Stimulation of autoimmunity by toll-like receptor ligands

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Healthy individuals are known to harbour self-specific T cells against a variety of immunodominant self-peptides—for example, myelin-basic protein (MBP) peptides. Yet, autoimmunity in a given individual is a rare event indicating that the self-reactive cells are somehow prevented from inflicting damage.1 Various experiments in which transgenic animals express both peripheral antigen and its cognate T cell receptor (TCR) have demonstrated that these self-antigens and their respective T cells are able to coexist without eliciting tissue damage.2–4 Similarly, in patients with tumours that express specific antigens the tumours continue to grow in the presence of antigen specific T cells.5–7 These studies therefore suggest that the mere coexistence of an antigen and its cognate T cell is not in itself sufficient to produce an immune response which can subsequently eliminate the cells expressing the self-antigen. This coexistence raises several crucial questions—for example:

1. What does it take to shift the balance from tolerant coexistence of autoreactive T cells and self-antigens to autoimmune destruction?
2. Are the mechanisms underlying this process the same for autoimmunity and tumour destruction?
3. Can we exploit this knowledge to devise specific immunotherapies against autoimmune disorders and tumours?

Anergy and regulatory T cells are known mechanisms capable of preventing activation of self-reactive T cells. In this report, we present evidence that additional mechanisms are operative within the innate immune system. In particular, toll-like receptor (TLR) activation is required for destruction of tissue by activated autoreactive T cells.

A TWO STEP MODEL FOR ORGAN SPECIFIC AUTOIMMUNITY

Several years back, we asked the simple question whether overcoming self-tolerance and activation of self-reactive cells would be sufficient to cause tissue damage. For this purpose, mice were generated expressing major histocompatibility complex (MHC) class I Kb molecules as a model antigen outside the thymus and exclusively on hepatocytes owing to the use of the liver specific albumin or C-reactive protein promoters.8 In addition, the mice were transgenic for a T cell receptor (Des.TCR) with specificity for the Kb molecule. The rationale for using a class I molecule as an autoregion is that MHC class I molecules present peptides from endogenously synthesised antigens, and that the MHC class I peptide complex is not supposed to travel to other cells in a form that can still be recognised by Des.TCR T cells. Thus, T cells have to interact directly with the cell expressing the MHC I molecule. These double transgenic mice harbour large numbers of Kb reactive CD8 T cells, which are, however, tolerant as indicated by their failure to reject a Kb positive tumour or skin grafts.7 In this model, TCR unresponsiveness can be reverted to a responsive phenotype by simultaneous challenge with Kb and interleukin (IL)-2, thus demonstrating breakage of tolerance towards a peripheral antigen in vivo.9 Mice in which tolerance was reversed were then able to reject Kb positive grafts. Surprisingly, despite activation of self-reactive T cells and the rejection of Kb positive grafts, there was no autoaggression against Kb positive hepatocytes. Histological inspection revealed that Kb reactive T cells failed to infiltrate the Kb positive liver parenchyma. Thus, additional factors appear to be involved in extravasation and effector function of self-reactive lymphocytes.

Autoimmune diseases are frequently preceded by infections, which besides mimicking self-antigens may also cause inflammatory responses in the target organ. To investigate the importance of inflammatory responses as a cause of autoimmune pathology following breaking of tolerance, the double transgenic mice were treated with proinflammatory inducers such as Listeria, γ irradiation, or TLR9 stimulation by cytosine-phosphorothioate-guanine-rich oligodeoxynucleotides (CpG-ODN).10 Indeed, now the autoreactive T cells massively infiltrated the liver parenchyma and caused tissue damage. We conclude that activated self-reactive T cells do not cross an unstimulated endothelial barrier in numbers sufficient for tissue damage. Instead, a proinflammatory environment “opens” the endothelial barrier, thus allowing T cell infiltration and tissue destruction. Thus, the two step model of organ specific autoimmunity comprises an initial activation of autoreactive T cells and the presence of proinflammatory signals in the target organ for increased extravasation of self-reactive T cells resulting in tissue damage.

