Prior immunity to influenza A virus (IAV) in mice changes the outcome to a subsequent lymphocytic choriomeningitis virus (LCMV) infection and can result in severe lung pathology, similar to that observed in patients that died of the 1918 H1N1 pandemic. This pathology is induced by IAV-specific memory CD8^+ T cells cross-reactive with LCMV. Here, we discovered that IAV-immune mice have enhanced CD4^+ Foxp3^+ T-regulatory (Treg) cells in their lungs, leading us to question whether a modulation in the normal balance of Treg and effector T-cell responses also contributes to enhancing lung pathology upon LCMV infection of IAV-immune mice. Treg cell and interleukin-10 (IL-10) levels remained elevated in the lungs and mediastinal lymph nodes (mLN)s throughout the acute LCMV response of IAV-immune mice. PC61 treatment, used to decrease Treg cell levels, did not change LCMV titers but resulted in a surprising decrease in lung pathology upon LCMV infection in IAV-immune but not in naive mice. Associated with this decrease in pathology was a retention of Treg in the mLN and an unexpected partial clonal exhaustion of LCMV-specific CD8^+ T-cell responses only in IAV-immune mice. PC61 treatment did not affect cross-reactive memory CD8^+ T-cell proliferation. These results suggest that in the absence of IAV-expanded Treg cells and in the presence of cross-reactive memory, the LCMV-specific response was overstimulated and became partially exhausted, resulting in a decreased effector response. These studies suggest that Treg cells generated during past infections can influence the characteristics of effector T-cell responses and immunopathology during subsequent heterologous infections. Thus, in humans with complex infection histories, PC61 treatment may lead to unexpected results.

During a lifetime the immune system is shaped by a history of infections. Prior infections with one pathogen may influence the severity of disease outcome to a subsequent infection with an unrelated pathogen, a phenomenon known as heterologous immunity (1). Enhanced immunopathology, which can be mediated by the activation of cross-reactive memory T cells, is one of the harmful consequences of heterologous immunity. For instance, it has been proposed during human infections that cross-reactive IAV-specific memory CD8^+ T cells can contribute to the induction of severe fulminant hepatitis during hepatitis C virus (HCV) infection and induction of acute infectious mononucleosis during Epstein-Barr virus (EBV) infection (2–4).

Lung pathology is a common manifestation of respiratory infections and can vary greatly in severity in different individuals infected with the same pathogen. To investigate the role of altered immunopathology during heterologous immunity in a controlled experimental setting, we utilized a mouse model of IAV-immune mice infected with lymphocytic choriomeningitis virus (LCMV) (5). We initially chose these two viruses because they are phylogenetically unrelated and because they are naturally spread through infection of the respiratory mucosa and induce significant inflammation in the lung (6–11). Influenza virus is an extremely common respiratory pathogen in humans, and LCMV, which induces a flu-like illness in humans, is also a relatively common pathogen, with 5 to 14% of the general population being serologically positive (12). These IAV-immune mice infected with LCMV could develop acute lung injury similar to that seen in individuals that died during the H1N1 IAV pandemic in 1918, with enhanced bronchus-associated lymphoid tissue (BALT), mononuclear pneumonia, necrotizing bronchiolitis, vasculitis, and bronchiolization (13, 14). The severity of lung pathology varied among genetically identical mice from mild pneumonitis to severe mononuclear necrosis, necrotizing bronchiolitis, and bronchiolization, an abnormal alveolar epithelial repair process considered premalignant and associated with idiopathic pulmonary fibrosis in humans. Although counterintuitive, severity of pathology did not directly correlate with LCMV titers. Instead, increased pathology was dependent on cross-reactive IAV-specific memory CD8^+ T cells (15). Disease severity was directly correlated with and could be predicted by the frequency of two IAV epitope-specific CD8^+ T-cell populations, PB1703 and PA224, which are cross-reactive with LCMV-GP34 and -GP276, respectively. Eradication or functional ablation of these pathogenic populations of IAV-specific memory T cells using mutant viral strains, peptide-based tolerization strategies, or short-term anti-gamma interferon (IFN-γ) treatment prevented this pathology.

