Little is known about the potential role of T cells in the inflammatory renal disease glomerulonephritis (GN). GN has been historically viewed as a product of immune complex–mediated complement activation, and the presence of autoantibodies made identifying T cell–specific effector contributions difficult to elucidate. In this issue of the *JCI*, Heymann et al. generate what they believe to be a novel, transgenic murine model of GN, demonstrating a direct role for CD8+ T cells, activated CD4+ T cells, and DCs in the pathogenesis of GN (see the related article beginning on page 1286).

The glomerulonephritides represent a diverse pathological and clinical subset of kidney diseases sharing the common endpoint of glomerular inflammation. It has been long assumed that deposition of immune complexes and subsequent complement activation were requisite steps in initiating disease. Recent data using mouse models, however, suggest that the pathogenesis of glomerular injury is more complicated. Mice deficient in various complement proteins still generate inflammation in a classic model of immune complex–triggered inflammation, the Arthus reaction (1–3). The absence of certain Fc receptors, such as FcγRIII, abrogates glomerulonephritis (GN), even in the presence of immune complex deposition and complement activation in glomeruli (4), via loss of neutrophil (5) and macrophage (6) effector functions.

Over the last ten years, evidence has been slowly accumulating that suggests a potential role for T cells in GN (7–9). Nevertheless, the idea of a causal role for T cells in GN remains weakly accepted, since much critical information is lacking. For example, we still do not know what role these cells play in the induction of GN or what the potential autoantigens are. In this issue of the *JCI*, Heymann and colleagues take our understanding of GN a considerable step forward by developing

a model of experimental GN based on two model antigens (10). The authors generated transgenic mice expressing the model antigens OVA and hen egg lysozyme in kidney podocytes. This allowed for the use of the well-characterized OVA-transgenic T cells: OVA-specific CD8+ T cells (OT-I cells) and OVA-specific CD4+ T helper cells (OT-II cells). The model allowed them to establish a causal role for T cells in mediating glomerular destruction.

**Direct role of T cells in glomerular immunopathology**

Prior to this study (10), experimental autoimmune GN (EAG) represented the most relevant model for examining the contribution of T cells to GN. In this model, mice immunized with purified glomerular basement membranes (GBMs) developed GN. With the identification of the Goodpasture antigen (11)—the noncollagenous domain of the α3 chain of type 4 collagen (α3IV-NC1)—a specific antigen could also be used to induce disease. While initial studies demonstrated that transfer of serum from immunized animals could recapitulate disease, depletion of CD4+ T cells attenuated GN pathology and autoantibody production (12). These studies were interpreted to support a primary role for autoantibodies, with T cells functioning mainly to provide B cell help (7, 8). Subsequent work suggested that T cells played a requisite role in pathogenesis, as serum could not initiate disease in T cell receptor–deficient animals (9). Furthermore, adoptive transfer of GBM-specific T cells generated in immunized mice conferred GN in recipient mice (9).

Interference with the function (13), activation (14), or adherence (15) of T cells was also shown to attenuate EAG activity. The exact role of T cells in glomerular immunopathology is unclear, but two possible mechanisms include macrophage recruitment to the inflamed kidney mediated by Th1 cells (16) and epitope spreading caused by T cell–mediated damage and antigen release (17).

To begin analyzing the specific roles of T cells, Heymann et al. (10) first injected naïve, OVA-specific CD8+ T cells into their transgenic mice. These cells proliferated in kidney draining lymph nodes, suggesting that OVA expressed on podocytes is taken up by DCs that migrate to the draining renal lymph nodes. These activated CTLs, however, did not cause disease, presumably because they were unable to traffic back into the noninflamed kidney. The authors then tested whether the addition of CD4+ T cells to the CD8+ T cells would allow for the development of GN. While the addition of naïve CD4+ T cells had no effect, the addition of activated OVA-specific CD4+ T cells with the naïve CD8+ T cells resulted in accumulation of both CD8+ and CD4+ T cells in the kidney and development of GN.

Why the addition of ex vivo–activated CD4+ T cells was required in the current study is unclear (10), but a similar requirement has been reported in a model of diabetes in which OVA is expressed in pancreatic β cells (18–20). This could be related to the findings of Heymann et al. (10) that naïve, OVA-specific CD4+ T cells did not proliferate when transferred into the transgenic mice, explaining the need to activate these cells ex vivo. The inability to activate CD4+ T cells in vivo could be related to low antigen levels or to the type of DC carrying the podocyte antigens to the lymph node. Hopefully, this issue will be explored in future experiments. It will also be important to test the general conclusions of their study using another model antigen system.
Nonetheless, the data reported by Heymann et al. (10) clearly demonstrate that the infiltration of activated CD4+ T cells into the kidney is required for the trafficking of CTLs to the kidney. One mechanism for the recruitment of CTLs explored by the authors is the production of chemokines capable of recruiting CTLs (CCL3, CCL4, and CCL5). Indeed, they found that antigen-specific CD4+ T cells purified from the kidney produced greater amounts of Ccl3, Ccl4, and Ccl5 mRNA. As the CD4+ T cells in the kidney were also producing Th1 cytokines, the authors imply that this subset may be important to confer renal pathology. Nevertheless, it will be important to directly test this using other Th subsets, such as Th2 or Th17, and Tregs.

