T cells presenting viral antigens or autoantigens induce cytotoxic T cell anergy

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Introduction

Paraneoplastic cerebellar degeneration (PCD) is a rare type of paraneoplastic neurologic disease (PND) that can arise when patients develop tumors expressing cerebellar degeneration–related protein 2 (CDR2). CDR2 is a cytoplasmic protein that is robustly expressed in Purkinje cells of the cerebellum. Patients develop neurologic symptoms when tumor cells expressing CDR2 trigger a CDR2-specific immune response that crosses the blood-brain barrier, resulting in profound cerebellar dysfunction. The hallmark of the disease is the production of high-titer antibodies against CDR2, also known as Yo antibodies, and circulating effector cytotoxic T cells in the blood and cerebral spinal fluid (CSF) that produce IFN-γ in response to CDR2 (1–3). Interestingly, this CDR2-specific immune response also effectively lysed target cells in an HLA-restricted manner and recognizes tumor cells (1, 2, 4), resulting in occult or limited-stage tumors and relative tumor immunity (5). PCD is extraordinarily rare despite the fact that approximately 56% of ovarian tumors and approximately 22% of breast tumors ectopically express CDR2 (5, 6). Given that expression is normally limited to the Purkinje cells of the cerebellum, it stands to reason that the immune system should be relatively ignorant to CDR2, facilitating responses to this tumor antigen. However, a study of 553 serum samples of PND patients discovered that no patients with primary antibodies against any of the other PND antigens harbored additional antibodies against CDR2 (PCA-1) (7), though 19% to 50% of patients with PND harbored coexisting antibodies against other intracellular neuronal antigens. This suggests there is an increased threshold for breaking tolerance to CDR2, relative to other PND antigens. Understanding the mechanism of this clinical observation could uncover insights about tolerance to tumors and immune responses more generally. Here we report the surprising finding that T cells themselves express CDR2 and that the identity of the cell presenting CDR2 has a profound effect on effector CTL responses; the threshold for breaking tolerance to CDR2 is higher when it is expressed in T cells than in the CNS.
Results

Generation of Cdr2-knockout mice. A fundamental goal of studying PCD patients is to understand mechanisms underlying naturally occurring tumor immunity and autoimmune brain disease. In initial attempts to model tumor immunity to CDR2 in mice, we observed poor immune responses to CDR2 vaccination. To test whether this was related to development of tolerance to endogenous CDR2-derived antigens, we generated Cdr2-knockout (Cdr2-KO) mice for immunizations. Mice were generated using homologous recombination with a targeting vector containing EGFP (Figure 1A) replacing 12.5 kb of the Cdr2 gene (including all of the Cdr2 coding region after the first 14 bp of exon 2) with EGFP and an intact Cdr2 3′ UTR; the absence of Cdr2 expression and presence of EGFP expression was confirmed by Northern blot (Figure 1B) and Western blot (Figure 1C).

CDR2 expression in wild-type mice limits the generation of humoral and cellular immune responses to CDR2. Cdr2-KO mice mounted high-titer (>1:1,000) IgG responses to CDR2 vaccination, but wild-type (WT) mice did not, even when vaccinated with considerable doses of antigen (Figure 2A). Cdr2-KO mice were competent to produce antibodies against a control antigen (β-gal) (Figure 2A).

We also assayed T cell responses in WT and Cdr2-KO mice. We first sought to identify the naturally processed and immunodominant MHCI and MHCII epitopes in H-2b mice. We isolated CD4+ T cells from Cdr2-KO mice shown to have CDR2-specific antibodies in Figure 2A and in vitro stimulated the cells with MHCI (H-2Ab) predicted binding peptides. We found that CDR2-170 and -172 are naturally processed epitopes of CDR2 (Figure 2B). We next challenged Cdr2-KO and WT mice with adenovirus expressing CDR2 (AdV-CDR2) and adenovirus expressing β-gal (AdV-β-gal). WT CD4+ T cells produced poor responses to both CDR2-170 and -172 compared with CD4+ T cells from KO mice (Figure 2C). To identify MHCI epitopes, Cdr2-KO or WT mice were immunized with AdV-CDR2. CDR2-120 peptide–pulsed RMA-S cells...
elicited the highest level of IFN-γ production in Cdr2-KO hosts, and was not recognized in WT hosts (Figure 2D). Together, these data reveal a defect in antibody CD4+ and CD8+ T cell immune responses to endogenous CDR2 in WT hosts and that CDR2 tolerance is lacking in mice that do not express the CDR2 protein.