The scenario makes sense if one considers that the immune system wants to focus T cells at the site of an inflammatory response.

REPEATED INFLAMMATORY RESPONSES ALONE CAN CAUSE AUTOAGGRESSION

Liver dendritic cells, Kupffer cells, and liver sinusoidal endothelial cells among resident liver cells are known to express TLR9, explaining the strong effect of CpG-ODN on the liver microenvironment. The proinflammatory response in the liver was found to induce expression of costimulatory molecules, adhesion molecules, and to increase the amount of MHC molecules in hepatocytes, thus generating a phenotype similar to that of antigen presenting cells. Indeed, these activated hepatocytes were able to activate Kb specific T cells, which then, in turn, attacked the hepatocytes and caused transient liver damage.11 Two important conclusions can be drawn from this observation. Firstly, T cell priming can take place outside lymphoid organs, provided that the microenvironment and stimulatory conditions are appropriate. Secondly, T cells activated by inflamed non-lymphoid tissue can cause autoaggression. Thus, an imbalance in the microenvironment of an organ may be sufficient to cause autoaggression (fig 1). It should be noted, however,

Abbreviations: CpG-ODN, cytosine-phosphorothioate-guanine-rich oligodeoxynucleotides; MHC, major histocompatibility complex; TCR, T cell receptor; TLR, toll-like receptor

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that the autoaggression observed here so far is only transient and does not result in serious autoimmune damage. Similar results supporting ours have recently been reported by Lang et al.11 who show that TLR ligands conditioned the target organ via the induction of type I interferon, enabling activated T cells to become autoaggressive.

TLR9 STIMULATION AND TUMOUR IMMUNITY

Autoimmunity and tumour immunity can be regarded as comparable immune reactions, as they are both directed against a self-organ or self-organ-like tissue. In order to see whether the above mentioned two step model of autoimmunity (activation of self-reactive T cells plus proinflammatory response) would also apply to tumour immunity, we used Rip.Tag tumour mice. These mice express the SV40 T antigen under the control of the rat insulin promoter (Rip) and develop insulinomas. The tumour nodules are characterised by a highly irregular vasculature and lack of cellular infiltrates.12 13 Attempts to eradicate the tumours by vaccination or transfer of activated Tag specific T cells failed. Immunohistologic examination demonstrated that even the activated T cells did not transmigrate into the tumour tissue. To see if a proinflammatory stimulus would facilitate tumour rejection, we transferred tumour (Tag)-reactive CD4 and CD8 T cells into tumour bearing Rip.Tag mice in combination with CpG-ODN as proinflammatory stimuli. Interestingly, induction of the local inflammatory milieu by CpG-ODN “opens” the endothelial tumour barrier, thereby permitting tumour eradication. Interestingly, induction of the local inflammatory milieu by CpG-ODN may modify the tissue in a way that otherwise non-professional cells acquire the ability to activate T cells outside lymphatic tissues. Similar modifications in tumours by TLR-L may help to stimulate T cell dependent tumour eradication.

CONCLUDING REMARKS

The data presented here demonstrate that activation of self-reactive T cells is not necessarily sufficient to cause autoimmunity—for example, autoimmune hepatitis. An additional step is required, namely activation of endothelia, which facilitates infiltration of autoactivity cells into the liver parenchyma. Activation of endothelium can be achieved by the TLR9 ligand CpG-ODN. A similar mechanism is observed in tumour immunity. Tumour reactive T cells sometimes fail to infiltrate tumours. However, a proinflammatory stimulus such as CpG-ODN “opens” the endothelial tumour barrier, thereby permitting tumour eradication. Interestingly, induction of the local inflammatory milieu by CpG-ODN may modify the tissue in a way that otherwise non-professional cells acquire the ability to activate T cells outside lymphatic tissues. Similar modifications in tumours by TLR-L may help to stimulate T cell dependent tumour eradication.

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