New insights into the functional role of the rheumatoid arthritis shared epitope

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ABSTRACT

The shared epitope (SE) – an HLA-DRB1-encoded 5-amino acid sequence motif carried by the vast majority of rheumatoid arthritis (RA) patients – is a risk factor for severe disease. The mechanistic basis of RA-SE association is unknown. This group has previously demonstrated that the SE acts as a signal transduction ligand that activates nitric oxide and reactive oxygen species production. SE-activated signaling depends on cell surface calreticulin, a known innate immunity receptor previously implicated in immune regulation, autoimmunity and angiogenesis. Recent evidence that the SE enhances the polarization of Th17 cells, which is a key mechanism in autoimmunity, is discussed highlighting one of several potential functional effects of the SE in RA.

1. Introduction

Rheumatoid arthritis (RA) affects 0.5–1.0% of the population [1]. The disease is characterized by chronic inflammatory changes in both articular and extra-articular tissues. Due to its high prevalence and debilitating nature, RA incurs a major economic burden on society. In recent years it has been realized that in addition to causing pain and disability, the disease significantly shortens life expectancy due to accelerated atherosclerosis [2].

Although genes play a major role in RA risk, the disease appears to have low sibling occurrence with a concordance rate of 12–15% in monozygotic twins [3]. Overall, the contribution of genetic factors to RA risk is calculated at approximately 60%, while the remaining 40% are believed to be contributed by environmental factors. The observations that RA is more common in urban versus rural populations, a recent decline in the incidence of the disease in high-incidence of populations, and the effect of birth cohort on disease incidence are all indirectly supporting environmental influences. Importantly, over the past few years it has been conclusively shown that the disease is strongly associated with environmental pollutants, such as cigarette smoking [4].

Among the genetic risk factors, the HLA-DRB1 locus is the most significant one. RA has long been shown to associate with human leukocyte antigen (HLA) genes. The pioneering studies on HLA-RA association were carried out in the late 1970s by Peter Stastny [5] and Robert Winchester’s group [6], who independently concluded that HLA-DR4 is significantly more common among patients with RA. It was subsequently found that other HLA-DR serotypes, for example, HLA-DR1 in Mediterranean, or HLA-DR14 in Native Americans, are also associated with the disease. With the advent of modern DNA sequencing techniques it had become apparent in the 1980’s that there is no RA-specific HLA-DR sequence. Instead, it was found that the majority of RA patients share a short sequence motif coded by several HLA-DRB1 alleles. This revelation had prompted the Shared Epitope Hypothesis [7].

2. The RA shared epitope

The term “shared epitope” (SE) most commonly refers to a five amino acid sequence motif in residues 70–74 of the DRβ chain coded by several HLA-DRB1 alleles that are over-represented among RA patients (Fig. 1). The SE motif consists of three homologous amino acid sequence variants: (1) QKRAA, the SE variant that is the most common motif among Caucasian, is coded primarily by the HLA-DRB1*0401 allele; (2) The second most common motif, QRRAA, is coded by several alleles, among them HLA-DRB1*0404, HLA-DRB1*0101, and HLA-DRB1*0405; (3) The third motif, RRRAA, coded by allele HLA-DRB1*1001, is the rarest. In addition to increasing RA risk, SE-coding HLA-DRB1 alleles have been shown to associate with more severe disease [8] and to exhibit allele-dose...
effect, i.e., patients with 2 SE-coding alleles tend to experience more severe disease than patients with 1 allele, who, in turn, have more severe RA than SE-negative patients.

The mechanism underlying the effect of the SE is unclear. Based on the known role of MHC class II molecules in antigen presentation, the prevailing paradigms postulate that presentation of arthritogenic self-peptides [9], molecular mimicry with foreign antigens [10], or T cell repertoire selection [11] are involved. While these hypotheses are all plausible, they are difficult to reconcile with the fact that data supporting antigen-specific responses as the primary event in RA are inconclusive. Additionally, several other human diseases have also been shown to be associated with SE-encoding DRB1 alleles, including polymyalgia rheumatica [12], giant cell arteritis [12], Type I diabetes [13], erosive bone changes in psoriatic arthritis [14] and lupus [15], autoimmune hepatitis [16] and early-onset chronic lymphoid leukemia [17], among other conditions. The SE is also associated with spontaneous arthritis in dogs [18] and, in HLA-DRB1*0401 transgenic mice it increases the incidence of spontaneous diabetes [19] and the severity of both collagen-induced arthritis (CIA) [20] and experimental autoimmune encephalomyelitis (EAE) [21]. Thus, although it is best known for its involvement in RA, the SE associates with several pathogenically unrelated diseases and experimental disease models, and its effect seems to lack antigen- or species-specificity. These promiscuities are incongruent with fundamental tenets of MHC-restricted antigen presentation theory.