**CDR2 expression is more tolerant when expressed by T cells than by neurons.** To further explore the mechanism of tolerance to CDR2, we generated bone marrow chimeras and tested them for CDR2-specific CTL responses. Cdr2-WT or -KO mice were transplanted with Cdr2-WT or -KO bone marrow cells, and T cell responses were assessed 14 days after immunization with AdV-CDR2 by IFN-γ ELISPOT assay. CD8+ T cells isolated from KO mice transplanted with Cdr2-KO bone marrow produced IFN-γ in response to CDR2-120 directly ex vivo (Figure 3A) and after in vitro stimulation (Figure 3B). CD8+ T cells isolated from WT mice transplanted with Cdr2-KO bone marrow did not produce IFN-γ in response to CDR2-120 directly ex vivo (Figure 3A), but were able to be activated after in vitro stimulation (Figure 3B). In contrast, CD8+ T cells isolated from Cdr2-KO mice transplanted with WT bone marrow were unable to respond to CDR2 either directly ex vivo or with in vitro stimulation (Figure 3, A and B). Nonhematopoietic, radio-resistant cell (such as Purkinje cell) expression of CDR2 leads to partial tolerance to CDR2, as CDR2-specific T cell responses in this chimera (KO→WT) could be eventually demonstrated after in vitro stimulation. Interestingly, the most profound tolerance was seen in the Cdr2-KO hosts receiving WT bone marrow. In this scenario (WT→KO), CTL responses to CDR2 could not be recovered, even with subsequent in vitro...
stimulation. Taken together, these results suggest that CDR2 is expressed by a hematopoietic cell and that there are distinct thresholds of tolerance to CDR2, depending on its site of expression.

To address whether some hematopoietic cells express CDR2, we tested hematopoietic lineages for CDR2 expression, using knocked-in EGFP from Cdr2-KO mice as a surrogate marker. CD3+ T cells expressed the CDR2 marker, as measured by a clear shift in EGFP, whereas CD19, NK cells, MHCII+, and CD11c subsets did not (Figure 3C). A small population of CD11b+ cells expressed low levels of CDR2. These either represent phagocytic cells that have taken up dying T cells or a subpopulation of CD11b+ cells that express low levels of CDR2. To assess which of the hematopoietic cells mediate the robust tolerance seen in Figure 3, A and B, we compared CDR2-specific CTL responses in mice that were transplanted with Cdr2-KO bone marrow (positive control), WT bone marrow (negative control), or a mixture of recombinase-activating gene 1–KO (Rag1-KO) bone marrow and Cdr2-KO bone marrow. In this latter scenario, all lymphocytes are Cdr2-KO, while the nonlymphocytic hematopoietic cells are a mix of Cdr2-KO and Cdr2-WT. CDR2 vaccination of these mice, lacking Cdr2 specifically in lymphocytes, produced CTL responses that are not significantly different from Cdr2-KO mice transplanted with Cdr2-KO bone marrow (Figure 3D). Taking these results
together, we conclude that CDR2-expressing CD3+ T cells are sufficient to confer tolerance to CDR2.

**Human T cells also express CDR2 but are not targeted in patients with PCD.** To directly test for CDR2 expression in hematopoietic cells, we assessed CDR2 expression by Western blot of human CD4+ T cells. We observed that these cells express readily detectable levels of CDR2, as assessed by both a monoclonal antibody against CDR2 and PCD patient serum (Figure 4A). The results were supported by RNA analyses; CDR2 RNA was robustly expressed, with low mean cycle time values, in CD4+ and CD8+ T cells. These values were comparable to those seen in the positive control tissue types, cerebellum and HeLa cells. CDR2 cycle time was much higher in neutrophils, indicating that the expression in this cell type is lower than in T cells or cerebellum, but unlikely to be nonexistent (Figure 4B). To test whether the Western blot signal could be due to antibody cross reactivity with the CDR2 paralog, CDR3, we investigated RNA expression using PCR primers specific for CDR3 in human T cells. Quantitative PCR (qPCR) demonstrated that CDR3 mRNA is robustly expressed in cerebellar lysate and HeLa cells but undetectable in CD8+ and CD4+ T cells (Figure 4B). Taken together, these results demonstrate that CDR2 is expressed in human T cells.