Here, we continue to investigate this mouse model to determine if there are other contributing factors responsible for this variation in lung pathology and to define potential therapies. At major mucosal interfaces such as the lung, which is frequently

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exposed to foreign antigens, discrimination between innocuous and foreign antigen-specific immune responses is necessary to limit chronic inflammation. T-regulatory (Treg) cells have been shown to be key mediators in balancing inflammation and in inhibiting immune-mediated tissue damage, especially in organs like the lung and gastrointestinal tract (16–18). Both natural and induced Treg cells can suppress the function of many types of immune cells, including CD8<sup>+</sup> and CD4<sup>+</sup> T cells, B cells, dendritic cells (DC), NK cells, and NKT cells either by direct contact or by production of inhibitory cytokines, such as interleukin-10 (IL-10) and transforming growth factor beta (TGF-β) (19, 20). Treg cells have been intensively studied in autoimmunity, tumors, and persistent infections (19, 21–24). Increased numbers of Treg cells and a loss of functional virus-specific effector T cells are reported in persistent virus infections, such as hepatitis C virus (HCV), human immunodeficiency virus (HIV), Friend virus (FV), and herpes simplex virus (HSV), but not in acute virus infection (19, 25–28). Depletion of the suppressive Treg cells during a persistent retroviral infection resulted in enhanced effector T-cell function and reduced viral load (28, 29). Treg cells can also prevent extensive immunopathology during viral infections (24, 30, 31). Depletion of natural Treg cell responses, using PC61 treatment (anti-CD25) prior to infection, enhanced antiviral responses without any evidence of enhanced immunopathology if HSV-1 was injected into the footpad (32). However, Treg cell depletion prior to corneal HSV-1 infection resulted in severe T-cell-mediated tissue lesions (33). These results suggest that Treg cells influence disease outcome during viral infection differently depending on the virus and on the site of infection. Interestingly, compared to naive age-matched mice, we observed enhanced numbers of CD4<sup>+</sup> Foxp3<sup>+</sup> T reg cells in the lungs, spleens, and draining lymph nodes of mice immune to the nonpersistent virus IAV. Therefore, we were interested in determining whether Treg cells that expanded following a viral infection could also play a role during heterologous immunity, modifying immunopathology during a subsequent unrelated infection.

**MATERIALS AND METHODS**

**Viruses and infections of mice.** Six-week-old C57BL/6 (H-2b) male mice were obtained from The Jackson Laboratory (Bar Harbor, ME). The mice were anesthetized by inhalation of metofane (Pitan-Moore, Mundelein, IL) and infected intranasally (i.n.) with 70 PFU of the mouse-adapted IAV A/PR/8/34 (H1N1) or were inoculated with phosphate-buffered saline (PBS) as a control. This virus was prepared from eggs and was not exposed to bovine serum (5). After the immune system had returned to homeostasis (6 weeks or longer), immune and control mice were challenged i.n. with 1 × 10<sup>6</sup> PFU of LCMV clone 13 (Fig. 1e).

In experiments examining activation of cross-reactive memory CD8<sup>+</sup> T cells, mice were infected intraperitoneally (i.p.) with 5 × 10<sup>4</sup> PFU of LCMV Armstrong or were inoculated with PBS as a control. After the immune system had returned to homeostasis, LCMV-immune and con-
trol mice were challenged i.p. with $2 \times 10^5$ PFU Pichinde virus (PV; AN3739 strain) (see Fig. 6a). LCMV and PV were propagated in BHK21 baby hamster kidney cells. The LCMV stock was diluted 1:100 in serum-free media, and the stocks of PV were purified through a sucrose gradient and diluted in Hanks’ balanced salt solution (34). Immune mice were sex and age matched to control mice and housed under exactly the same pathogen-free conditions for the same time period. All experiments were done in compliance with institutional guidelines as approved by the Institutional Animal Care and Use Committee at the University of Massachusetts Medical School.

**Virus titration.** A 10% homogenate of lung and spleen tissue from each individual mouse was serially diluted and titrated on Vero cells to quantify LCMV PFU (35). Titers were reported as the arithmetic log10 PFU each individual mouse was serially diluted and titrated on Vero cells to quantify LCMV PFU (35). Titers were reported as the arithmetic log10 PFU

**Lung histological evaluation.** At days 7, 9, and 12 post-LCMV challenge, lungs from naive, IAV-immune, and Treg cell-depleted IAV-immune mice were collected, fixed in 10% neutral buffered formaldehyde, and embedded in paraffin. Tissue sections (5 μm) were stained with hematoxylin and eosin (H&E) and analyzed microscopically. Scoring of lung pathology by a pathologist blinded to the experimental design was graded based on a scale described in previously published studies (3, 15): 1, mild interstitial mononuclear infiltrates, disorganized BALT, and perivascular edema; 2, moderate interstitial mononuclear infiltrates, small amount of organized BALT, and pulmonary edema; 3, moderate interstitial mononuclear infiltrates, pulmonary edema, enhanced organized BALT, mild bronchiolization, and mild consolidation; 4, severe interstitial mononuclear infiltrates, greatly enhanced pulmonary edema, enhanced organized BALT, moderate bronchiolization, moderate consolidation, and moderate necrotizing bronchiolitis; 5, severe interstitial mononuclear infiltrates, greatly enhanced pulmonary edema, enhanced organized BALT, severe bronchiolization, severe consolidation, severe necrotizing bronchiolitis, and vasculitis involving more than half of the lung. The scoring on each individual mouse was done on 4 to 5 different sections of the lung, representing 4 to 5 different lobes of the whole lung assessing both qualitative and quantitative changes in histology. The histology photographs showing high-power 10× views of a small portion of one lobe of the lung demonstrate examples of the types of pathology observed in different treatment groups, but evaluations of the whole lung were used for scoring.