The beauty of the model is that it possesses tremendous flexibility in allowing the contribution of various Th cells in GN to be determined. In vitro differentiated Th subsets can be administered, yielding important conclusions regarding the role of Th subsets in autoimmune pathology.

**DCs play a central role in mediating periglomerular infiltrates**

DCs are a functionally diverse class of antigen-presenting cells found in many tissues that possess tremendous immunostimulatory potential due to their ability to activate T cells. DCs are constitutively present in the glomeruli (21), but their role in GN prior to the development of the model reported in this issue by Heymann et al. (10) was unknown. A striking observation made in this study was the absolute requirement for DCs within the kidney for CTL-mediated disease. Depletion of diphtheria toxin–sensitive DCs after establishment of GN completely reverses immune infiltration pathology and restores normal glomerular microanatomy.
addition of activated CD4+ T cells resulted in upregulation of CD40, CD86, and IL-12 on the kidney DCs, the authors additionally suggest that the activated CD4+ T cells can induce DC maturation. What mediates the recruitment of activated CD4+ T cells to the kidney? The authors suggested that activated CD4+ T cells home to kidneys but do not stay due to the absence of CTL-mediated antigen release from the glomeruli. Once CTLs induce glomerular damage, kidney DCs present antigen to CD4+ T cells and retain them in the kidney. More importantly, it will be valuable to determine what kind of DCs are involved and exactly what their role is. While the glomerular DCs might seem to be the most likely DC type involved in the processing of podocyte antigens, the periglomerular location of the inflammation suggests that the involvement of DCs is more complicated. The authors suggest that since there is a large increase in monocyte-derived inflammatory DCs (CD11c+/CD11b+/Gr1+1) in the inflamed kidney, inflammatory DCs and not resident DCs are the likely culprit. But much more work, including kinetic studies, is necessary to demonstrate this point.

From their data (10), the Heymann et al. deduce the following testable model (Figure 1). First, the autoantigen, a membrane-tethered form of OVA (mOVA) expressed on podocytes, is taken up by DCs that then traffic to draining renal lymph nodes, in which they can activate naive CD8+ T cells. Second, activated CD4+ T cells home to the kidney, presumably via the recognition of OVA on antigen-presenting cells in the kidney. Third, recognition of antigen induces the expression of chemokines (CCL3, CCL4, and CCL5) that function to recruit CTLs to the kidney. Fourth, activated CTLs generate tissue damage, causing more antigen release. Fifth, released antigen is taken up by DCs that present antigen to CD4+ T cells in the context of a model antigen, the report by Heymann et al. (10) continues to modify our understanding of T cell immunity in GN. This model represents the clearest example of the causative roles T cells play in GN pathology. Additionally, the data suggest that therapies aimed at DCs may be a promising approach for the control of acute and chronic inflammation. The design of new therapies for treating glomerular injury will need to account for cell-mediated contributions, thanks to clever models such as this.

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It is important to note also that the nephritis promoter is active in lymph nodes, including in DCs (22), which could render the podocyte-specific expression of antigen irrelevant. However, OT-I cells do not proliferate in nondraining lymph nodes, making it unlikely that antigen resides constitutively in the lymph nodes of these mice.

**A renewed interest in examining T cell–mediated damage in GN**

The data generated from this model (10) demonstrate that T cells and DCs can mediate GN in the absence of immune complex–mediated pathologies. DCs control the presence of T cell infiltration, while CD4+ T cells potentiate the severity of cellular infiltration and CD8+ T cells confer glomerular injury. Activated macrophages also infiltrate the kidney in a T cell–dependent manner, providing additional effector mechanisms for glomerular injury.

Although much can be learned from this model, podocyte-derived antigen expression may not represent a physiologically relevant system for examining GN. Virtually all putative antigens thought to play a role in GN trap themselves against the GBM, as with nucleic acids, or are part of the GBM itself. It remains to be determined whether GN induced by podocyte antigens shows similar immunopathologies to that induced by GBM-based antigens. It will also be important to determine how autoantibodies and immune complexes may contribute to the cellular mechanisms studied here.

As what is believed to be the first model to test the role of T cells in the context of a model antigen, the report by Heymann et al. (10) continues to modify our understanding of T cell immunity in GN. This model represents the clearest example of the causative roles T cells play in GN pathology. Additionally, the data suggest that therapies aimed at DCs may be a promising approach for the control of acute and chronic inflammation. The design of new therapies for treating glomerular injury will need to account for cell-mediated contributions, thanks to clever models such as this.

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