# Molecular Analysis of the Helper T Cell Response in Murine Interstitial Nephritis

# T Cells Recognizing an Immunodominant Epitope Use Multiple T Cell Receptor V $\beta$ Genes with Similarities across CDR3

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### Abstract

Anti-tubular basement membrane disease ( $\alpha$ TBM disease) produces T cell-mediated interstitial nephritis in SJL mice after immunization with renal tubular antigen. Initial mononuclear infiltrates appear in vivo after several weeks, with the subsequent progression to renal fibrosis and end stage renal disease over many months. We have analyzed the fine specificity of the autoreactive helper T cell repertoire in  $\alpha$ TBM disease through the isolation and characterization of a panel of CD4<sup>+</sup> Th1 clones harvested after 1-2 wk from animals immunized to produce disease. All clones capable of mediating  $\alpha$ TBM disease are directed towards a 14-residue immunodominant epitope (STMSAEVPEAASEA) contained within the target antigen, 3M-1. Evaluation of the T cell receptor (TCR) V $\beta$  repertoire used by these autoreactive T cells reveals the use of several V $\beta$  genes, but with some preference for V $\beta$ 14. Sequencing across the putative CDR3 region of the TCR  $\beta$  chains suggests that common amino acids at the V $\beta(N)D\beta$  junction and the D $\beta(N)J\beta$ junction may contribute to the specific ability of these cells to recognize the immunodominant epitope. (J. Clin. Invest. 1994. 94:2084-2092.) Key words: autoimmunity • autoimmune disease  $\cdot$  kidney diseases  $\cdot$  gene rearrangement  $\cdot \beta$ chain • T cell antigen receptor

#### Introduction

Recent advances in immunology have facilitated the examination of autoimmune T cell repertoires (1-3), and have provided important insights into the molecular basis of autoreactive T cell recognition of target antigenic immunodominant epitopes (3-7). Analyses of T cell receptor  $(TCR)^1$  variable (V) region genes used by autoreactive T cells in animal models of autoimmunity (3, 8–12) as well as in human autoimmune diseases (3, 13–16), have revealed numerous instances of autoreactive T cell repertoires containing the preferential use of individual V $\beta$  region genes. Such work has resulted in successful, specific anti-V $\beta$  gene immunotherapy (2, 3, 6, 17, 18). Despite these findings, there remains some controversy about the extent of restricted V region gene repertoires in autoimmune diseases (16), especially in systems lacking good information about epitope recognition.

We have been studying the cellular and molecular mechanisms involved in the expression of autoimmune interstitial nephritis (19, 20). In one such model, anti-tubular basement membrane ( $\alpha$ TBM) disease in the SJL mouse, autoimmune interstitial nephritis is induced by immunizing susceptible animals with a rabbit renal tubular antigen preparation in complete Freund's adjuvant (RTA/CFA) (21). Animals develop nonpathogenic anti-tubular basement membrane antibodies in 7-10 d (22), with a subsequent renal mononuclear cell infiltrate at 6-8 wk (21, 22), that progresses to interstitial fibrosis, end stage renal disease, and death (21, 23). The development of  $\alpha$ TBM disease depends upon the emergence of a CD4<sup>+</sup> helper T cell reactive to the target antigen 3M-1 (7), which is expressed on the surface of proximal renal tubular epithelial cells (24). These helper T cells induce the development of 3M-1-reactive CD8<sup>+</sup> effector T cells (25, 26), that mediate cytotoxicity and delayed-type hyper-sensitivity (DTH), and can adoptively transfer disease to naive syngeneic recipients (26, 27).

To further characterize the helper T cell repertoire in  $\alpha$ TBM disease, we have set out to evaluate the peptide antigen specificity, and the selection of T cell receptor V $\beta$  genes in CD4<sup>+</sup> nephritogenic helper T cells.

#### Methods

Antigens. Rabbit renal tubular antigen (RTA) was isolated by previously published methods (28), and emulsified at 8 mg/ml with complete Freund's adjuvant (4 mg/ml of purified protein derivative from Connaught Laboratories Ltd, Toronto, Ontario, Canada; mixed with incomplete Freund's adjuvant purchased from GIBCO BRL, Grand Island, NY) to produce RTA/CFA; ~ 0.2 ml was injected over multiple sites. Soluble renal tubular antigen (SRTA) was made from lyophilized RTA by collagenase digestion (29). Peptides P1 (LLRRRHGDRRSTMSA-EVP), P2 (ASAEEQKEMEDKVTSPEKAEEA), and P3 (STMSAE-VPEAASAE) were derived from the deduced amino acid sequence of the cloned cDNA for 3M-1 (7, 30) and were synthesized by standard peptide synthetic techniques (31), and HPLC purified (University of Pennsylvania peptide synthesis facility, Philadelphia, PA, and Research Genetics, Huntsville, AL).

Immunization. SJL mice (H-2<sup>s</sup>) 4-6 wk of age (Jackson Laboratory,

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<sup>1.</sup> Abbreviations used in this paper:  $\alpha$ TBM, anti-tubular basement membrane; DTH, delayed-type hypersensitivity; RTA, renal tubular antigen; SRTA, soluble RTA; TCR, T cell receptor.

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Bar Harbor, ME) were immunized with RTA/CFA in multiple sites to get  $\alpha$ TBM disease (21).

Isolation and maintenance of T cell lines and clones. T cell lines were established by standard methods (26, 32). Immune cells harvested from draining lymph nodes of SJL mice 10-14 d after immunization were placed in a 24-well culture plate in 2 ml of complete T cell media containing 20  $\mu$ g/ml of appropriate antigen and cultured at 37°C in 5% CO<sub>2</sub> for 4 d. Complete T cell media consists of RPMI 1640 (JRH Biosciences, Lenexa, KS), supplemented with 10% complement-depleted fetal calf serum, 5% NCTC (Whittaker Bioproducts, Walkersville, MD),  $2 \times 10^{-5}$  M 2-mercaptoethanol, glutamine, and antibiotics. After 4 d, and then weekly thereafter, the cells were harvested and placed in fresh media containing antigen, 20% MLA 144 supernatant as a source of IL-2 and other growth factors (33), and  $3-5 \times 10^6$ irradiated (2,500 rads) syngeneic splenic feeder cells (26). T cell lines were cloned twice by limiting dilution in 96-well plates (26), and then expanded and passaged weekly as described above. Some clones were stimulated weekly with SRTA (M33.25.6, M36.9.1, M36.6.1, and M30.2), while others were stimulated weekly with either the P1 (M34.7.3, M34.4.3, M31.1, M31B.3.3) or P2 (M35.2.7, M32.18.2.2) peptide, alone.

Cytokine transcript production by T cell clones. T cell clones were harvested at the end of a 7 d cycle, and were isolated from any remaining dead splenocytes with Lympholyte-M<sup>™</sup> (Cedar Lane Laboratories, Ltd., Hornsby, Ontario, Canada) as per manufacturer recommendations.  $2-4 \times 10^6$  T cells were then cultured in 2 ml of T cell media in 24 well plates, with Concanavalin A (Con A) at 5  $\mu$ g/ml, for 6 h. The supernatants were stored at -20°C for use in HT-2 cell assays, and the cells were washed with T cell media and pelleted. Total RNA was made from the T cells (as well as from snap frozen SJL mouse spleen) by the guanidinium isothiocyanate isolation procedure (34). First strand cDNA was made with mMLV (GIBCO BRL) and oligo-dT (34, 35). 500 ng of the resultant RNA/cDNA was used in a 100  $\lambda$  polymerase chain reaction, containing  $10\lambda$  of  $10 \times$  buffer (Perkin-Elmer Cetus, Exton, PA) 200  $\mu$ M dGTP, dCTP, dATP, dTTP, 0.25  $\lambda$  Taq polymerase (Perkin-Elmer Cetus) and 40 pmol of each oligonucleotide primer. Primer sequences were derived from the cDNAs found in Genbank for IL-2, IL-4, and  $\gamma$ IFN. All oligonucleotide primers were synthesized on an Applied Biosystems 391 DNA Synthesizer (Foster City, CA) and purified with an oligonucleotide purification cartridge as per manufacturer's recommendations. The sequences are as follows: IL-2 sense, 5'CCACTTCAAGCTCTACAGCCGG3'; IL-2 antisense, 5'CCTTAT-GTGTTGTAAGCAGGAGG3'; IL-4 sense, 5'GCTTTCTCTAAC-AGTAGAAACC3'; IL-4 antisense, 5'CCGACAGCGAGACCCAAA-TCTGTC3'; yIFN sense, 5'GGTCAGTGAAGTAAAGGTACA-AGC3'; yIFN antisense, 5'GGAAGAGATTGTCCCAATAAGAA-TA3'. Predicted product sizes are: yIFN, 308 bp; IL-2, 463 bp; and IL-4, 302 bp. cDNA amplification was performed in a Perkin-Elmer Cetus DNA thermal cycler (Perkin-Elmer Cetus) for 35 cycles at 92°C denature for 30 s, 52°C anneal for 60 s, and 72°C elongation for 120 s. 15% of the resultant products were run out on a 2% agarose gel and photographed.