3. Activation of innate signaling by the SE: a new paradigm

Given the inconsistencies of SE-RA association with antigen presentation-based theories, over the past few years, our laboratory has examined an alternative hypothesis concerning the role of the SE in RA [22–28]. Based the known tri-dimensional homology among products of the MHC gene family, we postulated that similar to class I MHC-coded molecules [29], the SE may be acting as a ligand that can trigger innate immune signaling. The rationale of this antithetic hypothesis relates to the fact that the SE is located near the apex of α helical tri-dimensional structural motif that has been preserved throughout the entire MHC gene family and seems to be enriched in signal transduction ligands.

The first crystal structure of a class II MHC molecule, published in 1993 by Don Wiley’s group [30], revealed a remarkable tri-dimensional similarity to a previously reported class I MHC molecule. The degree of the similarity was surprising, given a substantial evolutionary distance between the two molecules and the fact that the peptide-binding groove in class I molecules is coded by a single gene, while in class II it is formed jointly by the products of two distinct genes. The extent of evolutionary ‘choreography’ required to bring these two disparate MHC molecules to form a near-identical tri-dimensional structure, is staggering. One of the notable features of the similarity is a ‘kink’ in the α2 domain of the class I MHC molecule, which could be almost perfectly superimposed on a similar structure in the β1 domain of the class II molecule. The ‘kink’ region in both molecules involves allele-diversity regions. Subsequent crystal analyses have shown very similar tri-dimensional structures in the entire MHC gene product family, irrespective of whether or not they can present antigens [31]. In all cases, this region forms a sharp protrusion ‘above’ the MHC groove plane (Fig. 1).

The remarkable conservation of a similarly-shaped ‘kink’ in the midst of allele diversity regions in MHC molecules independent of their antigen presentation capabilities suggests that this region may possess important allele-specific, conformationally-dependent, non-antigen presentation functions. Indeed, there are some indications that this region performs such functions. For example, in both classical and non-classical (HLA-E) class I MHC molecules, this region contains ligands for natural killer (NK) cell receptors [32]: in HFE (an empty-grooved human class I-like molecule), it interacts with transferrin receptor [33]. In M10 (a mouse class I-like molecule), the same region has been proposed as an interaction site with a pheromone receptor [34].

These considerations have led us to pursue a novel hypothesis which postulates that similar to its structural homologue in the class I MHC molecule, the SE functions as a signal transduction ligand that interacts with an evolutionarily-conserved receptor. Our
studies [22–28] have indeed demonstrated that the SE, whether expressed in its native conformation on the cell surface; as a cell-free HLA-DR tetrameric molecule; engineered into large recombinant proteins; or as a short synthetic peptide, activated in all cases nitric oxide (NO)-mediated signaling in trans in a strictly allele-specific manner. A consensus motif comprising of the \(70^Q/R-K/R-x-x-A74^Q/R\) sequence was found to be necessary for triggering the signal.

Given the known pro-oxidative effect of NO and the proposed role of oxidative stress in the pathogenesis of RA, we have explored whether SE-triggered signaling can increase cellular oxidative stress. These studies have shown that cells exposed to cell surface SE-positive HLA-DR molecules, to cell-free recombinant proteins genetically engineered to express the SE motif, or to SE-positive synthetic peptide showed diminished cyclic AMP-dependent signaling, increased reactive oxygen species (ROS) levels, and higher vulnerability to oxidative DNA damage. The SE effect is critically dependent on amino acids Q/R70, K/R71 and A74 of the DRb chain. The pro-oxidative effect of the SE could be reversed by inhibiting NO production. Thus, these studies demonstrate that the SE acts as a signaling ligand that activates an NO-mediated pro-oxidative pathway.