**Cell surface and tetramer staining.** Single-cell suspensions were prepared from spleens, mediastinal lymph nodes (mLN), and lungs, and erythrocytes were lysed with 0.84% NH₄Cl solution. Splenocytes were stained as described previously (35) using CD8α (53-6,7), CD4 (GK1.5), CD44 (IM7), PD-1 (J43), and Vβ-specific monoclonal antibody (Mab). All surface antibodies were purchased from BD Biosciences (San Jose, CA). For tetramer staining, cells were incubated for 60 min with allophycocyanin (APC) or phycoerythrin (PE)-labeled tetramers. After 40 min of tetramer incubation, surface antibodies were added for 20 min. Cells were fixed in Cytofixation (BD Biosciences). Data were acquired on an LSRII fluorescence-activated cell sorter (FACS) (BD Biosciences) and were analyzed with the software FlowJo (Tree Star, Ashland, OR).

**Synthetic peptides.** Synthetic LCMV peptides D b NP396-404 (FQPQNGQFI), D b GP33-41 (KAVYNFATC), D b GP276-286 (GQVENPGYCL), and K b NP205-212 (YTVKYPNL), as well as PV peptides D b GP33-41 (KAVYNFATC), D b GP276-286 (GQVENPGYCL), and Kb NP205-212 (YTVKYPNL), were purchased from BioExpress Inc. (Kaysville, UT). Three days post-infection with an associated alteration in the T-cell receptor repertoire (Fig. 1d). There was also a 2-fold increase in absolute Treg cell numbers in the draining mLNs (Fig. 1b), with no difference in the nondraining inguinal lymph nodes (iLN) (naïve, 7.56 ± 10^4 to 3.4 ± 10^3; IAV immune, 11 ± 10^4 to 5.2 ± 10^3; P = 0.6; n = 4) and a small increase in the spleens (Fig. 1c). In contrast, infection with another nonpersistent virus, LCMV (Armstrong strain), did not enhance absolute number (naïve, 3.51 ± 10^4 to 1 ± 10^5; LCMV-immune, 2.31 ± 10^4 to 1.2 ± 10^4; P = 0.6; n = 4 to 11) or percentage (naïve, 0.68 ± 0.04%; LCMV-immune, 0.82 ± 0.11%; n = 4 to 12) of Treg cells in the lungs of LCMV-immune mice. These results suggest that some nonpersistent but potentially cytotoxic viruses that induce lung pathology, such as IAV (5, 13, 14), could significantly enhance the number of Treg cells, while other viruses which induce minimal lung pathology, such as LCMV, do not. Interestingly, these increased levels of Treg cells were still detectable long after the acute IAV infection during the memory phase in the lung, the site of the virus infection, and in the draining mLNs.

**IAV-expanded increase in Treg cells persisted during a subsequent heterologous virus infection with LCMV.** After finding increased Treg cells in the lungs and mLNs of IAV-immune mice, we questioned whether these expanded Treg cells participate in dysregulating normal immune responses, leading to either the increased lung pathology or viral titers that had been observed in IAV-immune mice acutely infected with LCMV (5, 15). First, we
determined if this increased number of Treg cells in IAV-immune mice persisted throughout the subsequent LCMV infection (Fig. 1e and 2). In naive mice, all T-cell subpopulations demonstrated a similar pattern of expansion in the mLN during acute LCMV infection, with significant 4- and 3-fold increases in total numbers of CD4+ (P = 0.002; n = 7) and CD8+ (P = 0.05; n = 7) T cells, respectively, by day 3. There also was a 2-fold increase in Treg cells (P = 0.06; n = 7) (Fig. 2a). By day 7, all three T-cell subpopulations had returned to preinfection levels (day 0). However, in IAV-immune mice challenged 6 weeks later with LCMV, Treg cells expanded and contracted in a manner discordant with CD4+ and CD8+ T cells (Fig. 2a, lower). Unexpectedly, the total number of Treg cells remained elevated from day 0 to 9 post-LCMV infection in the mLN of IAV-immune mice (Fig. 2a and b). In fact, a significant, 2-fold higher number of Treg cells was observed on days 7 and 9 after LCMV infection in IAV-immune compared to naive mice (Fig. 2b). At day 7 post-LCMV infection, there was a significant 2-fold increase in the Treg cell/CD4+ Foxp3+ T cell and Treg cell/CD8+ T cell ratios in IAV-immune mice compared to naive mice (Fig. 2c). Since the effect of Treg cells on effector T cells is dependent upon their levels compared to each other, these data suggest that the increased number of Treg cells is able to attenuate the activation of LCMV-specific T-cell responses during their priming in the mLN.