Patients with PCD and clear evidence of autoimmunity to CDR2 are not known to suffer from lymphopenia, prompting us to assess white blood cell counts from our previously described (8) cohort of 11 patients with high-titer antibodies against CDR2 and recent neurologic decline (suggesting acute flare of disease). We found that PCD patients had no leukopenia or lymphopenia in the course of 16 flares among 11 patients (Figure 4C). We conclude that patients with PCD and autoimmune targeting of CDR2 suffer neurologic decline without T cell depletion, consistent with in vivo tolerance to T cell CDR2 expression in humans.
Effector CTLs produce less IFN-γ in response to antigen presented by T cells compared with KECs. To explore whether T cell tolerance to CDR2 is due to differences in antigen presentation by T cells, we compared responses of murine effector CTL clones to either Cdr2-KO kidney epithelial cells (KECs) or T cells pulsed with various peptides. NP indicates pulsed with irrelevant control influenza peptide and CDR2-120 indicates pulsed with cognate peptide. Each bar represents the mean of triplicate wells and error bars are standard deviations. (A) Surface staining of MHCI and costimulatory molecules in DCs (black line), live T cells (green line), KECs (blue line), and isotype control (shaded). These data are representative of 3 experiments. ***P < 0.001, as calculated using unpaired Student’s t test. SFC, spot-forming cells.

T cell antigen presentation of virus also limits effector CTL responses. To test whether the lack of effector CTL activation in our system could be an idiosyncratic feature of the CDR2 protein or is generalizable to other antigens, we assessed T cell responses to T cells presenting viral antigens, using the well-characterized flu system. We challenged mice with live flu virus and transferred NP or OVA-pulsed CD8+ T cells and measured expansion of NP-specific host CD8+ T cells by tetramer assay. Peripheral blood from mice infused with NP-pulsed CD8+ T cells contained significantly fewer NP-specific CD8+ T cells than those that received OVA-pulsed CD8+ T cells (Figure 6A). As a control, we found no difference in transferred NP or OVA-pulsed CD8+ T cells recoverable at day 8 (data not shown). While intravenous injections of free peptide at high doses (0.1 mg) can limit CTL responses in vivo (9), the CD8+ T cells in this experiment were pulsed with much lower amounts (1 μM), making it unlikely that low amounts of dissociating free peptide could account for these results. Instead, we attribute the decrease in NP-specific CD8+ T cells to the transfer of NP-pulsed CD8+ T cells.

To test the effect of T cell presentation of flu peptides in vitro, we measured NP-specific effector CTL responses to NP- or OVA-pulsed and irradiated DCs, KECs, CD8+, or CD4+ cells. While NP-specific effector CTLs robustly produce IFN-γ in response to NP-pulsed DCs and even KECs, which are not professional antigen-presenting cells, they produce significantly less IFN-γ in response to NP-pulsed CD8+ and CD4+ lymphocytes (Figure 6B). Addition of IL-2 rescued the ability of CD8+ lymphocytes to activate CTL IFN-γ production in an antigen-specific manner (Figure 6C). Taken together, these data indicate that antigen presentation by T cells attenuates flu-specific CTL expansion in vivo and induces flu-specific effector CTL anergy in vitro that can be recovered with IL-2.

Discussion
PNDs provide a unique window to examine mechanisms of immunity and tolerance. Here, we describe our discovery that CTLs capable of responding to a variety of cell types presenting the CDR2 PND antigen do
not become activated in response to lymphocyte presentation of the antigen. The observation that T cell–mediated antigen presentation regulates CTL function is reminiscent of Jerne’s theory of network regulation of immune responses (10). The repertoire of B and T cell receptors represents a collection of stochastic DNA VDJ gene rearrangements as well as mutations. These somatic mutations enable the diversity required to counter a nearly infinite possibility of pathogens but also generates T cell receptors that are neoantigens and therefore risk autoimmunity. Mechanisms downstream from initial immune tolerance checkpoints