SE-triggered signaling is transduced via cell surface calreticulin (CRT), a known innate immunity receptor. The role of CRT and the dependence on CD91 have been published [25]. In brief, cell surface CRT was identified as the SE-binding molecule using affinity chromatography purification, cell-binding assays, surface plasmon resonance (SPR) and time-resolved fluorescence resonance energy transfer techniques. SE-triggered signaling could be blocked by anti-CRT antibodies or antibodies against CD91 and by CRT-specific anti-sense or small interfering RNA (siRNA) oligonucleotides. Murine embryonic fibroblasts from \(C^−\)/ or \(c91^−\)-deficient mice failed to transduce SE-activated signals. Exogenously added soluble CRT attached to the cell surface and restored signaling responsiveness in \(C^−\)/ cells.

More recently we have mapped the SE binding site on CRT [28]. SPR experiments with domain deletion mutants suggested that the SE binding site is located in the P-domain of CRT. The role of this domain as a SE-binding region was further confirmed by a photoactive cross-linking and mass spectrometry methods. In silico analysis of docking interactions between a conformationally intact SE ligand and the CRT P-domain predicted the region within amino acid residues 217–224 as a potential SE binding site. Site-directed mutagenesis demonstrated involvement of residues Glu217 and Glu223 and, to a lesser extent, residue Asp220 in cell-free SPR-based binding and signal transduction assays.

4. Potential functional consequences of SE-activated signaling

As illustrated in Fig. 2, the SE acts as a signal transduction ligand that interacts with cell surface CRT and activates signal transduction in trans. Although the SE-CRT pathway has not been fully mapped, our data to date indicate that it involves activation of NO synthase, and production of ROS. We have also demonstrated that the SE-activated pathway blocks a cyclic AMP-mediated pathway [23].

NO is a ubiquitous signaling molecule with versatile effects in the immune system. In RA, increased NO levels correlate significantly with inflammatory markers [35] and anti-rheumatic agents have been shown to suppress NO production [35,36]. NO has also been implicated in the pathogenesis of experimental autoimmune models in mice. For example, SJL mice (H-2Q) are known for their NO overproduction [37]. These mice are susceptible to many autoimmune diseases, including EAE, myasthenia, myositis, inflammatory bowel disease and CIA [38], and their autoimmune tendencies are attributed to their NO overproduction [39]. Similar to human RA, SJL mice display aging-associated increase in disease incidence [40], excessive DNA damage [41], higher mutation rates [42] and higher incidence of spontaneous lymphoma [43].

There are many ways by which NO could contribute to RA pathogenesis. The salient cellular mechanisms by which SE-activated NO could be involved in RA are depicted in Fig. 3. For example:

- NO has been shown to modulate apoptosis, a process that has been extensively implicated in RA and other autoimmune conditions [44].
- NO plays a major role in angiogenesis [45], a key pathogenic mechanism in the inflammatory pannus. Blood vessels are critically important for both nourishing the proliferative synovial tissue and for the ingress of inflammatory leukocytes into the joint. Anti-angiogenic agents have been shown to modulate arthritis in animal models of RA [46]. Additionally, certain disease-modifying anti-rheumatic drugs, such as methotrexate or anti-TNF\(\alpha\) antibodies, have been found to be angiostatic [47]. Thus, SE-activated NO overproduction could conceivably enhance angiogenesis in RA.
- The pathogenesis of RA pannus involves activation of matrix metalloproteinases (MMPs), with MMP-13 being of particular interest. The relevance of this MMP to RA relates to its high potency and specificity for type II collagen [48]. In addition to its direct tissue degrading effects, MMP-13 is a likely contributor to RA pathogenesis due to its central position in the MMP activation cascade [49] and its pro-angiogenic effect [50]. MMP-13 is upregulated by inflammatory cytokines, such as IL-1, IL-6, TNF\(\alpha\) and IL-17 [51]. Consistent with the model discussed here, NO has been shown to increase MMP-13 expression and activity [52].
- Directly relevant to the focus of this review, NO has been recently shown to potently inhibit indoleamine 2,3 dioxygenase (IDO), a tolerogenic enzyme [53,54]. Our recent studies have indeed demonstrated that the SE, which trigger NO signaling in dendritic cells (DCs) has an inhibitory effect on IDO activation, with immune dysregulatory effect, both in vitro and in vivo (see below).
As Fig. 3 shows, SE-activated NO production leads to oxidative stress. The role of ROS in RA has been extensively studied [55]. There are many potential RA-relevant cellular and molecular mechanisms that could be affected by excessive production of ROS. Among them: protein and DNA damage, epigenetic modifications, telomere attrition, cell senescence and T cell hyporesponsiveness, all of which have been implicated in the pathogenesis of RA [56–61]. Among the many potential pathogenic effects, two ROS-mediated disease processes are of particular interest: atherosclerosis (AS) and bone erosions.