HT-2 proliferation assay. HT-2 cells (kindly provided by Michael Madaio, University of Pennsylvania) were maintained with twice weekly passage in T cell media supplemented with 2 U/ml murine recombinant IL-2 (Boehringer Mannheim Biologicals, Indianapolis, IN). Before use, the cells were pelleted, washed in fresh T cell media, and rested for 3 h at 37°C in 5% CO<sub>2</sub>. 20,000 rested HT-2 cells were incubated with 100 $\lambda$  of T cell clone supernatant and 1  $\mu$ Ci [<sup>3</sup>H]-thymidine (Amersham Corp., Arlington Heights, IL) in a total of 200 $\lambda$ , in 96-well culture plates for 16–48 h. In some experiments, anti–IL-2 blocking antibody (36) (kindly provided by Dr. Glen Gaulton, Department of Pathology, University of Pennsylvania), or murine polyclonal IgM (Bionetics, Charleston, SC), was added at 4  $\mu$ g/ml. Cells were harvested and counted, and the results expressed as the mean of three experiments±SEM.

Cytofluorography. Splenic lymphocytes or T cell clones (at the end of a 7-d passage) were purified through Lympholyte- $M^{TM}$  and washed

with phosphate buffered saline (PBS) with 0.1% BSA.  $1-2 \times 10^6$  cells were suspended in 50 $\lambda$  of PBS/0.1% BSA and incubated with a 1:50 dilution of FITC-conjugated antibody for 30 min on ice. Specific FITCconjugated antibodies for CD4, CD8 (37), TcR $\alpha\beta$  (38), TcR V $\beta7$  (39), TcR V $\beta3$  (40), TcR V $\beta14$  (41), were purchased from Pharmingen (San Diego, CA). PE-conjugated  $\alpha$ CD4 antibody was kindly provided by Bruce Freedman, M. D. (Dept. of Pathology, University of Pennsylvania, Philadelphia, PA). FITC-conjugated goat anti-rabbit antibody was obtained from Boehringer Mannheim Biologicals. The cells were washed three times in PBS/0.1% BSA, and resuspended in 2% paraformaldehyde/PBS. Fluorescence was recorded on a FACScan@ cytofluorograph (24, 42), and analyzed with Lysis II software (Becton Dickinson and Co., San Jose, CA). 10,000 live gated cells were analyzed in each run.

Lymphocyte blast transformation assay. T cell clones were harvested at the end of a 7-d passage cycle, purified through Lympholyte- $M^{TM}$ , and rested in complete T cell media for 4 h at 37°C in 5% CO<sub>2</sub> (24).  $5 \times 10^6$  T cells were mixed with  $3 \times 10^6$  irradiated syngeneic splenocytes and antigen in a total of 1 ml of T cell media, in a 24-well flat bottom plate, and cultured for 72 h at 37°C in 5% CO<sub>2</sub>. 100- $\mu$ l aliquots were removed and placed in flat bottom 96-well plates, mixed with 1  $\mu$ Ci [<sup>3</sup>H]thymidine (Amersham Corp.) and incubated for an additional 24 h. The cells were harvested and counted and the results expressed as the mean of three experiments±SEM.

Effector cell induction assay. The assay was performed as previously described (25, 26, 43). Briefly,  $30-250 \times 10^6$  nonirradiated syngeneic splenocytes were incubated with 20% MLA supernatant, 20  $\mu$ g/ml of antigen, and 20% T cell clone supernatant (taken from the end of a 7-d culture) in a T25 or T75 culture flask (Falcon; Becton Dickinson Labware, Lincoln Park, NJ) for 5 d at 37°C in 5% CO<sub>2</sub>. At the end of the culture period, the induced effector T cells were washed extensively in standard media followed by PBS, and were used for local adoptive transfer of delayed-type hypersensitivity (DTH) or subcapsular transfer.

Local adoptive transfer of DTH. This assay has been previously reported in detail (25, 26). Induced effector cells, as described above, were resuspended in PBS at  $2.0 \times 10^7$  cells/ml, with or without antigen (10 µg/ml), and 25 $\lambda$  of each cell preparation was injected into the hind footpads of recipient naive SJL mice (three per group). Footpad swelling as an index of DTH was measured 48 h later using a spring-loaded engineer's micrometer (Schlesingers for Tools, Brooklyn, NY). The magnitude of swelling was expressed as the mean increment between the antigen-challenged footpad and that injected with PBS in inches  $\times 10^{-3}\pm$ SEM. All measurements were performed by an individual blinded to the experimental protocol.

Subcapsular transfer experiments. Naive syngeneic mice were anesthetized and  $10^8$  induced effector lymphocytes (derived from an effector cell induction as described above) in 75 $\lambda$  of PBS were injected under the kidney capsule with a 30-gauge needle. After 7 d, the kidneys were harvested, fixed in 10% buffered formalin, and stained with hematoxylin and eosin. The stained sections were interpreted by an investigator blinded to the experiment (25, 43).

TCR V $\beta$  gene analysis by cDNA amplification. Total RNA was prepared from Lympholyte-M<sup>TM</sup>-treated T cell clones or snap frozen whole spleen by the guanididium isothiocyanate method (34). Oligo dT-primed first-strand synthesis was performed with mMLV reverse transcriptase (34, 35). PCR was performed in  $100\lambda$  total volume, as described above, using 750 ng of RNA/cDNA template, 40 pmol of 3' antisense primer PH12 (C $\beta$ ; 5'CTCAAACAAGGAGACCTTG3') and 40 pmol of 5' sense V $\beta$  region primer. The 5' V $\beta$  amplification primers for V $\beta$ 1-V $\beta$ 19 were derived by William Smoyer and Carolyn Kelly (University of Pennsylvania, Philadelphia, PA and University of California, San Diego, CA) using a matrix analysis (Macvector sequence analysis software for the Macintosh, IBI, New Haven, CT) and the published murine V $\beta$  gene sequences (44-46) (manuscript in preparation). Each primer is a 20 mer designed to have minimal cross-reactivity to all other  $V\beta$  region genes and encodes a product of unique predicted size. The  $V\beta 20$  primer was designed later from the published sequence (47) using the same criteria, and has the following sequence: 5'GCAGTTACA-

CAGAAGCCAAG3'. Predicted sizes for PCR products using the 3' PH12 primer (C $\beta$ ) and each of the 5' V $\beta$  primers are as follows: V $\beta$ 1,387 bp; V $\beta$ 2,256 bp; V $\beta$ 3250 bp; V $\beta$ 4213 bp; V $\beta$ 6373 bp; V $\beta$ 7165 bp; V $\beta$ 10355 bp; V $\beta$ 14253 bp; V $\beta$ 15394 bp; V $\beta$ 16234 bp; V $\beta$ 17319 bp; V $\beta$ 18327 bp; V $\beta$ 19312 bp; and V $\beta$ 20376 bp. The actual sizes of each product may vary by several base pairs secondary to N region additions and deletions. A master mix of all reagents except the 5' primers was made and divided into 14 tubes each containing one specific V $\beta$  region primer. A fifteenth control tube was run containing two constant region primers 5'-C $\beta$  (5'TACCCTTGTGTGCTTGGC-CA3') and WAS-2 (5'GAACTGCACTTGGCAGCGGA3'), encoding a predicted product of 207 bp. Amplifications were performed in a DNA thermal cycler (Perkin-Elmer Cetus) for 35 cycles at 92°C denature for 30 s, 54°C anneal for 60 s, and 72°C elongation for 120 s. 15% of each reaction was run on a 2% agarose gel and photographed (34, 48).

A cDNA product encoding the T cell receptor  $V\beta$  region from one of the T cell clones was also obtained using anchored PCR (35). Briefly, first strand synthesis was performed using mMLV and a specific 3' TcR  $C\beta$  antisense primer C $\beta$ dan, 5'TCAGGAATTCTTTCTTTTGA3' (49). The cDNA was then G tailed, and amplifications were performed using an internal 3' primer, PH12, and a 5' poly C primer (35).

Southern hybridization. PCR products were transferred to a Zetabind (Cuno, Inc., Meriden, CT) membrane by capillary action using standard techniques, denatured for 10 min with 0.4 M NaOH, and UV cross-linked (Stratalinker; Stratagene, La Jolla, CA). Prehybridization and hybridization were performed as previously described (34) using 1  $\times$  10<sup>6</sup> counts per ml of a <sup>32</sup>P- $\gamma$ ATP-labeled (34) internal C $\beta$  primer, PH15 (5'GAGTCACATTTCTCAGATC3'). The blot was washed twice in 6  $\times$  SSC at 4°C for 20 min, and then washed twice in 3 M tetramethylammonium chloride (Aldrich Chemical Co., Inc, Milwaukee, WI), 50 mM Tris HCl, pH 8.0, 2 mM EDTA, and 0.1% SDS at 55°C for 30 min, then autoradiographed.

Cloning and sequencing of the  $V\beta$  PCR products.  $V\beta$  PCR products were cloned into the TA cloning system (Invitrogen, La Jolla, CA). DNA was prepared for sequencing using the dideoxy chain termination (34, 35). Most  $V\beta$  genes, including the product isolated by anchored PCR, were subcloned from two different PCR reactions, and a single colony from each of the ligations was sequenced, in order to minimize any potential Taq polymearse-induced errors. Sequence analysis was done using the Macvector<sup>TM</sup> sequence analysis program for the Macintosh (IBI). For the isolation of the cDNA for the TcR $\beta$  chain of clone M30.2, we screened an M30.2 cDNA library in Lambda Zap II with a 530-bp TcR C $\beta$  probe (inserted in a Pst1 site of PBR322) kindly provided by Dr. Mark Greene (Department of Pathology, University of Pennsylvania). A full length cDNA clone was obtained, and sequenced as well.