Similar to the mLN, the absolute number of Treg cells in the lung was increased in IAV-immune mice throughout LCMV infection compared to the controls, with significant increases at days 7 and 9 (Fig. 2d). Even in the spleen the absolute number of Treg cells was increased before and during LCMV infection, reaching a significant increase by day 9, the peak of the LCMV CD8+ T-cell response, in IAV-immune mice (Fig. 2e). By using paired t tests for time point and organ (thereby eliminating the variability in cell number that occurs with time after infection and organ), we found that there was a significant overall increase of both the percentage (Fig. 2f) and total number (data not shown) of Treg cells in the lung, mLN, and spleen throughout the LCMV infection from day 0 to 12 of the IAV-immune mouse. This resulted in a decreased ratio of effector cells to Treg cells, both CD8+ (Fig. 2g) and CD4+ (data not shown), during the LCMV infection in IAV-immune mice compared to those of controls throughout the infection in all three organs.

As Treg cells are capable of producing inhibitory cytokines, such as IL-10, we questioned whether there was any evidence of enhanced IL-10 levels during LCMV infection in IAV-immune mice. ELISA multiplex analysis of splenic, mLN, and lung cell lysates demonstrated that IAV-immune mice had significantly higher levels of IL-10 expression than naive mice throughout the LCMV infection (Fig. 2h). Thus, it is possible that the presence of IAV-expanded Treg cells in IAV-immune mice will suppress normal immune effector responses and either decrease LCMV clearance or alter immunopathology.

**PC61 treatment (anti-CD25)** led to decreased lung pathology during acute LCMV infection of IAV-immune mice. In order to determine if the increased levels of Treg cells played a role in mediating immunopathology or altering viral clearance during LCMV infection of IAV-immune mice, PC61 treatment (anti-
CD25), a well-established conventional technique, was used to remove Treg cell activity (Fig. 3a) (16, 32, 33, 37, 38). LCMV infection i.n. in naive mice induced mild interstitial mononuclear infiltrates (Fig. 3c), whereas in IAV-immune mice it induced enhanced bronchus-associated lymphoid tissue (BALT), mononuclear pneumonia consolidations, perivascular and pulmonary edema, bronchiolitis, and bronchiolization (Fig. 3d) (5, 15). Since the presence of Treg cells is frequently associated with decreasing immunopathology, we were somewhat surprised to observe a significant reduction in the severity of lung pathology on days 7 and 9 post-LCMV infection in IAV-immune mice receiving PC61 treatment prior to challenge (Fig. 3e and f). The lung histology in many of the PC61-treated mice resembled that in naive mice infected with LCMV (Fig. 3c), with only mild mononuclear infiltrates. The vast majority of the mice scored below level 3, when bronchiolization, the pathognomonic finding of severe pathology, is first observed. There was no significant difference in weight loss (data not shown) or LCMV titers in the lungs post-LCMV challenge of PC61-treated and nontreated IAV-immune mice, suggesting that the presence of enhanced levels of Treg cells in IAV-immune mice was not altering LCMV clearance (Fig. 3g).

PC61 (anti-CD25) treatment resulted in decreased LCMV-specific CD8⁺ T-cell responses in the spleen and lung of LCMV-infected IAV-immune mice. PC61 treatment prior to LCMV infection in IAV-immune mice led to a significant decrease in the overall total number of LCMV-specific CD8⁺ T cells (NP396- and GP33-specific response) in the spleen (Fig. 4a), associated with lower numbers in the lung, at days 7 and 9 postinfection compared to nontreated mice. In complete contrast, PC61 treatment resulted in a significant enhancement of the total number of LCMV-specific CD8⁺ T cells in the draining mLN on day 7 in IAV-immune mice during LCMV challenge (Fig. 4a). In the nontreated mice, by days 7 and 9 of LCMV infection effector CD8⁺ T cells were decreased in the mLN and might have already trafficked to the lung and spleen, sites of infection (Fig. 4a). Interestingly, there was no effect of PC61 treatment on the number of LCMV GP61-specific CD4⁺ T cells at days 7 and 9 of LCMV infection in the spleen and mLN in IAV-immune mice (Fig. 4b).