As mentioned above, RA patients have long been noticed to have shorter life expectancy. It is now becoming increasingly apparent that AS is the most common cause of premature death in RA with a relative risk of about 2, compared with age-matched control populations [2]. Several studies have noted the fact that AS risk in RA exists in a rate greater than would be expected from the profile of classical cardiovascular risk factors, such as diabetes mellitus, hypercholesterolemia, obesity or hypertension [62]. The prevalent hypothesis for the increased risk of AS in RA attributes the association to inflammation. However, there is substantial evidence to suggest that the inflammatory milieu may not be the sole culprit. For example, in RA patients without established cardiovascular risk factors, the erythrocyte sedimentation rate was found to lack correlation with the carotid intima-media thickness, a known biomarker of AS [63]. Additionally, in RA patients chronically treated with TNFα blockers, endothelial cell dysfunction continues to exist [64] and AS continues to progress [65] despite reversal of the inflammatory state and improvement of the arthritis. Finally, patients with many other inflammatory conditions are known to be at a higher AS risk, suggesting that disease-specific factors may contribute to the increased AS risk in RA. Given the unequivocal role of oxidative stress in AS on one hand, and anecdotal reports suggesting that SE association with AS may exist also in the non-RA population [66–68] on the other, the possibility that SE-activated ROS production might be a direct contributing factor to premature AS development in RA is worth consideration.

Osteoclast-driven destruction of juxta-articular bone is a hallmark of RA. The severity of bone destruction is RA has been linked to the SE [69] with indications for allele-dose effect in certain populations [70]. Interestingly, the SE has also been implicated in erosive changes in non-RA conditions, such as psoriatic arthritis [14], SLE [15] and periodontal disease [71]. Thus, the SE may be involved in bone destruction irrespective of the underlying disease. The mechanism discussed here involving SE-activated NO and ROS production might provide a mechanistic basis for these associations. It has been previously shown that NO activates osteoclasts in a bi-phasic dose-dependent fashion [72]. ROS, likewise, have been shown to activate osteoclastogenesis [73] and bone resorption [74]. Accordingly, one scenario to consider is that the SE may contribute to osteoclast activation by stimulating higher production of NO and ROS, thereby increasing osteoclast-mediated bone destruction.

5. A case in point: SE-activated immune dysregulation

Using the diagram shown in Fig. 3 as a ‘blueprint’, this laboratory has been investigating RA-relevant SE effects in several pathogenic system with encouraging results. Here we discuss our recent findings in one of the areas of our current research interest: SE-activated immune dysregulation.

As discussed in greater detail in a recent review [27] CRT has been previously implicated in immune regulation. CRT is a 60 kDa protein which is expressed on the surface of many cells [75,76] and functions as an important innate immune system receptor [77–79]. It serves as the signal-transducing receptor for members of the collectin family [80]. Collectins bind foreign organisms or apoptotic cells through their globular heads, while their collagen-like tails bind to cell surface CRT. This leads to CRT-dependent phagocytosis. Different from elimination of foreign organisms, events which are associated with an intense inflammatory reaction [81], safe elimination of apoptotic cells is critically dependent on suppressing the inflammatory response [82]. The decision whether a pro- or anti-inflammatory reaction should be activated depends on the presence or absence of second signals that are triggered uniquely by apoptotic cells [83]. Thus, CRT plays a pivotal role in the junction between tolerance and autoimmunity due to its critical role in elimination of apoptotic cells [80]. Aberrant activation of the CRT-mediated pathway can lead to autoimmunity as exemplified by conditions that involve defective CRT-mediated clearance of apoptotic cells [84].

Our research interest in the effect of the SE on DCs is partly based on the fact that these cells are known to express functional CRT receptors on their surface [76]. DCs are professional antigen presenting cells strategically positioned in the interface between the innate and adaptive immune systems. In addition to their role in antigen presentation, DCs also induce tolerance through a vari-
In DCs by the Th1 cytokine IFN-γ [85]. A growing body of evidence indicates that the tolerogenic effect of DCs is mediated partly by IDO, an enzyme that catalyzes the catabolism of tryptophan [86,87]. The precise mechanism by which IDO exerts its effect is unknown, but may involve tryptophan depletion and/or pro-apoptotic or anti-proliferative effects of tryptophan downstream metabolites [88]. IDO is inducible in DCs by the Th1 cytokine IFN-γ [89] and by Treg-expressed CTLA4 through ligation of cell surface CD80/CD86 molecules [90]. Activation of IDO in DCs by Treg has been shown to inhibit IL17-producing helper T (Th17) cells [91], a T cell subset that is believed to play a key role in autoimmunity (discussed below).