Figure 6. T cell antigen presentation induces effector CTL split anergy. (A) Flow cytometry of circulating CD45.2+ NP-specific tetramer+ CD8+ T cells in a flu-infected host transferred with either NP-pulsed (n = 10) or OVA-pulsed (n = 10) CD45.1+ CD8+ T cells. Each bar represents the mean and error bars are standard deviations. These data are representative of 2 experiments. (B) IFN-γ ELISPOT assay of NP-specific effector CTLs cultured with DCs or kidney epithelial cells (KECs) (stimulator to effector ratio of 1:30), or CD8+ or CD4+ T cells (stimulator to effector ratio of 1:1). Each bar represents the mean of triplicate wells and error bars are standard deviations. These data are representative of 2 experiments. (C) IFN-γ ELISPOT assay of NP-specific effector CTLs cultured with KECs (stimulator to effector ratio of 1:30), CD8+ T cells (stimulator to effector ratio of 1:1), or CD8+ T cells plus IL-2. Stimulator cells were pulsed with either NP or OVA peptide. Each bar represents the mean of triplicate wells and error bars are standard deviations. These data are representative of 2 experiments. *P < 0.05, ***P < 0.001; ns, statistically not significant as calculated using unpaired Student’s t test. (D) Proposed model of the plasticity of CTL responses to antigen encounters and how the data presented in this paper may relate to human diseases. In the steady state (peripheral blood), armed effector CTLs respond to other T cells presenting cognate antigen by killing the stimulator T cell, but do not secrete IFN-γ. The effector CTL itself survives and is anergic. In the context of an inflammatory microenvironment (tumor or other inflamed tissue), exogenous IL-2 licenses the effector CTL to become fully activated and it secretes IFN-γ in addition to killing targets. IFN-γ augments immune responses upregulating antigen processing and presentation as well as promoting the recruitment of leukocytes to the tissue, promoting tumor immunity and autoimmunity seen in paraneoplastic neurologic disease. SFC, spot-forming cells.
CD4+ T cells in which latent HIV-1 infection has been reversed. In contrast, CD8+ T cells from patients infected with HIV maintain a CD4+ T cell count greater than 500 without antiretroviral therapy and T cell reservoirs of HIV are a major obstacle to HIV eradication (24). A small percentage of people essential for successful eradication (26). CD8+ T cells derived from elite controllers kill autologous are referred to as elite controllers. CTL activation is a correlate of HIV elite control (25) and may be previous work evaluating cytotoxic CD8+ T cell responses to T cell antigens, with the exception of one group that showed that T cell presentation of the H-Y antigen induces anergy and apoptosis in a noncytotoxic CD8+ T cell line (18). Through our efforts to better characterize the clinical phenotype of patients with PCD, we demonstrate that intracellular antigen expression by lymphocytes in vivo induces effector CTL anergy.

**Implications for PCD.** CTL anergy to lymphocyte antigen expression may be a necessary element of immune homeostasis. In this way, autoimmune disease may be suppressed, including autoimmunity to CDR2, which rarely develops in patients with breast or ovarian cancer, despite the fact that 22%–56% of these tumors express CDR2 (6). Lymphocyte expression and CDR2 tolerance may therefore help explain the low incidence of PCD (19). The vast majority of patients harboring gynecologic tumors expressing CDR2 are likely anergic to CDR2. We demonstrate here that T cell CDR2 expression underlies a particularly high threshold for mounting an immune response to CDR2. The robust tolerance to this antigen may make it a difficult tumor rejection antigen in vivo. Tumor antigens that are not expressed in T cells, and do not tolerize, are likely to be more effective targets for tumor vaccines.

Levels of tolerance to PND-associated antigens appear to vary depending on the antigen and its sites of expression. The high degree of tolerance to CDR2 appears to differ from that of other PND antigens such as Nova (the target antigen in paraneoplastic opsoclonus myoclonus ataxia) and HuD (the target antigen in the Hu syndrome). HuD and Nova expression are highly restricted to neurons, and effective priming of a high-affinity T cell population specific to the HuD antigen can be detected after in vitro stimulation (20, 21). This differs from CDR2-specific T cell responses, which cannot be detected in vivo, or after ex vivo stimulation in a Cdr2-WT mouse.

The finding that T cell clones produce less IFN-γ in response to CDR2 peptide–pulsed T cells than KECS (Figure 5B) is consistent with the clinical observation that patients with PCD and CDR2-specific autoimmunity (high-titer antibodies and cytotoxic CD8+ T cells capable of targeting CDR2-expressing cells [other than T cells]) do not suffer from autoimmune targeting of T cells or lymphopenia (1, 2). The discrepancy between the active immune clearing of CDR2-expressing Purkinje cells in the CNS and the lack of targeting lymphocytes in the circulation of patients with established PCD indicates that the local microenvironment is a critical component of CTL activation. Local inflammatory mediators, such as IL-2 and possibly others, in the tumor and CNS likely account for the distinct immune outcomes of CDR2-specific CTL interactions in CNS, tumor, and blood. A model of how the data presented in this manuscript may relate to human disease is presented in Figure 6D. Considering most of the data presented here describe in vitro experiments, further experiments in vivo are needed to refine a more definitive model.

