistry/in situ hybridisation†. T cell cytokine analysis (by ELISA/FACS) is done after short term in vitro culture of T cells. Increased [+ or ++], decreased [−] or equal [0] cytokine expression levels compared with control subjects are indicated. The summary in this table is based on references 4–6, 17–24, 29 and 33 among others.
association was observed between tuberculin-induced inflammatory responses and atopic disorders which were related to the mutual inhibition of Th1 (IFNγ, IL12) and Th2 mediated (IL4, IL10 and IL13) immunity (table 2). In the latter study it was shown that the balance between these responses changed over time, indicating that environmental factors influence the Th1/Th2 balance in genetically predisposed subjects. The abovementioned studies support the assumption that several autoimmune diseases, including RA, might benefit from Th2 mediated immune deviation. Therefore, studying the impact of well defined Th2 mediated diseases on arthritic conditions may help to clarify the role of this immune response in regulation of disease activity. Thus, Rudwaleit et al investigated the impact of atopic disorders on the seronegative spondyloarthropathy ankylosing spondylitis (AS). Additionally, the reverse—the effects of this disease on atopic disorders—was also studied.

**Th1/Th2 balance in AS and atopy-induced Th2 responses**

The seronegative spondyloarthropathies include a heterogeneous group of diseases characterised by inflammatory axial spine disease, asymmetric peripheral arthritis, enthesopathy, inflammatory eye disease, and overlapping mucocutaneous features occurring in the absence of serum rheumatoid factor. In adults this group of diseases includes AS, reactive arthritis (ReA), some forms of psoriatic arthritis (PsA), undifferentiated forms of spondyloarthropathy (uSpA), and arthropathies accompanying inflammatory bowel disease. These arthropathies share features with RA, such as inflammation of the peripheral joints. To specifically intervene in these arthritides it is important to define systemic and intra-articular immune responses. Several studies have defined these responses in AS to a certain extent. The profile of circulating proinflammatory cytokines in AS has similarities with RA but differs in some aspects (table 1). Serum levels of TNFα and IL6 in AS are increased compared with controls, although cytokine levels are generally lower than in patients with RA. Increased TNFα production was also found in the synovial tissue of patients with AS. In contrast with RA, IL1β was not increased in AS serum and synovial tissue. IFNγ also in contrast with RA, could not be detected in the serum of patients with AS.

Although a large number of studies have confirmed the Th1/Th2 imbalance in RA, only a few studies have focused on Th cytokine profiles in AS. Cytokine profiles of circulating T cells in AS are similar to those found in patients with RA (table 1). Both in patients with RA and those with AS, decreased peripheral IFNγ and TNFα production by T cells is observed, whereas unchanged and even increased IL4 production is documented. Similar observations were made in patients with ReA and seem to hold true for PsA and uSpA. Analysis of peripheral T cell cytokine profiles might be helpful but should be interpreted with caution. For example, reduced peripheral Th1 activity and TNFα production in patients with RA and ReA are associated with increased local production of these cytokines during active disease (table 1). Furthermore, reduced peripheral T cell cytokine secretion is at least partly a consequence of the strong ability of activated Th1 cells to migrate to sites of inflammation. This is associated with high antigen reactivity at sites of inflammation and low antigen reactivity in the peripheral blood of patients with RA, AS, and ReA. Therefore, the reduced peripheral Th1 activity in patients with AS, as well as in other spondyloarthropathies, cannot be considered as evidence for local impaired Th1 cytokine production. This is supported by the reduction in TNFα secretion by peripheral T cells in AS, observed to be associated with increased TNFα production in synovial tissue (table 1). These findings suggest that further investigations are required to define, in particular, the intra-articular inflammatory response in AS, including the Th1/Th2 balance.
with RA.\textsuperscript{29} Although together the data are suggestive of an impaired or absent Th1 cytokine production in joints of patients with AS, more data need to be collected to confirm this suggestion. The assumption that AS is characterised by a Th2 cytokine profile,\textsuperscript{30} however, is not scientifically sound. Increased production of IL4 has not been shown in AS. In two patients with AS it was even found that IL4 mRNA expression in the synovial tissue was lower than the IL4 expression in patients with RA, which in many studies has been shown to be very low.\textsuperscript{12}

To gain further insight in the role of the Th1/Th2 balance in patients with AS, Rudwaleit \textit{et al} studied the impact of Th2 driven atopy on AS in comparison with the impact of atopy on RA.\textsuperscript{10} The reverse—the effects of RA and AS on atopy—was also studied. The atopy-induced amelioration of disease severity in patients with RA was not seen in patients with AS.