It is worth noting that in contrast to the IDO-inducing effect of IFN-γ and CTLA4, NO is a potent inhibitor of IDO [53,54]. Given our findings that the SE activates NO production in many cell types, including DCs, we examined whether the SE ligand could affect IDO enzymatic activity. Using murine fibroblast L-cells transfectants expressing structurally intact and functionally HLA-DR/β2m heterodimeric molecules through cDNA transfection we demonstrated that transfectants expressing SE-positive HLA-DR molecules on their surface produced significantly less kynurenine (an IDO-dependent tryptophan metabolite) in response to IFN-γ, compared to transfectants expressing SE-negative HLA-DR molecules. An identical pattern was seen when human fibroblasts were stimulated with a soluble SE ligand in the form of SE-expressing peptides. Importantly, the SE ligand IDO inhibitory activity was restricted to CD11c+CD8− DCs, a subset known to express IDO [92]. Thus, the SE ligand effectively and specifically inhibits the activity of the tolerogenic enzyme IDO in DCs.

In addition to IDO-mediated T cell regulation, DCs can affect immune reactions by production of cytokines that can activate or expand particular subsets of T cells. In mice, the combination of IL-6 and TGF-β facilitates differentiation of Th17 cells, while IL-23 is involved in the expansion of this subset [93]. Accordingly, we have studied supernatants of SE-stimulated DCs. Our data showed that in the CD11c+CD8− DC subset, but not in the CD11c+CD8+ subset, a SE peptidic ligand activated a robust production of IL-6. An SE-negative ligand did not trigger any cytokine production. Other cytokines (IL-4, IL-10, IL-12, IL-17, TGF-β) did not show any increased production, indicating the specificity of SE effect. Interestingly, IL-23 levels in DCs did not increase following stimulation with the SE ligand. However, in the presence of suboptimal concentrations of LPS (100 ng/ml), the SE had a prolonged synergistic effect on IL-23 production. The effect was specific for IL-23, since no synergism was found in the production of another LPS-inducible cytokine, IL-6.

It has been previously demonstrated that IDO inhibition [94] or increased IL-6 levels [95] inhibit Treg cells. As discussed above, the SE inhibited IDO activity in CD11c+CD8+ DCs and increased IL-6 production in CD11c+CD8− DCs. We therefore determined whether the SE interferes with Treg differentiation. Mouse CD11c+DCs were first incubated overnight with SE-positive or SE-negative peptides, or with medium. DCs were then co-cultured with purified syngeneic CD4+ T cells or CD4+CD25+CD62L−CD44− naïve T cells in the presence of TGF-β and anti-CD3 Ab. After 5 days, CD4+CD25+Foxp3+ Treg abundance was determined by FACS analysis. Our data showed that the SE-positive, but not an SE-negative ligand, significantly inhibited Treg cell differentiation. Similarly, splenic CD4+CD25+CD62L−CD44− naïve T cells cultured with CD11c+DCs, treated with a SE-positive HLA-DR tetramer, but not with SE-negative HLA-DR tetramers demonstrated markedly reduced Treg cell differentiation.

Since SE ligands were found to enhance production of Th17-promoting cytokines, IL-6 and IL-23, we determined whether they can facilitate Th17 differentiation. To this end, CD11c+DCs were first stimulated overnight with SE-expressing peptides or tetramers. CD4+CD25−CD62L−CD44− naïve T cells were added and cultured in the presence of a Th17-polarizing cocktail of cytokines and antibodies. After 6 days, cells were collected and analyzed by flow cytometry. The results showed that SE-positive ligands, in particular when presented as HLA-DR tetramers, had a robust enhancement of Th17 differentiation. The SE showed similar effect when expansion of Th17 cells was studied [26].

To determine the biologic significance of the in vitro data shown above, we have undertaken to characterize the SE polarizing effect in vivo. Our findings showed that mice immunized with collagen type II in the presence of a cell-free SE ligand displayed much higher abundance of Th17 cells in the draining lymph nodes, compared to mice immunized with the same antigen in the presence or absence of a control ligand. Similarly, splenocytes from mice immunized with collagen in the presence of the SE ligand produced much higher levels of IL-17 compared to the control groups [26]. Thus, the SE facilitates Th17 polarization both in vitro and in vivo.