**Implications for lymphotropic viruses.** We also demonstrate that effector CTL anergy in response to lymphocyte-derived antigen is a more broadly relevant phenomenon, as we find the observations regarding CTL anergy extend to lymphocytes presenting viral flu antigens. T cell antigen presentation of flu peptide fails to activate effector CTL IFN-γ production (Figure 6). Defective IFN-γ production by CTLs leads to defective viral clearance (22, 23) likely because an orchestrated immune response is required for clearance and IFN-γ production is important for recruitment of other innate effector immune cells. We show that addition of IL-2 rescues T cell IFN-γ activation, which defines an anergic phenotype. The discovery that antigen presentation by T cells induces anergy in effector cytotoxic T cells is of particular clinical relevance as it suggests that viruses that infect lymphocytes, such as HIV, may preferentially inhabit a lymphocyte niche because it is a relatively immune-privileged site. CD4+ T cell reservoirs of HIV are a major obstacle to HIV eradication (24). A small percentage of people infected with HIV maintain a CD4+ T cell count greater than 500 without antiretroviral therapy and are referred to as elite controllers. CTL activation is a correlate of HIV elite control (25) and may be essential for successful eradication (26). CD8+ T cells derived from elite controllers kill autologous CD4+ T cells in which latent HIV-1 infection has been reversed. In contrast, CD8+ T cells from patients maintained on chronic highly active antiretroviral therapy (HAART) do not produce IFN-γ ex vivo. Usual (non-elite controller), chronically HIV-infected patient–derived CD8+ T cells required prestimu-
lation with IL-2 and HIV-derived (gag) peptides to overcome functional defects in order to proliferate and become activated in response to HIV-infected T cells (26). Future efforts to overcome effector CTL energy to antigen expressed in lymphocytes may help overcome this challenge.

**Methods**

**Generation of Cdr2-KO mice.** The Cdr2 targeting vector consisted of Cdr2 intron 1 and the beginning of exon 2, EGFP, the human GAPDH gene, a neomycin resistance cassette flanked by loxP sites, and the Cdr2 3′UTR and 3′ flanking sequence. The vector was electroplated into embryonic stem (ES) cells. Neomycin-resistant lines were expanded and recombination was confirmed using long-range PCR. An ES cell line with the desired genomic structure was injected into blastocysts, resulting in male chimeras. For the FLP recombination, chimeras were bred with 129S4/SvJae-Gt(Rosa)26Sor<sup>tm1(FLP1)Dym</sup>/J mice (The Jackson Laboratory, strain 003946) to excise the GAPDH and NEO cassette. Resulting mice were backcrossed for 13 generations to C57BL/6 mice (The Jackson Laboratory). The colony was maintained by breeding heterozygotes. Both Cdr2-KO and control WT mice were obtained from this breeding strategy. All mice were housed in pathogen-free facilities at Rockefeller University.

**Northern blot.** Total RNA from mouse cerebellum was prepared by TRIzol (Invitrogen) extraction and column purification using a High Pure RNA Isolation Kit (Roche). Total RNA (10 µg per lane) was separated with 0.8% SeaKem Gold agarose (Lonza) along with Millennium RNA Markers (Ambion). The gel was treated with 0.05 M NaOH and 1.5 M NaCl for 10 minutes, and then with 0.5 M Tris, pH 7.5 containing 1.5 M NaCl for 10 minutes, before being equilibrated for blotting in 20× SSC for 20 minutes. The gel was blotted onto a Hybond-N filter (Amersham). Probes were prepared using a Prime-It II Random Primer Labeling Kit (Agilent) with [α-<sup>32</sup>P]dCTP. For hybridization, labeled probes were used at 1 × 10⁶ cpm/ml of hybridization solution (1% BSA fraction V, 7% SDS, 0.5 M NaH₂PO₄, pH 7, 1 mM EDTA, pH 8). The filter was hybridized overnight at 65°C and washed with 2× SSC with 0.05% SDS at 60°C 2 times for 10 minutes each, and then with 0.1× SSC with 0.1% SDS at 60°C 2 times for 30 minutes each.