Sequence comparison of T cell receptor deduced amino acid sequences. 52 random murine TcR $\beta$  junctional sequences were obtained from the October 15, 1993 Entrez:Sequences program (NCBI, Bethesda, MD) using the keywords "T-cell receptor" and "T-cell receptor beta." Only sequences encoding functional TcRs in which the V $\beta$ /D $\beta$  and D $\beta$ /J $\beta$  junctions were identified were analyzed. Statistical comparisons were done using  $\chi^2$  analysis.

## Results

Selection of potential antigenic peptides. Two potential antigenic epitopes, P1 and P2 were chosen from the deduced amino acid sequence of the cloned cDNA for the target antigen of  $\alpha$ TBM disease, 3M-1 (7), using a software algorithm integrating a variety of determinants of protein conformation to assist in the prediction of peptide immunogenicity (50) (Fig. 1). P1 is an 18-amino acid peptide, separated by two amino acids from the 22-residue peptide, P2. We later chose a third, overlapping candidate 14 residue peptide, P3 (Fig. 1), that starts at a potential Cathepsin B site (after Arg-Arg), because Cathepsin B

#### LLRRRHGDRR**STMSAEVPEAA**SAEEQKEMEDKVTSPEKAEEA



Figure 1. Amino acid sequences of the potential antigenic epitopes P1, P2, and P3. Abbreviations are the standard single letter amino acid codes. The eight residues in bold type represent the core amino acids of the immunodominant epitope recognized by the nephritogenic T cell clones.

has been shown to be relevant in processing of other antigens presented in the context of MHC class II molecules (51).

General characterization of the T cell clones. To evaluate the relevance of peptides P1, P2, and P3 to the T helper cell response in  $\alpha$ TBM disease, we prepared T cell clones from the draining lymph nodes of several SJL/J animals immunized to get  $\alpha$ TBM disease. The T cell clones were stimulated weekly with either P1, P2, or SRTA. 10 clones were isolated from 8 different animals: 4 were stimulated with SRTA (M33.25.6, M36.9.1., M36.6.1, and M30.2), 4 were stimulated weekly with P1 (M34.7.3, M34.4.3, M31.1, and M31B.3.3), and 2 were stimulated weekly with P2 (M35.2.7, M32.18.2.2).

Previous work by our laboratory has shown that CD4<sup>+</sup> T cells overgrow all other T cell subtypes when isolated by our methods (26). The CD4<sup>+</sup>, CD8<sup>-</sup>, TcR  $\alpha\beta^+$  phenotype of each T cell clone was confirmed with FACS analysis using specific FITC-conjugated antibodies. A representative analysis of one clone is shown is Fig. 2. All of the isolated T cell clones expressed this identical phenotype.

Cytokine expression profile of T cell clones. The reverse transcription polymerase chain reaction (RT-PCR) was initially used to analyze the cytokine profile of each T cell clone. Pilot studies were done using total spleen RNA to determine optimal parameters for the detection of each transcript, and to show that amplification was in the linear range without evidence of plateau (data not shown). The cytokine profile of two representative T cell clones is shown in Fig. 3 A, and reveals appropriate sized products for  $\gamma$ IFN and IL-2, but not IL-4, while all three transcripts are found in splenic RNA. All of the clones exhibited the same cytokine transcript profile. Confirmation of IL-2 production by selected Con A stimulated T cell clones is shown in Fig. 3 B. Supernatants induced proliferation of HT-2 cells that was almost completely blocked by anti-IL-2 antibody, but was unaffected by a polyclonal, control, isotype matched, mouse IgM. All clones tested behaved in a similar fashion (data not shown).

Antigen specificity as determined by local adoptive transfer of DTH. We initially chose to determine the antigen specificity of each clone using a functional assay, the ability to induce an antigen-specific, DTH-producing effector cell population. Each clone was tested with SRTA, P1, P2, and P3. As can be seen in Table I, the four clones stimulated weekly with SRTA (M26.9.1, M33.25.6, M30.2, M36.6.1) were able to induce an antigen-specific DTH response with SRTA, P1 and P3, but not with P2. Similarly, the four clones stimulated weekly with P1 (M34.7.3, M34.4.3, M31B.3.3, M31.1) had the same specificity for SRTA, P1, and P3, but not P2 (Table I). Notably, all eight clones reactive to SRTA were also reactive to P3, implying that P1/P3 is the sole antigenic epitope within the 3M-1 protein in



Figure 2. FACS analysis of clone M36.9.1 using FITC-conjugated antibodies to CD4 (*upper right panel*), CD8 (*lower left panel*), and TCR $\alpha\beta$  (*lower right panel*). A FITC-conjugated goat anti-rabbit antibody served as a negative control (*upper left panel*).

the SJL mouse. The two clones stimulated weekly with P2, however, were only able to induce a DTH response with P2, and not to the intact antigen, SRTA, or to the peptides P1 or P3 (Table I).

Antigen specificity as determined by lymphocyte proliferation assay. Selected T cell clones were additionally tested for their ability to proliferate specifically to the peptide antigens in a standard lymphocyte proliferation assay. As can be seen in Fig. 4, clone M36.9.1 and M33.25.6 proliferate to peptides P1 and P3, but not to P2, consistent with our DTH results.

Subcapsular transfer of induced effector cells. Previous work has shown that the M30.2 clone (SRTA, P1 and P3-reactive) can induce effector cells that cause renal interstitial injury within 5 d of renal subcapular transfer (7, 43). We next wished to test this finding on other P3-reactive clones and to evaluate whether P2-reactive clones could mediate renal inter-



Figure 3. Cytokine profile of selected T cell clones. (A) Representative PCR results for  $\gamma$ IFN (308 bp), IL-2 (463 bp), and IL-4 (304 bp) from total spleen RNA (*left*), clone M36.9.1 RNA (*middle*), and clone M35.2.7 RNA (*right*). Phi is a nucleotide marker. (B) HT-2 cell proliferation assay. ( $\Box$ ) Media alone; ( $\blacksquare$ ) rIL-2 at 0.2 U/ml; ( $\blacksquare$ ) supernatants from Con A-stimulated T cell clone M36.9.1; ( $\blacksquare$ ) supernatants from Con A-stimulated T cell clone M35.2.7. Experiments with rIL-2 and the T cell clone supernatants were performed with no additives (*none*), antiIL-2 antibody at 4  $\mu$ g/ ml (*anti-IL-2*), or a control polyclonal IgM at 4  $\mu$ g/ ml (*IgM*). n = 3 in each experiment; P < 0.05 between anti-IL-2 effect and the IgM control.

Table I. Antigen Specificity of T Cell Clones by DTH

	Weekly stimulus	DTH				
Clone		SRTA	<b>P</b> 1	P2	P3	
M30.2	SRTA	18.7±2.1	19.3±1.2	3.0±2.0	18.3±3.1	
M33.25.6	SRTA	18.3±1.5	18.6±5.5	$3.0 \pm 0.0$	17.0±0.0	
M36.6.1	SRTA	16.3±0.6	19.3±0.6	3.7±1.2	16.7±1.5	
M36.9.1	SRTA	19.3±1.2	17.7±1.2	4.7±0.6	17.0±2.6	
M34.4.3	P1	19.3±1.2	19.0±0.3	3.0±1.0	17.3±2.5	
M34.7.3	P1	19.0±2.6	16.7±0.3	$3.7 \pm 0.6$	21±1.7	
M31.1	P1	16.0±1.7	17.7±1.5	3.0±1.4	17.5±0.7	
M31B.3.3	P1	18.0±1.0	20.0±1.0	3.3±1.2	18.0±1.7	
M32.18.2	P2	3.3±1.2	$2.3 \pm 2.1$	18.0±3.1	3.7±2.5	
M35.2.7	P2	3.3±1.2	3.7±0.9	15.3±1.0	3.0±1.7	

stitial injury. Fig. 5 shows that clone M34.4.3 (P3 reactive) can mediate acute renal injury when P1 peptide is used in the effector cell induction culture (Fig. 5 A). A significant renal infiltrate with tubular degeneration and atrophy occurred within 5 d of subcapsular injection of M34.4.3/P1-induced effector cells (Fig. 5 A). The same clone was unable to cause renal injury when P2 was used in the induction culture (not shown), confirming the antigen specificity of this phenomenon. Similar interstitial renal injury was produced by all eight P3-reactive clones when the P3 peptide was used in the induction culture (data not shown). In contrast, a P2 reactive clone (M35.2.7) was unable to induce nephritogenic effector cells when either P1 (not shown) or P2 (Fig. 5 B) was used in the induction cultures. In these kidneys, we were able to find injected lymphocytes under the capsule of the kidneys, but the histology was indistinguishable from normal controls.

Analysis of TcR V $\beta$  gene usage by nephitogenic helper T cell clones. We next wished to evaluate the TcR V $\beta$  gene used by individual P3-reactive T cell clones. Specificity of the V $\beta$ PCR primers was first demonstrated by Southern analysis of V $\beta$  cDNA amplifications from total spleen RNA using an inter-



Figure 4. Lymphocyte blast transformation assay of selected T cell clones. (**•**) Clone M36.9.1; (**•**) clone M33.25.6. Peptide antigens P1, P2, or P3 were added at 50  $\mu$ g/ml. n = 3 for each group.