These results suggest that there is a delay in effector CD8⁺ T-cell trafficking out of the lymph node in the absence of Treg cells from the LCMV-infected IAV-immune mice, which is consistent with previous reports (39). This delay in effector CD8⁺ T-cell trafficking to the spleen and lung after PC61 treatment may play some role in decreasing lung pathology.

PC61 treatment during acute LCMV infection in naive mice showed no differences in immunopathology and viral titer but did decrease LCMV-specific CD8⁺ T-cell responses. In order to determine if the above-described findings are a phenomenon related to IAV-expanded Treg cells in the lungs and mLN following sequential infection of IAV-immune mice with LCMV or just an

**FIG 3** PC61 (anti-CD25) treatment in IAV-immune mice prior to LCMV infection resulted in significantly decreased lung pathology. (a) Experimental setup of sequential IAV and LCMV infections with PC61 treatment. Three days prior to LCMV infection, IAV-immune mice were treated i.p. with PBS/isotype or PC61 (anti-CD25) (IAV and PC61) and compared to PBS/isotype-treated naive mice. Lung pathology was analyzed at days 7 and 9 post-LCMV infection, and LCMV viral titers were determined at days 7 to 12 postinfection. (b to e) Lung histology of representative mice: naive untreated, uninfected lung (b); LCMV-infected IAV-immune mice (IAV + PC61Tx) with consolidation, enhanced BALT (small arrow), and bronchiolization (large arrow) (d); and Treg-depleted LCMV-infected IAV-immune mice with mild interstitial infiltrates (IAV + PC61Tx + LCMV) (c). (f) Scored lung pathology in naive, IAV-immune, and PC61-treated IAV-immune mice post-LCMV infection (pooled data from days 7 and 9 from 4 independent experiments). (g) LCMV viral titers in lungs at days 7, 9, and 12 post-LCMV infection in LCMV-infected IAV-immune and PC61-treated IAV-immune mice (representative experiment of two similar experiments). Open circle, LCMV-infected naive mice; closed, black circle, LCMV-infected IAV-immune mice; half-filled circle, LCMV-infected, PC61-treated IAV-immune mice. Statistical analysis was done using a two-way ANOVA.
PC61 (anti-CD25) treatment prior to LCMV infection, naive mice were treated with PC61 prior to LCMV infection (Fig. 5a). Unlike the IAV-immune mice, PC61-treated naive mice had lung pathology and virus titers similar to those of control treated mice at days 5, 7, and 9 post-LCMV infection (Fig. 5b and c). However, similar to the results for IAV-immune mice (Fig. 4), PC61-treated naive mice had decreased LCMV-specific CD8+ T-cell responses in the spleen and lung at days 7 and 9 post-LCMV infection compared to control-treated mice (Fig. 5d). They also had increased LCMV-specific CD4+ T cells in the mLNs at day 9. However, while PC61 treatment led to retention of Treg cells in the mLNs, it had no effect on pathology or virus titers during acute LCMV infection of naive mice. This suggests that PC61 treatment in IAV-immune mice where Treg cells are elevated affects CD8+ T cell responses during LCMV infection differently than in naive mice.

PC61 treatment during PV infection in LCMV-immune mice did not alter activation of cross-reactive NP205-specific CD8+ memory T cells. PC61 treatment prior to virus infection in naive mice usually results in increasing the activation and frequency of virus-specific CD8+ T cells (16, 25, 32, 38). However, it is possible that anti-CD25 treatment interferes with the activation of the cross-reactive IAV-specific memory CD8+ T cells responsible for the lung pathology upon LCMV infection. In order to test this hypothesis, we utilized a heterologous virus infection model, LCMV-immune mice infected with PV, where there is no confounding variable, such as immunopathology, and there is a strong activation of cross-reactive NP205-specific memory CD8+ T cells which become immunodominant during the second virus infection (40, 41) (Fig. 6a). PC61 treatment of LCMV-immune mice prior to PV infection resulted in equally effective reactivation of the cross-reactive LCMV-NP205 memory CD8+ T cells, which instead of being very subdominant during PV infection became codominant with the PV-NP38 response (Fig. 6b). Also, the NP205-specific T-cell response became immunodominant in the LCMV-specific memory response (Fig. 6c), as we have previously reported (40). These results suggest that PC61 treatment does not
interfere with reactivation of cross-reactive memory cells, which is consistent with data suggesting that CD25 does not play a major role in activation of memory CD8+ T cells (42).