**IP Western blot.** PCD patient antibodies were purified from serum by column chromatography with a 1:1 mixture of protein A and protein G (Invitrogen) (27). Purified PCD patient antibodies were then conjugated to M-270 Epoxy Dynabeads (Invitrogen). Conjugation was performed using the method described by Cristea and Chait (28).

Cerebella from WT mice and Cdr2-KO littermates were homogenized in 1× PBS with 1% NP-40, 0.5% Na deoxycholate, 0.1% SDS, 20 mM N-ethylmaleimide, and Complete EDTA-free protease inhibitors (Roche). Protein concentrations were adjusted to 6 mg/ml and proteins were used as input for each IP reaction with 6 mg of PCD antibody–conjugated Dynabeads. After a 30-minute immunoprecipitation at 4°C, beads were washed once with lysis buffer, and then 4 times with 1× PBS/0.5% NP-40. Captured proteins were then eluted with 0.5 M NH₄OH/0.5 mM EDTA. Eluates were frozen in liquid nitrogen and then dried completely in a SpeedVac. These eluates were then analyzed by Western blotting using affinity-purified chicken antibodies directed against the CDR2 peptide RKPLKRSSSET, corresponding to amino acids 302–312.

**Peptides, antibodies, and mice.** CDR2 p120 (CLQTNIDHL), CDR2 p151 (EKPAFSFSCL), CDR2 p285 (PSQSLLEEM), CDR2 p348 (VDTQYSAL), CDR2 p64 (QEIEYLTKQVELLRQMN), CDR2 p170 (LRQHFYDHVAEIKITS), CDR2 p172 (QHFVYDHVFAEIKITSLQ), CDR2 p242 (ATDAYRARALELEAEVA), CDR2 p352 (VDTQYSALVKYEELLK), β-gal p726 (NLSVTLPAASHAIPH), β-gal p96 (DAPITYTNV), A/PR/8 NP p366 (ASNENMETM), and OVA p323 (ISQAVHAAHAEINEAGR) were HPLC purified. Peptides for CDR2 (>95% pure) were obtained from Life Technologies and all other peptides were from American Peptide, Inc. All antibodies used in the analysis of cells by FACS were obtained from BD Pharmingen; antibodies used in ELISPOTs were from Mabtech. The Cdr2-KO colony was maintained by breeding heterozygotes. Both Cdr2-KO and control WT mice were obtained from this breeding strategy. C57BL/6 mice (stock number 000664), Rag1-KO mice (B6.129S7-Rag<sup>tm1Sor</sup>/J, stock number 002216), and CD45.1 mice (B6.SJL-Ptprc<sup>Pepc</sup>/BoyJ, stock number 002014) were purchased from The Jackson Laboratory. CD45.1 Rag1-KO mice were bred for bone marrow transplantation studies. In all experiments, mice were 6–8 weeks old and housed in pathogen-free facilities at Rockefeller University.
**CDR2 inclusion bodies.** The full-length coding sequence of mouse Cdr2 was ligated into the bacterial expression vector pET-28c (EMD Millipore) to create pET28-CDR2. This plasmid was transformed into Rosetta 2 (DE3) (EMD Millipore) competent cells and the resulting cells were plated on LB plates with kanamycin and chloramphenicol. After overnight growth at 30°C, a single colony was used to inoculate an overnight starter culture. For protein production, the starter culture was grown to an OD600 of 0.6 and cells were induced with 0.4 mM IPTG and grown for another 40 minutes. Cells were harvested by centrifugation and the cell pellet was frozen in a dry ice/ethanol bath. The CDR2 protein present in these cells as inclusion bodies was purified using BugBuster Master Mix (EMD Millipore) according to the manufacturer’s protocol. The purified CDR2 inclusion bodies were resuspended in PBS by sonication. Integrity of the protein was verified by SDS-PAGE analysis and the protein concentration was estimated by comparison to a BSA standard curve.