Figure 5. Subcapsular transfer of induced effector lymphocytes. (A) Effector cells induced with M34.4.3 (P1- and P3-reactive) and P1 peptide transfer florid interstitial nephritis with a mononuclear cell infiltrate, tubular degeneration and atrophy. (B) Effector cells induced with M35.2.7 (P2-reactive) and P2 peptide are not capable of transferring disease.

nal C $\beta$  region primer, PH15 (Fig. 6). All V $\beta$  PCR products were of predicted size and bound specifically to the internal primer. The genes encoding V $\beta$ 5, V $\beta$ 8, V $\beta$ 9, V $\beta$ 11, V $\beta$ 12, and V $\beta$ 13 are deleted from the genome of SJL mice (52), and were therefore not analyzed.

Fig. 7 shows the results of the TcR V $\beta$  analysis for two representative P3-reactive T cell clones. RT-PCR for TcR V $\beta$ genes performed on RNA from clone M31.1 reveals a product for V $\beta$ 14 and a C $\beta$  control (Fig. 7 A, *left*). Surface expression of V $\beta$ 14 was confirmed by FACS (Fig. 7 A, right). Similarly, clone M36.9.1 expresses V $\beta$ 7 (Fig. 7 B). The V $\beta$  genes used by all 10 T cell clones are shown in Fig. 8. The V $\beta$  genes expressed by six of the clones (those expressing  $V\beta$ 3, 7, and 14) were confirmed by FACS. No antibodies for V $\beta$ 1, 10, 15, and 16 were available. Six V $\beta$  genes were used by the eight different P3-reactive T cell clones, with three clones, M33.25.6, M31.1, and M31B.3., using V $\beta$ 14. The two P2-reactive clones used V $\beta$ 7 and V $\beta$ 1 (Fig. 8). To assess the baseline frequency of V $\beta$ 3, V $\beta$ 7, and V $\beta$ 14 within the peripheral T cell repertoire of the SJL mouse, we performed FACS analysis on freshly isolated splenic lymphocytes derived from unimmunized animals. Double labeling experiments using a PE-conjugated  $\alpha$ CD4 antibody, and FITC-conjugated  $\alpha$ V $\beta$  antibodies revealed that V $\beta$ 3, V $\beta$ 7, and V $\beta$ 14 make up 5%, 5%, and 6.5% of the CD4<sup>+</sup> T cell population, respectively (data not shown).

Analysis of the  $V\beta/D\beta/J\beta$  junctional sequences. The individual V $\beta$  PCR products were then cloned into the PCR 3000 vector (Invitrogen), sequenced, and compared with the known



Figure 6. Southern blot of V $\beta$  PCR products from splenic RNA probed with an internal C $\beta$  oligonucleotide, PH15. Predicted product sizes are listed in Methods.



 $V\beta$  (44–46),  $D\beta$  (53), and  $J\beta$  (54, 55) gene sequences. In all cases, the sequence of the  $V\beta$  PCR product corresponded to the known nucleotide sequence of that particular  $V\beta$  gene, further confirming the specificity of the cDNA amplification products. The nucleotide sequences, D gene, J gene, and deduced amino acid sequences of the 10 T cell clones are shown in Fig. 8. An appropriate open reading frame was present from the V region, through the VDJ junction and into the constant region of the T cell receptor in each case. The eight P3-reactive clones use five