Partial exhaustion of LCMV-specific CD8+ T-cell responses might contribute to decreasing lung pathology in PC61-treated IAV-immune mice. To further investigate why PC61 treatment during LCMV infection of IAV immune mice resulted in decreased lung pathology, we questioned if decreased Treg cell function induced by PC61 treatment had any effect on the functional efficiency of the LCMV-specific CD8+ T-cell responses. Loss of effector CD8+ T-cell function, called exhaustion, has been described for different persistent viruses (43–45). Exhaustion of CD8+ T cells can be divided into different stages with a gradual loss of T-cell functions in the following order: IL-2 production and proliferation, TNF production and cytotoxicity, and IFN-γ production (45). Partial exhaustion can occur during early periods of chronic infection or in chronic infections when the antigen load is not exceptionally high. Partially exhausted CD8+ T cells are characterized by loss of TNF-α production and reduced proliferation potential (45). Additionally, upregulation of PD-1 and changes in the immunodominance hierarchy of the antigen-specific CD8+ T-cell responses can be found, as some epitope-specific responses are more sensitive to exhaustion (45–49). For instance, NP396-specific CD8+ T-cell responses during chronic LCMV infection are the most sensitive to exhaustion, and this normally immunodominant response becomes subdominant or disappears completely (45). We began to question whether partial exhaustion was occurring in the IAV-immune mice treated with PC61 prior to LCMV infection, when we noticed that in the decreased LCMV-specific CD8+ T-cell response (Fig. 4) there was a disproportionately greater loss of NP396-specific CD8+ T cells (Fig. 7). In fact, upon closer examination, we noted that in these PC61-treated IAV-immune mice infected with LCMV, the NP396-specific TNF-α-producing CD8+ T cells were significantly decreased, in contrast to the GP33-specific CD8+ T cells (Fig. 7 to c). Significantly fewer NP396-specific multicytokine-producing (TNF+ IFN-γ−) to single cytokine-producing (TNF− IFN-γ+) CD8+ T cells were observed in PC61-treated mice (ratio, 5.9 ± 1) compared to nontreated IAV-immune mice (2.5 ± 0.3) (P = 0.025; n = 14) (Fig. 7a). This was also associated with a decrease in the

FIG 5 PC61 treatment (anti-CD25) in naive mice prior to LCMV infection had no effect on lung pathology and viral clearance. (a) Experimental setup of acute LCMV infection with PC61 treatment. To deplete Treg cells in naive mice prior to LCMV infection, PC61 treatment (anti-CD25) was injected i.p. 3 days prior to LCMV infection. As a control, naive age-matched mice were treated with PBS or the isotype control prior to LCMV infection. Lung pathology scoring (b) and viral titers (c) from LCMV-infected naive and PC61-treated naive mice at days 5, 7, and 9 post-LCMV infection are shown. (d) Total number of LCMV-specific T-cell responses (NP396 + GP33) was determined in spleens, lungs, and mLNs at days 7 and 9 post-LCMV infection in PC61-treated or nontreated mice. Naive nontreated (white bar) and PC61-treated (horizontal striped bar) mice post-LCMV infection are shown. Data shown are from 1 to 3 pooled experiments with n = 5 to 15 mice/group. Statistical analysis was done using Student’s t test.

Partial exhaustion of LCMV-specific CD8+ T-cell responses is characterized by loss of TNF-α production and reduced proliferation potential. Additionally, upregulation of PD-1 and changes in the immunodominance hierarchy of the antigen-specific CD8+ T-cell responses can be found, as some epitope-specific responses are more sensitive to exhaustion. For instance, NP396-specific CD8+ T-cell responses during chronic LCMV infection are the most sensitive to exhaustion, and this normally immunodominant response becomes subdominant or disappears completely. We began to question whether partial exhaustion was occurring in the IAV-immune mice treated with PC61 prior to LCMV infection, when we noticed that in the decreased LCMV-specific CD8+ T-cell response there was a disproportionately greater loss of NP396-specific CD8+ T cells. In fact, upon closer examination, we noted that in these PC61-treated IAV-immune mice infected with LCMV, the NP396-specific TNF-α-producing CD8+ T cells were significantly decreased, in contrast to the GP33-specific CD8+ T cells. Significantly fewer NP396-specific multicytokine-producing (TNF+ IFN-γ−) to single cytokine-producing (TNF− IFN-γ+) CD8+ T cells were observed in PC61-treated mice (ratio, 5.9 ± 1) compared to nontreated IAV-immune mice (2.5 ± 0.3) (P = 0.025; n = 14). This was also associated with a decrease in the

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mean fluorescence intensity (MFI) of NP396-specific CD8$^+$ T-cells, making both TNF and IFN-γ levels consistent with decreased cytokine production per cell (Fig. 7a and b).