It is worth noting that IL-17 has been shown to enhance several pro-arthritogenic processes, including angiogenesis, MMPs production, osteoclastogenesis, leukocyte recruitment and inflammation [96]. Both IL-17-producing cells and IL-17 are abundantly expressed in the RA joint [97], and neutralizing IL-17 prevents experimental arthritis development, while deficiency of Treg cells has been shown to increase cellular and humoral immune responses and disease severity in CIA [98]. Thus, SE-activated Th17 polarization, observed by us both in vitro and in vivo, suggests a potential mechanism by which the SE could affect RA pathogenesis.

6. A proposed model

Different from the prevailing paradigms, which attribute the role of the SE in RA to presentation of a putative self or foreign antigen, the considerations discussed above and our experimental findings implicate an allele-specific, antigen presentation-independent mechanism. We have shown that the SE activates RA-relevant signaling events in several immune and non-immune cell types. The SE signaling effect is independent of the antigen presentation function of the parent HLA-DR molecule and could be seen using synthetic SE ligands, which do not possess any antigen presentation capability. It is worth mentioning, however, that using X-ray crystallography-based analyses others have predicted that amino acid residues K 71 [99], or Q 70 and K 71 [100] of the SE protein could participate in the binding to human collagen type II epitopes [101]. We therefore determined whether (1) The identity of the target antigen in RA is unknown. The role of collagen type II as a target self antigen has been disputed by many. It is therefore unclear whether the possible interactions between SE residues and collagen II epitopes are pathogenically relevant; (2) Different groove peptides may have different affinities to the SE. It is therefore conceivable that CRT could displace certain types of peptides; (3) Another scenario to consider is that CRT interacts primarily with ‘empty’-grooved HLA-DR molecules; (4) Our preliminary data using SE-positive HLA-DR tetrameric molecules loaded with two distinct groove peptides suggest that the identity of the peptide does not have any significant impact on SE-CRT interaction or signaling (unpublished results); (5) Finally, as discussed below, the antigen presentation paradigm and the SE signaling hypothesis are not mutually exclusive. We cannot rule out a scenario in which certain groove peptides and CRT could co-interact with the SE.

Based on the experimental and theoretical considerations discussed above, we propose the following model: In healthy individuals carrying SE-coding HLA-DRB1 alleles, the SE ligand, expressed on antigen presenting cells or lymphocytes interacts at low affinity with cell surface CRT. As our data show, SE activation of DCs pro-
motes Th17 polarization, which in healthy individuals could be advantageous against pathogens. We further propose that over time, due to environmentally-triggered stochastic events, the affinity of SE-CRT interaction could increase with resultant aberrant activation of the SE pathway, excessive Th17 polarization and development of RA. It is worth mentioning here that different from humans, mice do not express class II MHC molecules on the surface of activated T cells. We therefore posit that the reported SE-promoted autoimmunity in HLA-DRB1 transgenic mice could be due to activation of pro-inflammatory signaling by SE expressed on cells other than T lymphocytes (i.e., macrophages, B lymphocytes or DCs).

This model is non-exclusive with other hypotheses in the field. One scenario to consider is that while SE-expressing HLA-DR molecules are uniquely capable of presenting joint-specific antigens, thereby determining the tissue-specificity of the immune response and its anatomical distribution, it is the SE ligand function that determines the outcome of this response by facilitating Th17 polarization. Finally, we wish to clarify that given the focus of this review on Th17, the simplified model proposed here highlights the SE effect on DCs. However, the SE could conceivably activate RA-associated functional consequences in other cell systems (some of which are listed in Fig. 4).

7. Summary

We have previously demonstrated that the SE acts as a signal transduction ligand that activates NO and ROS production in other cells. SE signaling is transduced by cell surface CRT, a known innate immunity receptor previously implicated in immune regulation, autoimmunity and angiogenesis. There are multiple RA-relevant cellular and molecular mechanisms that could conceivably exert pathogenic influences secondary to activation of the newly discovered pathway. An example of SE-activated aberrant immune regulation that involves inhibition of the tolerogenic enzyme IDO on one hand and enhanced Th17 expansion on the other is presented here for illustration. Other conceivable SE-activated pathogenic mechanisms are being currently studied by this group.

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