#### VS/DS/JS Junctional Sequences

V814D81.1J81.2M33.25.6TGT GCT TGG AGT CYS ALA TTP SerCGA CAG GGA GCA Arg GLn GLY ALAAAC TCC GAC Asn Ser AspP3M31.1TGT GCC TGG AGT Cys ALA TTP SerLys GLn GLY ALA Lys GLn GLY ALAJ81.2 AAC TCC GACP3M31.1TGT GCC TGG AGT Cys ALA TTP SerLys GLn GLY ALA Lys GLn GLY ALA Asn Ser AspJ81.2 AAC TCC GACP3M31B.3.3TGT GCC TGG AGT Cys ALA TTP SerLys GLn GLY ALA Lys GLn GLY ALAJ81.2 Asn Ser AspP3M30.2TGT GCC AGC AGT Cys ALA SET SerD81.1 Ch GLY ALA Cys ALA SET SerD81.1 Ch GLY ALA Asn GGG GTC Cys ALA SET SerJ81.4 Ch GLY ALAP3M30.2TGT GCT AGC AGC Cys ALA SET SerD81.1 Ch GLY ALA Cys ALA SET SerJ81.1 Ch GLY ALA Asn GLU ATGJ81.2 P3N36.9.1TGT GCT AGC AGC Cys ALA SET SerD81.1 Cys ALA SET SerJ81.1 Cys ALA SET SerJ81.2 Cys ALA SET SerJ81.2 Cys ALA SET SerN36.6.1TGT GCT AGC AGC Cys ALA SET Ser Cys ALA SET Ser Arg GLU GLY LeuJ82.5 GGC AGC GLA GCYJ82.4 Cys ALA SET Ser Arg GLU GLY LeuJ82.4 Cys ALA SET SET PTO Arg Leu GLY GLY Arg GLU ALA THTJ81.1 J82.1 J82.4J83.5 Cys ALA SET SET PTO Arg Leu GLY GLY Arg GLU ALA Arg GLU ALAJ81.1 CAC AGC AGC CYs ALA SET SET Leu TYT Arg GLY Arg Ala SET SET Leu TYT Arg GLY Arg Arg GLU Alp HeeJ81.1 <br< th=""><th>Clone</th><th>95 96 97 98 99</th><th>100 101 102 103</th><th>104 105 106</th><th>Peptide Specificity</th></br<>	Clone	95 96 97 98 99	100 101 102 103	104 105 106	Peptide Specificity
Cys Ala Trp SerArg Gin Gly AlaAsn Ser AspN31.1TGT GCC TGG AGT Cys Ala Trp SerDB1.1 AMA CAA GGA GCA AMA CAA GGA GCA 		V814			
VB14DB1.1JB1.2M31.1TGT GCC TGG AGTAMA CAA GGA GCAAAC TCC GACP3M31.1TGT GCC TGG AGTLys GLn Gly AlaAan Ser AapP3M318.3.3TGT GCC TGG AGTCTA CAG GGG GCAAAC TCC GACP3M318.3.3TGT GCC TGG AGTCTA CAG GGG GCAAAC TCC GACP3M30.2TGT GCC AGC AGTCTA CAG GGG GCAAAC TCC GACP3N30.2V83D81.1J81.4J81.2N36.9.1TGT GCC AGC AGCCQC AGG GGCAAC GAA AGAP3N36.9.1TGT GCT AGC AGCCQC AGG GGCAAC GCA CACP3N36.6.1TGT GCC AGC AGCCQC AGG GGCAAC TCC GACP3N36.6.1TGT GCC AGCAGC GAC GGA CTGGAA GAC ACCP3N36.6.1TGT GCT AGC AGCGGG GGGGGC GGGGAA CAC CCP3N34.9.9TOT GGT GCT AGCGGG GGGGGG GGGGAC ACCP3N34.4.3TGT GCC AGC AGC CCC CGA CTG GGG GGGGGC GAA ACCP3P3N34.4.3TGT GCC AGC AGC CCC CGA CTG GGG GGG AGGArg Gln ThrP3N32.18.2V81D81.1J82.5Arg Gln AgcArg Gln AgcN32.18.2V81D81.1J82.5AgC GAA GACP2N35.2.7GCT AGC AGC CTC TAC AGG GGG AGG AGC GATGAA GGC TTCP2	M33.25.6				P3
M31.1 TOT GCC TGG AOT Cys Ala Trp Ser AMA CAA GQA GQA Lys Gln Gly Ala AAC TCC GAC P3   M31.1 TOT GCC TGG AOT Cys Ala Trp Ser D81.1 J81.2 J81.2 P3   M31B.3.3 TGT GCC TGG AOT Cys Ala Trp Ser D81.1 J81.2 AAC TCC GAC P3   M30.2 TGT GCC AGC AOT Cys Ala Ser Ser D81.1 J81.4 J81.4 P3   M30.2 TGT GCC AGC AOT Cys Ala Ser Ser D81.1 J81.4 P3   M36.9.1 TGT GCC AGC AOC Cys Ala Ser Ser D81.1 J81.2 P3   M36.6.1 V87 D81.1 J81.2 P3   M36.6.1 TGT GCC AGC AOC Cys Ala Ser Ser D81.1 J81.2 P3   M36.6.1 TGT GCC AGC AGC GAG GAC Cys Ala Ser Ser D81.1 J82.5   M34.9.9 TOT GGT GCT ACG Cys Gly Ala Thr D82.1 J82.4 P3   M34.4.3 TGT GCC AGC AGC CCC CGA CAG GGG GGG Cys Gly Ala Thr D82.1 J82.3 CCC AA ACC   M34.4.3 TGT GCC AGC AGC CCC CGA CAG GGG GGG CGC AA ACC P3   M32.18.2 V816 D81.1 J82.5 ACC AA ACC   M32.18.2 V81 D81.1 J82.5 P2   M32.18.2 V81 D81.1 J82.5 P2		Cys Ala Trp Ser	Arg Gln Gly Ala	Asn Ser Asp	
M31.1 TOT GCC TGG AOT Cys Ala Trp Ser AMA CAA GGA GCA Lys Gin Gly Ala AAC TCC GAC P3   M318.3.3 TGT GCC TGG AOT Cys Ala Trp Ser D81.1 J81.2 AAC TCC GAC P3   M318.3.3 TGT GCC TGG AOT Cys Ala Trp Ser Leu Gin Gly Ala Asn Ser Asp P3   M30.2 TGT GCC AGC AOT Cys Ala Ser Ser D81.1 J81.4 J81.4 P3   M30.2 TGT GCC AGC AOT Cys Ala Ser Ser D81.1 J81.4 P3   M36.9.1 TGT GCC AGC AOC Cys Ala Ser Ser D81.1 J81.2 P3   M36.6.1 TGT GCC AGC AOC Cys Ala Ser Ser D81.1 J81.2 P3   M36.6.1 TGT GCC AGC AOC Cys Ala Ser Ser D81.1 J82.5 P3   M36.6.1 TGT GCC AGC AGG GAG GGG Cys Alg Ser Arg Arg Glu Gly Leu Glu Gly Leu Glu Gly Clu J82.4 P3   M34.9.9 TOT GGT GCT ACG CC Cys Gly Ala Thr D82.1 J82.3 J82.3   M34.4.3 TGT GCC AGC AGC CCC CGA CMG GGG GGG Cys Gly Ala Thr D82.1 J82.3   M34.4.3 TGT GCC AGC AGC CCC CGA CMG GGG GGG Cys Ala Ser Ser Pro Arg Leu Gly Gly J82.5 Arg Gln Thr   M32.18.2 V81 D81.1 J82.5 Arg Gln Agc Arg Gln Agc Arg Gln Agc P3   M32.18.2 V81 D81.1 J82.1 J82.5		V614	D81.1	J81.2	
Cys Ala Trp SerLys Gin Gly AlaAsn Ser AspN31B.3.3TGT GCC TGG AGT Cys Ala Trp SerD81.1 Cys Ala Trp SerJ81.2 Cys Ala Ser SerJ81.2 AAC TCC GACP3N30.2V83 Cys Ala Ser SerD81.1 CAG GG GCC Cys Ala Ser SerJ81.1 CAG GGG GCC ASG GG GCC ASG GGG GCC ASG GGG GCC ASG GGG GCC ASG GGG GCC ASG GGG GCC ASG GGG GCC Cys Ala Ser SerJ81.1 J81.2 J81.2 ASG GG GCC ASG GGC GCC Cys Ala Ser Ser Arg Glu Gly Lau Glu Glu Ser ThrJ82.3 J82.4 CAC Cys Ala Ser Ser Pro Arg GGG GGG Cys Gly Ala ThrJ82.1 GCC AGC AGC CCC GGA CTG GGG GGG CCC GAA AGG Arg Glu Gly Glu Glu Glu ThrJ82.3 FTGN34.4.3TGT GCC AGC AGC CCC CGA CTG GGG GGG Cys Ala Ser Ser Pro Arg Leu Gly Gly Ala Ser Ser Pro Arg Glu Agg GAA GAC Cys Ala Ser Ser Pro Arg Glu Agg GAA GAC Arg Glu Agg CAA AGG Arg Glu Ang FTJ82.3 FTG GCT GCA AGC CCC CGA CTG GGG GGG Arg Glu Ang FTJ82.3 FTN34.4.3TGT GCC AGC AGC CCC CGA CTG GGG GGG Cys Ala Ser Ser Pro Arg Leu Gly Gly Ala Ser Ser Pro Arg Leu Gly Gly Ala Ser Ser Leu Tyr Arg Gly Arg Arg Glu AngJ82.5 Agg GAA GAC GC TTC FTP2N35.2.7V87 GCT AGC AGC CTC GAC GGG GGG GGG GGG GGG AGA Ala Ser Ser Leu Tyr Arg Gly Agg AGA GGC TTC Cyc AGC GGG GGA CA CAA AGC TTC Cyc AGC GGG GGA CA CAA GC CTC TTCP2	M31.1				P3
W31B.3.3 TOT GCC TGG AGT Cys Ala Trp Ser CTA CAG GGG GCA Leu Gln Gly Ala AAC TCC GAC Asn Ser Asp P3   N30.2 V83 D81.1 J81.4 J81.4 J81.4 J81.4   N30.2 TGT GCC AGC AGT Cys Ala Ser Ser Gln Gly Val Asn Glu Arg P3   N30.2 V87 D81.1 J81.4 J81.4 P3   N36.9.1 TGT GCC AGC AGT Cys Ala Ser Ser D81.1 J81.2 J61.4 P3   N36.6.1 TGT GCC AGC AGT Cys Ala Ser Ser Arg Gly Asn Ser Asp P3   N36.6.1 TGT GCC AGC AGG GAG GGT Cys Ala Ser J82.1 J82.5   N34.9.9 V815 D82.1 J82.4 J82.4   N34.4.3 TGT GCT AGC AGC CCC CGA CAG GGG GGG Cys Ala Ser Ser Pro Arg Leu Gly Gly J82.3 J82.3   N34.4.3 TGT GCC AGC AGC CCC CGA CAG GGG GGG Cys Ala Ser Ser Pro Arg Leu Gly Gly J82.5 J82.3   N34.4.3 TGT GCC AGC AGC CCC CGA CAG GGG GGG AGG AGG		Cys Ala Trp Ser			