LCMV-specific CD8$^+$ T-cell responses have a highly predictable hierarchy during acute LCMV infection in naive mice which is not altered in IAV-immune mice (Fig. 7d). During acute LCMV infection, either NP396 or GP33/34 CD8$^+$ T-cell responses may be more dominant, or they can be fully codominant (50). During the phases of T-cell exhaustion, the LCMV NP396-specific CD8$^+$ T-cell response becomes nonfunctional before the GP33/34 response (45). Thus, changes in immunodominance hierarchy during LCMV infection are one way to assess the severity of clonal exhaustion. Thus, as the NP396-specific CD8$^+$ T-cell response was becoming less functional during LCMV infection in PC61-treated IAV-immune mice, the GP33/34-specific CD8$^+$ T-cell response became dominant more often than the NP396-specific response (Fig. 7d). This change in immunodominance hierarchy was unique to PC61-treated IAV-immune mice, as there was no significant difference in hierarchy in PC61-treated naive mice infected with LCMV (Fig. 7e).

Also consistent with ongoing partial CD8$^+$ T-cell exhaustion, significantly enhanced PD-1 expression was observed on NP396-specific CD8$^+$ T cells at days 7 and 9 of LCMV infection in PC61-treated IAV-immune mice compared to nontreated naive or IAV-immune mice (Fig. 7f). This difference in PD-1 expression had disappeared by day 12, when virus had cleared from these mice.

These changes in the characteristics of the LCMV-specific CD8$^+$ T-cell responses (Fig. 7a to f), including alteration in immunodominance hierarchy (Fig. 7d), decreased TNF production (Fig. 7a to c), and enhanced PD-1 expression (Fig. 7f), was not observed in PC61-treated naive mice infected with LCMV. Thus, Treg cell depletion using PC61 treatment did not lead to any evidence of partial clonal exhaustion upon LCMV infection in naive mice (Fig. 7e). Only when the increased numbers of Treg cells present in IAV-immune mice were depleted by PC61-treatment did the LCMV-specific CD8$^+$ T cells demonstrate these features of partial clonal exhaustion. This decreased CD8$^+$ T-cell response would result in decreased IFN-γ production, one of the mediators of severe lung pathology (15). These results suggest that the increased levels of IAV-expanded Treg cells is attenuating LCMV-specific CD8$^+$ T-cell activation and preventing partial T-cell exhaustion in the presence of the higher viral loads in IAV-immune mice (5).

In the absence of functional Treg cells, the ratio of LCMV-GP33/34 to NP396 response directly correlated with the severity of lung pathology. In IAV-immune mice infected with LCMV, the severity of the lung pathology directly correlated with and was predicted by the frequency of IAV-PB1$_{703}$- and IAV-PA224-specific memory responses, which cross-reacted with LCMV-GP33/34 and -GP276, respectively (15). Eradication or functional ablation of these pathogenic memory T-cell populations, using mutant virus strains, peptide-based tolerization strategies, or short-term anti-IFN-γ treatment, inhibited severe lesions, such as bronchiolization, from occurring. In PC61-treated IAV-immune mice, the ratio of GP33/34 to NP396 CD8$^+$ T-cell responses directly correlated with the severity of the residual lung pathology, which suggests that the greater the cross-reactive GP33/34-specific and the lower the non-cross-reactive NP396-specific CD8$^+$

FIG 6 PC61 treatment (anti-CD25) in LCMV-immune mice prior to PV infection does not affect reactivation and immunodominance by cross-reactive NP205-specific memory CD8$^+$ T cells. (a) Experimental setup of sequential LCMV and PV (LCMV+PV) infections with PC61 treatment. Three days prior to PV infection, LCMV-immune (LCMV) mice were treated with PBS/isotype or PC61 (anti-CD25) (LCMV+PC61) and compared to PBS/isotype-treated naive mice. LCMV and PV epitope-specific responses were determined using peptide stimulation in an intracellular cytokine assay at day 8 post-PV infection. (b) Percentage of PV-specific CD8$^+$ T-cell responses (NP38, NP205, and NP122) was determined in spleens at day 8 post-PV infection in PC61-treated or nontreated LCMV-immune mice. (c) Percentage of LCMV-specific CD8$^+$ T-cell responses (NP396, GP33, GP276, or NP205) was determined in spleens at day 8 post-PV infection in PC61-treated or nontreated LCMV-immune mice.
stimulation (gated on CD8^+ T cells) in spleens of LCMV-GP34 CD8^+ T cell responses were determined in naive, IAV-immune, and PC61-treated IAV-immune mice post-LCMV infection (see the experimental set up described in the legend to Fig. 3a). (a to c) IFN-γ and TNF production was measured using an intracellular cytokine assay in response to LCMV-NP396 or GP33 stimulation (gated on CD8^+ T cells) in spleens of nontreated (IAV) and PC61-treated IAV-immune (IAV + PC61Tx) mice at day 7 post-LCMV infection. The partial exhaustion phenotype is demonstrated by FACS plots of two representative mice showing the NP396-specific CD8^+ T-cell response (a), by graphing the MFI of the TNF-α IFN-γ NP396-specific CD8^+ T cell response (b), and by the percentage of TNF-α IFN-γ-producing cells in either group (c). (d and e) The dominant epitope hierarchies of LCMV-specific CD8^+ T-cell responses in spleens at day 7 post-LCMV infection are shown from naive (white bars), IAV-immune (black bars), and PC61-treated IAV-immune mice (diagonally striped bars) (the experimental set-up is described in the legend to Fig. 3a) (d) and LCMV-infected naive (white bars) and PC61-treated naive (horizontally striped bars) mice (n = 14 to 19 mice/group from panels b and c) (the experimental set-up is described in the legend to Fig. 5a) (e). Statistical analysis used the chi-square test. (f) MFI of PD-1 on LCMV NP396-specific CD8^+ CD44^+ IFN-γ^+ T cells of LCMV-infected naive, IAV-immune, and PC61-treated IAV-immune mice at days 7, 9, and 12 post-LCMV infections in the spleen. (For this, data from Fig. 1 to 2 for pooled independent experiments with 5 to 10 mice/group and time point were used; Student’s t-test was used for statistical analysis.) (g and h) Direct correlation of severity of lung pathology and the ratio of LCMV-specific GP33/NP396 responses in the spleen of day 9 LCMV-infected, PC61-treated IAV-immune mice (g) and day 9 LCMV-infected control nontreated IAV-immune mice (h).

