TLR-4 ligation of dendritic cells is sufficient to drive pathogenic T cell function in experimental autoimmune encephalomyelitis

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Abstract

Background: Experimental autoimmune encephalomyelitis (EAE) depends on the initial activation of CD4+ T cells responsive to myelin autoantigens. The key antigen presenting cell (APC) population that drives the activation of naïve T cells most efficiently is the dendritic cell (DC). As such, we should be able to trigger EAE by transfer of DC that can present the relevant autoantigen(s). Despite some sporadic reports, however, models of DC-driven EAE have not been widely adopted. We sought to test the feasibility of this approach and whether activation of the DC by toll-like receptor (TLR)-4 ligation was a sufficient stimulus to drive EAE.

Findings: Host mice were seeded with myelin basic protein (MBP)-reactive CD4+ T cells and then were injected with DC that could present the relevant MBP peptide which had been exposed to lipopolysaccharide as a TLR-4 agonist. We found that this approach induced robust clinical signs of EAE.

Conclusions: DC are sufficient as APC to effectively drive the differentiation of naïve myelin-responsive T cells into autoaggressive effector T cells. TLR-4-stimulation can activate the DC sufficiently to deliver the signals required to drive the pathogenic function of the T cell. These models will allow the dissection of the molecular requirements of the initial DC-T cell interaction in the lymphoid organs that ultimately leads to autoimmune pathology in the central nervous system.

Keywords: Multiple sclerosis, Experimental autoimmune encephalomyelitis, Dendritic cells, Myelin basic protein
Thus, one possible reason for the lack of a robust DC-driven model of EAE may be that LPS-activated DC cannot provide all of the signals to the naïve myelin-response T cells to engender strong autoaggressive function.

To formally test this possibility, we sought to induce EAE using bone marrow-derived DC (BMDC) that had received only the TLR4-signal in vitro, by activation with LPS. To maximize our chances of success we first sought to obviate two possible confounding issues – the size of the naïve myelin-responsive repertoire and the ability of the DC to present the autoantigen. We therefore started by developing a transgenic two-cell transfer model. We have previously reported that C57BL/6xB10.PL mice are resistant to EAE induction using the Ac1-9 peptide of myelin basic protein (MBP), unless they are first seeded with a cohort of naïve MBP-responsive T cells derived from the Tg4 T cell receptor (TCR) transgenic mouse [6]. This approach therefore allowed us to control the size of the Ac1-9-responsive T cell repertoire. To remove the possibility that BMDC may for some reason fail to present the Ac1-9 peptide efficiently, we made use of a second transgenic line, the AMK35 mouse, in which major histocompatibility molecule (MHC) class II-expressing cells constitutively express the Ac1-9 peptide, covalently bound to the AU molecule [7].

BMDC generated from AMK35 mice showed elevated expression of AU, CD80 and CD86 after overnight exposure to LPS (Figure 1A) and supernatants from these cultures contained elevated levels of pro-inflammatory cytokines (Figure 1B), notably IL-1β, IL-6 and IL-23, each of which has been described as being required for EAE development using gene-deficient mice [8-10]. These LPS-activated AMK35 DC also efficiently induced (LPS). Thus, one possible reason for the lack of a robust DC-driven model of EAE may be that LPS-activated DC cannot provide all of the signals to the naïve myelin-response T cells to engender strong autoaggressive function.

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the in vitro proliferation of naïve Tg4 CD4+ T cells (Figure 1C), and their production of IFN-γ, IL-17, TNF-α and GM-CSF (Figure 1D), without any addition of exogenous antigen to the culture. Thus, LPS-activated DC appeared to have many of the key attributes associated with EAE pathology. To test this in vivo, we seeded C57BL/6xB10.PL mice with a cohort of naïve Tg4 CD4+ T cells and one-day later subcutaneously administered LPS-activated AMK35 DC. Ten out of 12 mice developed clinical signs of EAE (Figure 2A). However, the extent and, importantly, the timing of onset of clinical signs showed variation.

The use of BMDC derived from mice in which antigen presentation is driven by a transgene, of course, has limitations. For example, it can be envisaged that it would be desirable to study DC expressing multiple autoantigenic peptides, either separately or together. To extend our study in this direction we therefore generated BMDC from Aα− non-transgenic BMDC, from B10.PL mice. These BMDC were activated with LPS and loaded with the Ac1-9(4Tyr) peptide in vitro. This variant peptide binds to the Aα molecule with high affinity, allowing stable antigen presentation (unlike the wild type Ac1-9 peptide). These antigen-loaded non-transgenic DC could efficiently activate naïve Tg4 T cells in vitro (Figure 2B) and induced a robust disease course in the T cell/DC transfer model in C57BL/6xB10.PL mice (Figure 2B). Of note, there was a trend for the onset of disease to be earlier and more synchronous than achieved with AMK35 DC (Figure 2C).