M31B.3.3 TOT GCC TGG AGT Cys Ala Trp Ser CTA CMG GGG GCA Leu Gln Gly Ala AAC TCC GAC Asn Ser Asp P3   N30.2 VB3 TGT GCC AGC AGT Cys Ala Ser Ser DB1.1 Gln Gly Val JB1.4 Asn Glu Arg JB1.4 Asn Glu Arg P3   N30.2 VB7 Cys Ala Ser Ser DB1.1 Gln Gly Val JB1.4 Asn Glu Arg P3   N36.9.1 TOT GCC AGC AGT Cys Ala Ser Ser DB1.1 Asn Glu Arg JB1.2 Asn Glu Arg P3   N36.6.1 TGT GCC AGC AGC Cys Ala Ser Ser DB1.1 Arg Gly Glu Glu Asp Thr JB2.5 Glu Asp Thr P3   N36.6.1 TGT GCT AGC AGC Cys Ala Ser Arg Glu Gly Crg Glu Glu App Thr JB2.4 SE2.4 P3   N34.9.9 TOT GGT GCT AGC Cys Gly Ala Thr DB2.1 Gly Gly Gly Bis Thr JB2.3 Coc CAA ACC P3   N34.4.3 TGT GCC AGC CAGC CCC CGA CRG GGG GGG Cys Ala Ser Ser Pro Arg Leu Gly Gly Ala Ser Ser Pro Arg Leu Gly Gly JB2.5 Arg Gln Thr JB2.3 Arg Gln Thr   M32.18.2 VB1 GC AGC AGC CCC CGA CRG GGG AGG Ala Ser Ser Leu Tyr Arg Gly Arg Ala Ser Ser Leu Tyr Arg Gly Arg Arg Gln App P2   M35.2.7 GCT AGC AGT CAC AGC GGG GGA CA AGT GGA GGA CT TC D81.1 AGC GGA GGA CT D81.1 AGA GTC TTC P2					
Cys Ala Trp Ser Leu Gin Gly Ala Ass Ser Asp   V\$3 D\$1.1 J\$1.4   N30.2 TGT GCC AGC AGT CAG GGG GTC AAC GAA AGA P3   Cys Ala Ser Ser Gin Gly Val Ass Glu Arg P3   N36.9.1 TGT GCT AGC AGC CQC AGG GGC AAC CC GAC P3   N36.9.1 TGT GCT AGC AGC CQC AGG GGC Ass Ser Asp P3   N36.6.1 TGT GCT AGC AGC CQC AGG GGG GAA GGA CCC P3   N36.6.1 TGT GCT GCT ACG AGG GA GGG GAA GGA CCC P3   N36.6.1 TGT GCT GCT ACG GGG GGG GAA GCC P3   N36.6.1 TGT GCT GCT ACG GGG GGG GAA GAC CC P3   N34.9.9 TGT GCT GCT ACG GGG GGG GAA GAC CC P3   N34.9.9 TGT GCT ACG AGC CCC CGA CTG GGG GGG GCC CAA ACG P3   N34.4.3 TOT GCC AGC AGC CCC CGA CTG GGG GGG CGC CAA ACG P3   N34.4.3 TOT GCC AGC AGC CCC CGA CTG CGG GGG GCC CAA ACG P3   N34.4.3 TOT GCC AGC AGC CCC CGA CTG CGG GGG AGG Arg Gln Thr P3   N32.18.2 V81 D\$1.1 J\$2.5 AGG CAA GAC   N32.18.2 GCC AG	M318 3 3				53
VB3 DB1.1 JB1.4 JB1.4   N30.2 TGT GCC AGC AGT CAG GGG GTC AAC GAA AGA P3   VB7 DB1.1 JB1.2 Ass Glu Arg P3   N36.9.1 TGT GCT AGC AGC CQC AGG GGC AAC CGAC AGA P3   N36.9.1 TGT GCT AGC AGC CQC AGG GGC AAC CGAC GAC P3   N36.6.1 TGT GCC AGC AGG GAG GGG GAA GAC ACC P3   N36.6.1 TGT GCT AGC AGG GAG GGG GAA GAC ACC P3   N34.9.9 VB15 DB2.1 JB2.4 JB2.4   VB15 DB2.1 JB2.4 JB2.4 JB2.3   N34.4.3 TGT GCT AGC AGC CCC CGA CTG GGG GGG CCC CAA ACC P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG CCC CAA ACG P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG CCC CAA ACG P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG CCC CAA ACG P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG Arg Gln Abc P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG AGG Arg Gln Abc P2   N32.18.2 VB1 DB1.1 JB2.5 AcG CAA GAC   N32.18.2 GC1 AGC AGC CCC CCC AGC AGC GCC CCG AGG AGG	M318.3.3				PS
M30.2 TOT GCC AGC AGT Cys Ala Ser Ser CAG GGG GTC Gln Gly Val AAC GAA AGA Asn Glu Arg P3   N36.9.1 V87 TGT GCT AGC AGC Cys Ala Ser Ser D81.1 Arg Gly J81.2 Asn Ser Asp J81.2 Asn Ser Asp P3   N36.9.1 TGT GCT AGC AGC Cys Ala Ser Ser Arg Arg Gly Arg Gly Asn Ser Asp P3   N36.6.1 TGT GCT AGC Cys Ala Ser Arg Glu Gly Teu Glu Asp Thr J82.5 GGA GAC ACC P3   N34.9.9 TGT GGT GCT ACG Cys Gly Ala Thr GGG GGG Cys GGA GGC AGC GGG GGG Cys Gly Ala Thr J82.1 Glu Asp Thr J82.3 FGT GCC AGC AGC CCC CGA CTG GGG GGG Cys Ala Ser Ser Pro Arg Leu Gly Gly Arg Gln Thr J82.3 Arg Gln Ang P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG Cys Ala Ser Ser Pro Arg Leu Gly Gly Ala Ser Ser Leu Tyr Arg Gly Arg Arg Gln App J82.5 Arg Gln App P2   N32.18.2 V81 Ala Ser Ser Leu Tyr Arg Gly Arg Ala Ser Ser Leu Tyr Arg Gly Arg Ala Ser GGG GGG AGC Ala GGC GGG AGC ARA GGC TTC J81.1 J81.1 J81.1 J81.1 J81.1   N35.2.7 GCT AGC AGC CCC CAC AGC GGG GGG AGA AGA GGC TTC D81.1 CAC AGC GGG GGG AGA AGA GGC TTC P2		Cys Ala IIP Sel	Ded Gin Giy Ala	Nen Ger Veb	
Cys Ala Ser Ser Gin Gly Val Asn Glu Arg   V87 D81.1 J81.2   N36.9.1 TGT GCT AGC AGC CQC AGG GGC AAC TCC GAC P3   N36.9.1 TGT GCT AGC AGC CQC AGG GGC AAC TCC GAC P3   N36.6.1 TGT GCC AGC AGG GAG GGA CTG GAA GAC ACC P3   N36.6.1 TGT GCC AGC AGG GAG GGA CTG GAA GAC ACC P3   N34.9.9 TGT GGT GCT ACG GGG GGG GGG GGG CAC ACC P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG GGC GAG AGC CC CAA ACG P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG GC CAA ACG P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG GC CAA ACG P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG GC CAA ACG P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG GC CAA ACG P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG Arg Gln Thr P3   N34.4.3 TGT GCC AGC AGC CCC TAC AGG GGG AGG Arg Gln Abc P3   N32.18.2 V81 D81.1 J82.5 Arg Gln Abc   N32.18.2 GCC AGC AGC CCC CT TAC AGG GGG AGG Arg Gln Abp   N35.2.7 GCT AGC AGC			D\$1.1	J81.4	
V87 D81.1 J81.2   H36.9.1 TGT GCT AGC AGC GQC AGG GGC AAC TCC GAC   Cys Ala Ser Ser Arg Arg Gly Ass Ser Asp   H36.6.1 TGT GCC AGC AGG GG GGC GAA GAC ACC   H36.6.1 TGT GCC AGC AGG GG GGG GAA GAC ACC   H36.6.1 TGT GCC AGC AGG GG GGG GAA GAC ACC   H34.9.9 TGT GCT GCT ACG GGG GGG GAC ACC   V815 D82.1 J82.4   H34.4.3 TGT GCC AGC AGC CCC GA CTG GGG GGG GCC CAA ACG   V816 D82.1 J82.3   V816 D82.1 J82.3   H34.4.3 TGT GCC AGC AGC CCC GA CTG GGG GGG CGC CAA ACG   V916 D81.1 J82.5   H32.18.2 V81 D81.1 J82.5   H32.18.2 V81 D81.1 J82.5   H32.18.2 GCC AGC AGC CTC TAC AGG GGG AGG Arg Gln Asp   H32.18.2 V81 D81.1 J82.5   H32.18.2 GCT AGC AGC CTC TAC AGG GGG AGA Arg Gln Asp   H32.18.2 GCT AGC AGC CTC TAC AGG GGG AGA Arg Gln Asp   H32.18.2 GCT AGC AGC CTC TAC AGG GGG AGA Arg Gln Asp   H35.2.7 GCT AGC AGT CAC GCG GGG GGG AGA GAA GTC TTC	M30.2				P3
N36.9.1 TOT GCT AGC AGC OGC AGC GGC AAC TCC GAC P3   V810 D81.1 J82.5 GAA GAC ACC P3   N36.6.1 TGT GCC AGC AGG GAG GGC GAA GAC ACC P3   N36.6.1 TGT GCC AGC AGG GAG GGC GAA GAC ACC P3   N34.9.9 TGT GGT GCT AGC AGC GGG GGG GAC ACC P3   N34.4.3 TOT GCC AGC AGC CCC CGA CTG GGG GGG GCC CAA ACC P3   N34.4.3 TOT GCC AGC AGC CCC CGA CTG GGG GGG CGC CAA ACG P3   N34.4.3 TOT GCC AGC AGC CCC CGA CTG GGG GGG CGC CAA ACG P3   N34.4.3 TOT GCC AGC AGC CCC CGA CTG GGG GGG CGC CAA ACG P3   N34.4.3 TOT GCC AGC AGC CCC CGA CTG GGG GGG CGC CAA ACG P3   N34.4.3 TOT GCC AGC AGC CCC TAC AGG GGG AGG Arg Gln Thr P3   N34.4.3 TOT GCC AGC AGC CCC TAC AGG GGG AGG Arg Gln Agp P2   N32.18.2 V81 D81.1 J82.5 AgG GAA GAC P2   N32.18.2 V87 D81.1 J81.1 J81.1 P2   N35.2.7 GCT AGC AGC CAGC CCC CAA GG GG GAA GA GAA GA		C <b>ys Ala Ser Ser</b>	Gln Gly Val	Asn Glu Arg	