**Generation of antibody in mice.** Incomplete Freund’s Adjuvant (Difco) was reconstituted with 5 mg/ml *Mycobacterium tuberculosis* (Difco) to make Complete Freund’s adjuvant (CFA). A 1:1 emulsion was made by mixing 10 mg/ml β-gal protein (Roche) or 5 mg/ml CDR2 derived from bacterial inclusion bodies and CFA. Each mouse received 4 × 50 μl emulsate in their lower back and pertussis toxin on days 0 and 2. Mice were bled and tested for antibody reactivity using in vitro translated/transcribed (TNT Quick Coupled Transcription/Translation Rabbit Reticulocyte Lysate system [T7 promoter version, Promega]) CDR2 and β-gal 21 days after immunization.

**Cell lines and influenza virus.** The RMA-S cell line was obtained from Stanley Nathenson (Albert Einstein College of Medicine, Bronx, New York, USA). KECs were obtained by organ culture for 5 days in DMEM with 10% FCS (GIBCO). Adherent cells were given recombinant murine IFN-γ (50 ng/ml, R&D Systems) for 48 hours prior to use in assays. Human influenza A/PR/8 virus in allantoic fluid was obtained from Charles River Laboratories. HeLa cells were obtained from the American Type Culture Collection.

**Immunization of mice with AdV-CDR2, AdV-β-gal, or AdV-Trk.** Serum-free viruses were obtained from PureSyn, Inc. The virus was provided at a concentration of 10⁶ particles/ml. Virus (100 μl) was injected intradermally. Pertussis toxin (PTx, Sigma-Aldrich) 400 ng was injected i.p. in each mouse on day 0 and day 2.

**Bone marrow chimeras.** Host mice were lethally irradiated with a cumulative dose of 900 cGy (2 × 450 cGy). Four hours after the last irradiation, host mice were reconstituted with a total of 10 × 10⁴ bone marrow cells (Thy1.2⁺ cells depleted). Mixed bone marrow chimeras were generated using 5 × 10⁶ cells from each donor bone marrow. Seventy to 95 days after reconstitution, 50:50 reconstituted mice were injected with 100 μl purified AdV-CDR2 (10⁷ particles/ml) intradermally and were treated with PTx (400 ng/mouse) i.p. on days 0 and 2. Thirteen days later, CD8⁺ cells were isolated from the spleens and CD8⁻ cells were assayed in a 40-hour IFN-γ ELISPOT.

**T cell isolation and ELISPOT Assay.** CD4⁺ and CD8⁺ T cells were isolated by positive selection using MACS beads (Miltenyi Biotec). ELISPOT assays were performed as previously described (29). Briefly, 200,000 isolated CD4⁺ or CD8⁺ T cells were plated respectively with 250,000 irradiated Thy1.2⁻ depleted cells. Mixed bone marrow chimeras were generated using 5 × 10⁶ cells from each donor bone marrow. Seventy to 95 days after reconstitution, 50:50 reconstituted mice were injected with 100 μl purified AdV-CDR2 (10⁷ particles/ml) intradermally and were treated with PTx (400 ng/mouse) i.p. on days 0 and 2. Thirteen days later, CD8⁺ cells were isolated from the spleens and CD8⁻ cells were assayed in a 40-hour IFN-γ ELISPOT.

**Patient samples.** Patients positive for the presence of Yo antibodies (PCD) were enrolled in this Rockefeller University IRB-approved study at The Rockefeller University Hospital (RDA-572). All patients gave written consent. PCD patients and healthy controls were leukapheresed and CD4⁺ T cells were isolated from peripheral blood mononuclear cells by positive selection using MACS beads (Miltenyi Biotec).

**Western blot of CD4⁺ T cells from PCD patients and healthy donors.** Pelleted CD4⁺ T cells were lysed using 5× Passive Lysis Buffer (Promega) supplemented with Complete Mini, EDTA-free protease inhibitors (Roche) and protein concentration was determined using a BCA Protein Assay kit (Pierce). For each sample, 37 μg of total protein from CD4⁺ T cells was resolved by 10% SDS-PAGE and transferred to a PVDF membrane. Membranes were first probed with anti-CDR2 rabbit monoclonal antibody diluted 1:1,000 (clone EPR9464, Abcam), stripped, and reprobed with PND patient serum 1:10,000, and then stripped and incubated with normal serum diluted 1:10,000, and finally stripped and reprobed with anti-GAPDH diluted 1:20,000 (clone 6C5, Life Technologies). HRP-conjugated secondary antibodies diluted 1:10,000 (The Jackson Laboratory) and ECL (PerkinElmer) were used to detect membrane-bound antibodies.
qPCR. mRNA was isolated using a HiPure kit (Roche). First-strand cDNA synthesis of total RNA was performed with iScript (Bio-Rad). Primer pairs used to detect gene expression in human tissue were CDR2F, 5′-CGTCACAGCAAGGGAACTGGAA-3′ and CDR2R, 5′-CACTTGGCTCTGGAGGTGATCA-3′; CDR3F, 5′-CAGGACCTGGACAGCAGACT-3′ and CDR3R, 5′-ACCTGTTTCTCATTGTTGGA-3′; HRP14F, 5′-CGGAGCTGACCAGACTTTTC-3′ and HRP14R, 5′-GGTTCGACCGTCATACTTCTTC-3′.