The two novel models of DC-driven EAE we have described have a number of advantages over the few earlier reports of EAE following transfer of autoantigen-loaded DC. Previous studies using BMDC pulsed with the 35–55 peptide of myelin oligodendrocyte glycoprotein (MOG) in C57BL/6 mice have required repeated injections of BMDC to induce EAE [3] or have required concurrent administration of CFA to drive robust disease [5]. A previous report of EAE involving MBP-pulsed DC differed markedly from ours in that five times as many...
MBP-reactive T cells were transferred and the host mice were irradiated prior to cell transfer [4]. Of note, the wild type MBP peptide (lacking the insertion of a Tyr residue at position 4) has such low affinity for $\mathrm{A}^\mathrm{u}$ that APC cannot be pulsed with that peptide to allow efficient loading into the $\mathrm{A}^\mathrm{u}$ peptide binding groove [13]. It is therefore difficult to reconcile that observation with the reported ability of DC loaded with the wild type peptide to activate MBP-responsive T cells for the initiation of EAE [4]. Unlike that previous report, we used two systems in which the presentation of the autoantigenic peptide-MHC complex by the DC was highly stable. AMK$_{35}$ DC, expressing transgenic peptide-MHC [7] induced variable degrees of EAE, particularly in terms of day of onset, but this difficulty was overcome using non-transgenic DC pulsed with the Ac1-9(4Tyr) peptide, which shows extraordinary affinity for the $\mathrm{A}^\mathrm{u}$ molecule [14]. DC that are loaded with this peptide in vivo can maintain expression of the peptide-MHC complex for at least seven days (SMA, unpublished). Given that the consensus is that BMDC probably persist for only a few days after administration, we can be confident that the DC we transferred were presenting the peptide to T cells for a sustained length of time. It is conceivable that the MOG(35–55)-$\mathrm{A}^\mathrm{b}$ complex might be prone to more rapid degradation in DC transfer experiments. Nevertheless, it is evident from other studies that the use of MOG(35–55)-loaded C57BL/6 DC can be sufficient to provide TCR-signaling in vivo, as we ourselves have shown [15,16], but it was not sufficient to trigger robust EAE, which still required subsequent immunization with CFA in those models.

A key paradigm is that autoreactive T cells can develop into autoaggressive T cells if they recognize (self, or cross-reactive non-self) antigen presented during infection (that is, on activated APC). The use of CFA to induce EAE mimics infection due to the presence of heat-killed mycobacteria. However, CFA contains multiple moieties capable of stimulating APC through a range of PRRs. Although LPS is not one of these moieties, heat shock proteins 65 and 70 have been described as TLR-4 agonists [17], as has pertussis toxin [18], which is routinely used during EAE induction. Less complex adjuvants, including LPS, have been reported to be able to substitute for CFA [19–22], but results have been variable between studies. Indeed, exposure to LPS can protect against EAE if it occurs before immunization with autoantigen in CFA [23]. Those studies have not provided clear information on which APC need to be activated through a given PRR to drive pathology, or which of the multiple consequences of PRR-ligation are non-redundant for the induction of EAE. Furthermore, it is clear that expression of the same TLR on different immune cells can have markedly different effects on disease outcome [24,25].

From our experiments described here, we can conclude that a) DC are sufficient as APC to effectively drive the differentiation of naïve myelin-responsive T cells into autoaggressive effector T cells and b) exposure to a TLR-4 agonist can activate the DC sufficiently to deliver the signals required to drive the pathogenic function of the T cell. These new model systems will therefore allow comparison of a range of PRR stimuli upon the DC and the use of gene-knockout/knockdown DC to help to probe the key molecular events in the initial DC-T cell dialogue that lead ultimately to autoimmune CNS inflammation in EAE.

Abbreviations

APC: Antigen presenting cell; BMDC: Bone marrow derived dendritic cell; CFA: Complete Freund’s adjuvant; CNS: Central nervous system; DC: Dendritic cell; EAE: Experimental autoimmune encephalomyelitis; ELISA: Enzyme-linked immunosorbent assay; FACS: Fluorescence activated cell sorting; GM-CSF: Granulocyte-macrophage colony-stimulating factor; IFN: Interferon; IL: Interleukin; LPS: Lipopolysaccharide; MBP: Myelin basic protein; MHC: Major histocompatibility complexes; MOG: Myelin oligodendrocyte glycoprotein; PRRs: Pattern-recognition receptors; TCR: T cell receptor; TLR: Toll-like receptor; TNF: Tumor necrosis factor; Tyr: Tyrosine.

Competing interests

The authors declare that they have no competing interests.

Authors’ contributions

RJM, ASM and SMA designed the experiments. RJM, HC, DT, ROC and MDL performed the experiments. RJM and SMA drafted the manuscript. FCK and BA generated the AMK$_{35}$ transgenic mouse. All authors have read and approved the final manuscript.

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