N36.9.1 TOT GCT AGC AGC OGC AGC GGC AAC TCC GAC P3   V810 D81.1 J82.5 GAA GAC ACC P3   N36.6.1 TGT GCC AGC AGG GAG GGC GAA GAC ACC P3   N36.6.1 TGT GCC AGC AGG GAG GGC GAA GAC ACC P3   N34.9.9 TGT GGT GCT AGC AGC GGG GGG GAC ACC P3   N34.4.3 TOT GCC AGC AGC CCC CGA CTG GGG GGG GCC CAA ACC P3   N34.4.3 TOT GCC AGC AGC CCC CGA CTG GGG GGG CGC CAA ACG P3   N34.4.3 TOT GCC AGC AGC CCC CGA CTG GGG GGG CGC CAA ACG P3   N34.4.3 TOT GCC AGC AGC CCC CGA CTG GGG GGG CGC CAA ACG P3   N34.4.3 TOT GCC AGC AGC CCC CGA CTG GGG GGG CGC CAA ACG P3   N34.4.3 TOT GCC AGC AGC CCC TAC AGG GGG AGG Arg Gln Thr P3   N34.4.3 TOT GCC AGC AGC CCC TAC AGG GGG AGG Arg Gln Agp P2   N32.18.2 V81 D81.1 J82.5 AgG GAA GAC P2   N32.18.2 V87 D81.1 J81.1 J81.1 P2   N35.2.7 GCT AGC AGC CAGC CCC CAA GG GG GAA GA GAA GA		V87	D81.1	381.2	
Cys Ala Ser Ser Arg Arg Gly Asn Ser Asp   V810 D81.1 J82.5   H36.6.1 TGT GCC AGC AGG GAA GGA CTG GAA GAC ACC P3   K34.9.9 TGT GGT GCT AGG GGG GGG GGG GGG GAC ACC P3   K34.4.3 TGT GCC AGC AGC D82.1 J82.4 J82.4   K34.4.3 TGT GCC AGC AGC CCC CGA CGG GGG GGG GGG GCA CACC P3   K34.4.3 TGT GCC AGC AGC CCC CGA CGG GGG GGG CGC CAA ACG P3   K34.4.3 TGT GCC AGC AGC CCC CGA CGG GGG GGG CGC CAA ACG P3   K34.4.3 TGT GCC AGC AGC CCC CGA CGG GGG GGG CGC CAA ACG P3   K34.4.3 TGT GCC AGC AGC CCC CGA CGG GGG GGG CGC CAA ACG P3   K34.4.3 TGT GCC AGC AGC CCC CGA CGG GGG GGG CGC CAA ACG P3   K34.4.3 TGT GCC AGC AGC CCC CGA CGA GGG GGG GGG CGC CAA ACG P3   K32.18.2 VB1 D81.1 J82.5 Acg CAA GAC   K32.18.2 GCC AGC AGC CCC CTC TAC AGG GGG AGA Acg CAA GAC P2   K35.2.7 GCT AGC AGT CAC AGG GGG GGA CA GAA GGC TTC P2	M36.9.1				P3
M36.6.1 TGT GCC AGC Cys Ala Ser AGG GAG GGA CTG Arg Glu Gly Leu GAA GAC ACC Glu Asp Thr P3   N34.9.9 TGT GGT GCT Cys Gly Ala Thr DB2.1 JB2.4 JB2.4   N34.4.3 TGT GGT GCT AGG Cys Gly Ala Thr Gly Gly Bis Thr JB2.3   VB16 DB2.1 JB2.3 CGC GAA ACC P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG CGC CAA ACG P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG CGC CAA ACG P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG CGC CAA ACG P3   N34.4.3 TGT GCC AGC AGC CCC TG ACG GGG AGG Arg Gln Thr P3   N32.18.2 VB1 DB1.1 JB2.5 AGG CAA GAC   N32.18.2 VB7 DB1.1 JB1.1 JB1.1   N35.2.7 GCT AGC AGT CAC AGG GGG ACA GAA GTC TTC P2		Cys Ala Ser Ser	Arg Arg Gly	Asn Ser Asp	
M36.6.1 TGT GCC AGC Cys Ala Ser AGG GAG GGA CTG Arg Glu Gly Leu GAA GAC ACC Glu Asp Thr P3   N34.9.9 TGT GGT GCT Cys Gly Ala Thr DS2.1 JS2.4 JS2.4   N34.4.3 TGT GGT GCT AGC Cys Gly Ala Thr Gly Gly Gly Bis Thr JS2.3   N34.4.3 TGT GCC AGC CCC CGA CTG GGG GGG Cys Ala Ser Ser Pro Arg Leu Gly Gly JS2.3   N32.18.2 VS1 DS1.1 JS2.5   N32.18.2 VS7 DS1.1 JS2.5   N35.2.7 GCT AGC AGT CAC AGT GAA GGA CAC					
V\$15 D\$2.1 J\$2.4   N34.9.9 TGT GGT GGT ACG GGG GGG CAC ACC   Y\$15 D\$2.1 J\$2.4   Y\$16 D\$2.1 J\$2.4   Y\$17 GGT ACG GGG GGG   Y\$18 TGT GGT GCT ACG D\$2.1   Y\$19 His Thr His Thr   Y\$16 D\$2.1 J\$2.3   Y\$16 D\$2.1 J\$2.3   Y\$16 D\$2.1 J\$2.3   Y\$17 GCT AGC AGC CCC CGA CTG GGG GGG CGC CAA ACG   Y\$18 Ser Ser Pro Arg Leu Gly Gly Arg Gln Thr	N36 6 1				
VB15 DB2.1 JB2.4   H34.9.9 TOT GOT GOT GCT ACG GOG GOG CAC ACC   YB16 DB2.1 JB2.3   H34.4.3 TOT GOT AGC AGC AGC CCC CGA CRG GOG GOG GC CCC CGA CRG GOG GOG GC CAA ACC P3   H34.4.3 TOT GCC AGC AGC CCC CGA CRG GOG GGG GG CAA ACC P3   H34.4.3 TOT GCC AGC AGC CCC CGA CRG GOG GGG GG CAA ACC P3   H32.18.2 VB1 DB1.1 JB2.5   H32.18.2 GCC AGC AGC CCC CTC TAC AGG GGG AGG AGG ACG ACA GAC P2   H35.2.7 GCT AGC AGT CAC GCA CAC AGA GGG GGG AGA GAA GTC TTC P2	H30.0.1				P3
M34.9.9 TOT GGT GCT ACG Cys Gly Ala Thr GGG GGG Gly Gly GL ACC Bis Thr P3   M34.4.3 TOT GCC AGC AGC Cys Ala Ser Ser Pro Arg Leu Gly Gly JB2.3 CGC AAA ACG Arg Gln Thr JB2.3 CGC AAA ACG Arg Gln Thr P3   M32.18.2 VB1 ACC AGC AGC CCC CTC TAC AGG GGG AGG Ala Ser Ser Leu Tyr Arg Gly Arg JB2.5 AGG CAA GAC Arg Gln App P2   M32.18.2 VB1 Ala Ser Ser Leu Tyr Arg Gly Arg Ala Ser Ser Leu Tyr Arg Gly Arg JB1.1 JB1.1 JB1.1 JB1.1 JB1.1 JB1.1   M35.2.7 GCT AGC AGT CAC AGG GGG GGG ACA GAA GTC TTC P2		- <i>i</i> ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	my ore ory bee	ord map the	
VB16 DB2.1 JB2.3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG CGC CAA ACG P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG GGG CGC CAA ACG P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG GGG CGC CAA ACG P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG GGG CGC CAA ACG P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG GGG CGG CAA ACG P3   N34.4.3 TGT GCC AGC AGC CCC CGA CTG GGG AGG AGG CGG CAA GAC P3   N32.18.2 VB1 D81.1   N32.18.2 GCC AGC AGC CTC TAC AGG GGG AGG AGG AGG CAA GAC P2   N132.18.2 GCC AGC AGC CTC TAC AGG GGG AGG AGG AGG CAA GAC P2   N132.18.2 GCC AGC AGC CTC TAC AGG GGG AGG AGG AGG AGG AGG CAA GAC P2   N132.18.2 GCC AGC AGC CTC TAC AGG GGG AGG AGG AGG AGG AGG AGG AGG A					
VB16 DB2.1 JB2.3   M34.4.3 TOT GCC AGC AGC CCC CGA CTG GGG GGG GG CCC CAA ACG P3   M34.4.3 TOT GCC AGC AGC CCC CGA CTG GGG GGG GG CCC CAA ACG P3   M32.18.2 VB1 DB1.1 JB2.5   M32.18.2 GCC AGC AGC CTC TAC AGG GGG AGG AGG AGG AGG AGG AGG AGG A	M34.9.9				P3
W34.4.3 TOT GCC AGC AGC CCC CGA CTG GGG GGG GGG CCC CAA ACG P3   M34.4.3 TOT GCC AGC AGC CCC CGA CTG GGG GGG GGG CCC CAA ACG P3   M32.18.2 VB1 DB1.1 JB2.5   M32.18.2 GCC AGC AGC CTC TAC AGG GGG AGG AGG AGG CAA GAC P2   M35.2.7 GCT AGC AGC CTC CAC AGG GGG AGG AGG AGG AGA CTC TTC P2		Cys Gly Ala Thr	Gly Gly	His Thr	
V\$1 D\$1.1 J\$2.18.2   W\$1 D\$1.1 J\$2.5   M32.18.2 GCC AGC AGC CTC TAC AGG GGG AGG AGG CAA GAC   P2 Ala Ser Ser Leu Tyr Arg Gly Arg Arg Gln Asp   V\$7 D\$1.1 J\$1.1   M35.2.7 GCT AGC AGT CAC GGG GGG AGG GGA AGT CTC TPC		V816	D82.1	J82.3	
vši pši.i jš2.5   M32.18.2 GCC AGC AGC CTC TAC AGG GGG AGG AGG CAA GAC P2   Ala Ser Ser Leu Tyr Arg Gly Arg Gln Asp   VS7 DŠ1.1 JŠ1.1   N35.2.7 GCT AGC AGG CAC AGG GGG ACG GAA GTC TTC	M34.4.3	TGT GCC AGC AG <u>C CC</u>	<u>C C</u> GA CTG GGG GGG	CGC CAA ACG	P3
H32.18.2 GCC AGC AGC CCC TAC AGG GGG AGG AGG AGG CAA GAC P2 Ala Ser Ser Leu Tyr Arg Gly Arg Arg ClA App V87 D81.1 J81.1 H35.2.7 GCT AGC AGT CAC AGG GGG ACA GAA GTC TTC P2		Cys Ala Ser Ser Pr	o Arg Leu Gly Gly	Arg Gln Thr	
H32.18.2 GCC AGC AGC C <u>TC T</u> AC AGG GGG AGG AGG CAA GAC P2 Ala Ser Ser Leu Tyr Arg Gly Arg Arg ClA AgA CPC VB7 DB1.1 JB1.1 H35.2.7 GCT AGC AGT <u>C</u> AC AGG GGG ACA GAA GTC TTC P2					
Ala Ser Ser Leu Tyr Arg Gly Arg Arg Gln Asp V87 D81.1 J81.1 M35.2.7 GCT AGC AGT <u>CAC AGG GGG ACA</u> GAA GTC TTC P2					
VB7 DB1.1 JB1.1 H35.2.7 GCT AGC AGT CAC AGG GGG ACA GAA GTC TTC P2	M32.18.2				P2
M35.2.7 GCT AGC AGT CAC AGG GGG ACA GAA GTC TTC P2		Ala Ser Ser Le	u Tyr Arg Gly Arg	Arg Gln Asp	
M35.2.7 GCT AGC AGT CAC AGG GGG ACA GAA GTC TTC P2		<b>VB</b> 7	D\$1.1	J61.1	
	M35.2.7				P2
		Ala Ser Ser	His Arg Gly Thr	Glu Val Phe	