CDR2 T cell clone. The CDR2-120–specific T cell clone was generated by immunizing Cdr2-KO mice with AdV-CDR2. After in vitro stimulation, CD8+ T cells were isolated and cloned by limited dilution. A high-affinity clone was selected by screening for responses to greater than 10⁻⁹ M CDR2-120 peptide–pulsed RMA-S cells. The clone was assayed and restimulated weekly with 10⁻⁹ M CDR2-120 peptide–pulsed irradiated splenocytes and 10 U/ml rhIL-2 (Chiron).

DCs. Bone marrow–derived DCs were generated as previously described (30). Briefly, bone marrow was isolated from Cdr2-KO mice and cultured for 6 days in 10% supernatant from GM-CSF–transfected J558L cells. On day 6, Cdr2-KO DC cultures were harvested and cultured in the presence of GM-CSF and 80 ng/ml recombinant mouse TNF-α (R&D Systems) was added as a maturation stimulus. Forty-eight hours later, CD11c+ cells were isolated and used as stimulators in the ELISPOT.

In vivo transfer of antigen-pulsed CD8+ T cells. C57BL/6 mice were simultaneously given 300 hemagglutination units (HAU) A/PR/8 virus i.p. and 5 × 10⁶ NP-pulsed and 3-times-washed CD45.1 CD8+ T cells intravenously. On day 8, 100 μl of blood was red blood cell lysed with ACK (GIBCO) and then coincubated with NP-D₃-tetramer (MBL), CD45.2 (clone 104), and CD8a (clone 53-6.7) for 30 minutes at 6°C. Dead cells were excluded using TO-PRO-3 (Thermo Fisher Scientific) and immediately acquired by FACS.

Generation of NP-specific effector CTLs. C57BL/6 were immunized with 300 HAU of A/PR/8 virus i.p. Fifteen days after immunization, spleens were harvested and 5 × 10⁶ cells/ml were stimulation with 1 μM NP-specific peptide. On day 5 of NP peptide stimulation, CD8+ cells were isolated using MACs positive selection (Miltenyi Biotec). Where indicated, 20 U/ml of rhIL-2 (Chiron) was added to the assay.

Statistics. Data were analyzed using Prism software (GraphPad). Standard deviations are reported for ELISPOT data and unpaired Student’s t test (2-tailed) was used to compare 2 groups. A P value of less than 0.05 was considered significant.

Study approval. The present studies in humans were IRB approved at The Rockefeller University Hospital (RDA-572). All patients gave written informed consent. Animal procedures were performed in compliance and were approved by The Rockefeller University Institutional Animal Care and Use Committee regulations at The Rockefeller University.

Author contributions
NEB, DEO, ECG, BDS, GCC, and RBD conceived the study. NEB and GCC were responsible for the methodology. NEB, DEO, ECG, BDS, GCC, TRM, JF, KJO, ZR, SP, and MOF carried out the investigation. NEB and DEO wrote the original draft of the manuscript. NEB, DEO, ECG, BDS, GCC, TRM, JF, KJO, ZR, SP, MOF, MJM, and RBD reviewed and edited the manuscript. RBD acquired funding.

Acknowledgments
We would like to thank the patients with PCD and their families, for their participation in research as well as the staff of the Rockefeller University Hospital and grant UL1 TR000043 from the National Center for Advancing Translational Sciences (NCATS), NIH Clinical and Translational Science Award (CTSA) program. This work was also supported by the NIH grant R01 NS034389. We would like to thank the Rockefeller Gene Targeting Facility for generating the ES cells resulting in the generation of the mice incorporating the targeted Cdr2 gene KO. RBD is an Investigator of the Howard Hughes Medical Institute (HHMI). GCC was supported by an HHMI Predoctoral Fellowship.

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