“\textit{Atopic diseases are not decreased in patients with AS}”

Also, the decreased prevalence of atopic disorders, such as hay fever, asthma, and atopic dermatitis, in patients with RA was not found in patients with AS (table 2). Instead, the prevalence of atopy in patients with AS (24.6\%) was slightly increased compared with healthy controls (20.7\%). This difference is even more pronounced (23.4\% in AS v 15.8\% in controls, 48\% increase) when the bias introduced by latex allergies is taken into account, because these were observed more often in the control group which consisted of hospital workers who are more often exposed to latex (4.9\% for controls v 1.2\% for AS).

These data suggest that AS is not a disease in which Th1 predominates because atopic responses are not inhibited by AS. Furthermore, patients with AS are not likely to benefit from Th2 activity because atopy does not reduce the symptoms of AS.

Conclusions and discussion

Low Th1 activity in patients with AS is supported by the lower proinflammatory cytokine production (observed as lower TNF\(\alpha\) and IL1\(\beta\) levels) in AS compared with patients with RA. This is in line with the less aggressive form of arthritis seen in patients with AS. The slightly increased prevalence of atopy in AS might indicate that AS is associated with a predominant humoral immunity. Although there is no evidence for a typical Th2 response, joint inflammation in patients with AS may be associated with humoral responses. In patients with rheumatic fever, arthritis has been shown to be induced by such a response, dependent on bacteria-induced cross reactive autoantibodies to joint antigens.\textsuperscript{11} In a group of patients with spondyloarthropathies, including those with AS, IL10 serum levels, which can stimulate humoral responses and inhibit Th1 activity, correlated with disease activity.\textsuperscript{12} Although no separate data on patients with AS were given in this study it can be speculated that IL10-induced antibody production in AS (as, for example, in rheumatic fever) is not inhibited by Th2 activity but may even be stimulated. The association with bacterial infection is supported by the HLA-B27 predisposition found in AS and ReA (approximately 90\% and 50\% of the patients, respectively). The intra-articular presence of bacteria in ReA correlates with joint inflammation. If this is true for AS as well, effective clearance may require Th1-induced immunity and might be counteracted by Th2 activity.

The different effects of atopy-induced Th2 predominance on RA and AS raise the question as to what the impact of this Th2 mediated immune deviation is on other forms of arthritis such as ReA. The presence of Th2 activity in ReA has been thought to contribute to the remitting form of this arthritis in contrast with patients with RA who do not have a Th2 response and who have chronic arthritis.\textsuperscript{10} However, based on more recent studies it has been suggested that Th2 responses (together with IL10 production), which are detected in patients with ReA, may prevent effective clearance of arthritis-inducing bacteria and sustain the arthritis.\textsuperscript{13} The role of Th2 activity in this infection-induced arthritis becomes even more uncertain when one takes into account the fact that clearance of some bacteria (for example, \textit{Borreilia burgdorferi}) in Lyme arthritis) seems to require Th2-induced immunity.\textsuperscript{14} Studying the impact of atopic disorders on disease activity in patients with ReA may provide an insight into the role of Th2 polarisation in this arthritic response.

Together, the effect of Th2 mediated atopic disorders on RA and other Th1 mediated (common non-atopic) inflammatory conditions suggests a potential beneficial role of Th2 induction in these patients. In contrast, a similar approach seems less useful in the treatment of patients with AS. This emphasises the diversity of the different arthropathies and the importance of documenting the actual intra-articular inflammatory responses in relation to clearly defined disease characteristics of the separate arthritic forms. Investigation of the impact of naturally occurring Th2 mediated immune deviation on the various forms of arthritis may prove to be a tool to clarify the role of Th1/Th2 balances in these diseases. It might also clarify whether Th2 mediated diseases like atopic disorders have a favourable impact on the prevalence of several arthritic conditions, including RA, because this so far has not been convincingly proved. Because of the prevalence of the arthritic conditions, collaborative efforts are required because large numbers of patients need to be included.