Figure 8. Aligned nucleotide and deduced amino acid sequences of the ten T cell clones across the  $V\beta/D\beta/J\beta$  junction. The  $V\beta$ ,  $D\beta$ , and  $J\beta$  genes used by each clone are in bold type above each sequence. Underlined nucleotides represent positions of N region additions/deletions. Amino acid residue numbers 95–106 are listed across the top of the figure.

Figure 7. T cell receptor  $\nabla\beta$  expression by individual T cell clones. (A) RT-PCR for  $\nabla\beta$  genes using RNA from clone M31.1 revealing a PCR product for  $\nabla\beta14$  (253 bp) and for  $C\beta$  (207 bp), with confirmatory surface expression by FACS using a specific  $\nabla\beta14$  antibody. (B) RT-PCR for  $\nabla\beta$  genes using RNA from clone M36.9.1 revealing a PCR product for  $\nabla\beta7$  (165 bp) and for  $C\beta$  (207 bp), with confirmatory surface expression by FACS using a specific  $\nabla\beta7$  antibody. An FITC conjugated goat anti-rabbit antibody served as a negative control for the FACS studies.

different J $\beta$  genes, with a preponderance of J $\beta$  1.2 (used by M33.25.6, M31.1, M31B.3.3, and M36.9.1). Six of the eight P3-reactive clones use D $\beta$ 1.1.

The deduced amino acid sequences of the TcR $\beta$  chains from the eight P3-reactive T cell clones are remarkably similar across the  $V\beta/D\beta/J\beta$  junction, with clones M33.25.6, M31.1, M31B.3, and M30.2 showing identical or conservatively substituted amino acids in 5 or 6 positions. All eight P3-reactive clones have a small nonpolar amino acid (Ala, Val, Leu, or Gly) at the D $\beta$ /J $\beta$  junction (residue 102 or 103), and five of the eight clones have a cationic, basic residue (Arg or Lys) at the  $V\beta/D\beta$  junction (residue 100). Residues in both of these positions are encoded primarily by N region deletions or additions (underlined in Fig. 8), and are thus potentially highly variable. Five of the eight P3-reactive clones additionally have an Asn residue 104. In the lower portion of Fig. 8 are the deduced amino acid sequences of the TcR  $\beta$  chains from the two, P2-reactive clones. The sequence of the  $V\beta/D\beta/J\beta$  junctions shows some homology to the P3-reactive clones, but position 103 contains a charged residue (Arg in M32.18.2 and Glu in M35.2.7) as opposed to a small nonpolar residue. Additionally, position 100 is Tyr or His in the P2 reactive clones, and not Arg as found in most of the P3-reactive clones.

To confirm the relevance of the sequence findings, we analyzed a random sample of 52  $V\beta/D\beta/J\beta$  sequences for the presence of Ala, Val, Leu or Gly residue at the  $D\beta/J\beta$  junction, or an Arg or Lys at the  $V\beta/D\beta$  junction. 14 of 52 sequences contained an Ala, Val, Leu, or Gly at the  $D\beta/J\beta$  junction, compared to 8/8 of our P3 reactive clones (P < 0.001), while 14/52 of the random sequences contained an Arg or Lys at the  $V\beta/D\beta$  junction, compared to 5/8 of our P3-reactive clones (P < 0.05). Additionally, 6/8 of the P3-reactive  $V\beta/D\beta/J\beta$  sequences are cationic (with the other two having an overall neutral charge), compared with the 5/52 incidence found in the randomly chosen T cell sequences (P < 0.001).

#### Discussion

We have successfully isolated a panel of CD4<sup>+</sup>, TcR  $\alpha\beta$ -expressing T cell clones from mice immunized to get  $\alpha$ TBM disease. The cytokine profiles and the functional characteristics (i.e., the ability to provide T cell help for the induction of DTH-

producing effector cells) of these clones are consistent with a Th1 phenotype (56). We have not tested these clones for their ability to induce a humoral immune response, however, and cannot fully exclude B cell help as a potential function of these cells. Additionally, we have not yet attempted to preferentially isolate Th2-like cells from these animals.

Antigen specificity was demonstrated functionally in all clones by an established biological assay (25, 43), and confirmed in selected clones through the use of a proliferation assay (24, 26). All clones stimulated weekly with SRTA or P1 specifically reacted to the P3 peptide, and not to P2, suggesting that these T cell clones recognize a core sequence of eight amino acids (STMSAEVP, bold letters in Fig. 1). Interestingly, this core sequence, as well as the entire P3 epitope, is anionic, and contains no positively charged residues (Fig. 1). Future work involving the elution of peptide from MHC II molecules (57), and a systematic evaluation of T cell reactivity to overlapping and mutated peptides, will be required to potentially identify the optimal peptide size, and the importance of individual residues in the recognition of this immunodominant epitope by the different T cell clones.

We also isolated two CD4<sup>+</sup> Th1 T cell clones which recognized the P2 peptide in vitro, but were unable to recognize the intact SRTA from which this peptide was derived. We find it curious that these T cell clones, isolated from animals immunized with RTA, can recognize the P2 peptide, but not the intact antigen. Although the explanation for this phenomenon is not readily apparent, the relative difficulty in isolating these clones (we were only able to isolate two) suggests that the target antigen of  $\alpha$ TBM disease, 3M-1, is only rarely processed into P2 (or a peptide containing a significant portion of P2) by I-A<sup>s</sup> antigen presenting cells in the SJL mouse.

The relevance of P3 (but not to P2) to the development of  $\alpha$ TBM disease was demonstrated through showing that the clones reactive to P3 were able to transfer disease to naive recipients, while a P2-reactive clone was not. In sum, these findings suggest that CD4<sup>+</sup>, Th1 helper T cells derived from the peripheral lymphoid system of animals immunized to produce  $\alpha$ TBM disease, and directed towards an immunodominant epitope contained within the P3 peptide, are nephritogenic. Interestingly, both the  $\alpha$ TBM antibody response and the CD8<sup>+</sup> effector T cell response are also directed at P1 (7, 27) (reactivity towards P3 has not been tested), confirming the importance of this antigenic epitope in the pathogenesis of  $\alpha$ TBM disease. Although our results demonstrate a single immunodominant peptide epitope relevant to the development of disease, it is possible that other, secondary, cryptic epitopes emerge within the kidney as the disease process progresses (58). Studies evaluating the peptide specificity of nephritogenic T cells derived from diseased organs will be required to address this issue.

Analysis of the TcR V $\beta$  genes used by the P3 epitopespecific T cell clones revealed the use of several V $\beta$  genes, with a recurrent use (3/8) of V $\beta$ 14. This 38% incidence of V $\beta$ 14 expression by P3-reactive clones, compared with 6.5% of the peripheral CD4<sup>+</sup> T cell population, provides evidence for the preferential use of the V $\beta$ 14 gene by the nephritogenic T cell clones. Collectively, our findings suggest that the anti-3M-1 TCR repertoire is oligoclonal. Whether these findings are representative of the TcR repertoire within diseased organs is an area of ongoing investigation. It should be noted however, that interpretation of TCR gene usage analyses in diseased whole organs is often hampered by the lack of definition of peptide specificity (16), and results may not correlate with findings derived from peptide specific T cell clones (16).

Perhaps more intriguing than the oligoclonality of the TCR response is the similarity of the deduced amino acid sequences across the  $V\beta/D\beta/J\beta$  junction. Based initially on comparative analyses between immunoglobulin and TcR sequences (59, 60), and later confirmed by statistical (61), and mutational (62-64) analyses, the V $\beta$ /D $\beta$ /J $\beta$  junction of the TcR in CD4<sup>+</sup> cells has been proposed to constitute the putative CDR3 region, and thus make direct contact with its peptide/MHC class II ligand. The statistically significant overrepresentation (compared with a random set of functional TcR $\beta$  chains) of a small nonpolar amino acid at the  $D\beta/J\beta$  junction, and an Arg or Lys at the  $V\beta/D\beta$  junction, both determined by N region additions or deletions, suggests that these residues may be important in the specific recognition of P3. Additionally, the CDR3 region of all of the P3-reactive clones is neutral or cationic (Arg/Lys), with a distinct lack of negative charges (Glu in position 101 of M36.6.1 is the only one, and is counteracted by an adjacent Arg, Fig. 8), suggesting a direct interaction with the anionic P3 peptide. Analogous findings have been noted in another model of autoimmune renal injury (8). The precise determination of the role played by individual residues within the CDR3 in the specific recognition of the P3/MHC class II complex will require a mutational analysis of the TCR. Finally, the sequence of the TCR  $\alpha$  chain needs to be defined before any definitive statements regarding the molecular basis of P3 recognition by autoreactive T cells can be made. Based on these experiments, however, it may be possible to direct specific immunotherapy at a common epitope within the TCR $\beta$  CDR3 region of nephritogenic, P3-reactive T cells.

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