T-cell response, the greater the lung pathology (Fig. 7g). This correlation was not observed in control nontreated IAV-immune mice at day 9 post-LCMV infection (Fig. 7h). These data suggest that the partial exhaustion of the NP396-specific CD8^+ T-cell response in PC61-treated IAV-immune mice during LCMV challenge resulted in a greater dependence on the cross-reactive GP33/34-specific response to induce the residual pathology. This is an interesting observation, as the LCMV-GP34 CD8^+ T-cell response is cross-reactive with the IAV-PB1 response, and we have shown that the expansion of memory IAV-PB1-specific CD8^+ T cells during LCMV infection directly correlated with the severity of lung pathology (15). However, it would appear that only when we deplete the confounding influence of the T cells in IAV-immune mice can we observe a direct correlation between cross-reactive LCMV epitope-specific responses and severity of lung pathology. This is another indication that PC61 treatment did not affect the activation and proliferation of cross-reactive memory cells.

**DISCUSSION**

Recently, antigen-specific IAV-induced Treg cells have been reported to attenuate subsequent T cell responses and decrease pathology during a secondary challenge with heterosubtypic IAV infection (51, 52). Here, we show for the first time that virus-expanded Treg cells could attenuate immune responses and influence induction of lung pathology during a subsequent unrelated nonpersistent virus infection. The presence of increased Treg cells during persistent viral infections in naive mice is well documented and is known to influence viral clearance and immunopathology (19, 25–28). Our results suggest that Treg cells generated during an acute infection that clears but where tissue damage occurs can influence the qualitative characteristics of effector T-cell responses and their ability to contribute to lung pathology during a subsequent heterologous virus infection. Thus, an individual’s history of infection and specific sequence of infection can alter Treg cell populations, resulting in greatly altered disease outcome during subsequent new infections.

Whereas Treg cell inactivation with PC61 treatment prior to an acute LCMV infection had no influence on the LCMV-induced immunopathology in the lung, depletion prior to LCMV infection in IAV-immune mice resulted in significantly decreased immunopathology. Increased numbers of Treg cells in IAV-immune mice appeared to attenuate activation of both non-cross-reactive and cross-reactive LCMV-specific CD8^+ T cells, and their presence was detrimental for the host in terms of immunopathology. In the absence of Treg cells there was a delay in effector CD8^+ T-cell trafficking to the lung and spleen, and the non-cross-reactive NP396 CD8^+ T-cell response was so strongly activated that it became partially clonally exhausted, leading to decreased immunopathology. Furthermore, depletion of Treg cells in these sequen-
tially infected mice led to revealing a direct correlation between the ratio of cross-reactive LCMV GP33/34 to non-cross-reactive NP396 CD8+ T-cell responses and the severity of the residual lung pathology. These data suggest that prior shaping of the immune system with unrelated infections is as important as the characteristics of a specific pathogen in terms of influencing Treg cell induction and their beneficial or detrimental effects for the host.

Treg cells were increased in the lung and draining lymph node (mLN) of IAV-immune mice even 6 weeks after resolution of the acute infection. This is consistent with recent reports (51, 52) that IAV can induce IAV-antigen-specific Treg cells, which can directly attenuate the function of IAV-specific effector T cell function, including proliferation and IFN-γ production. Increased Treg cells have been found in chronic virus infections like hepatitis B/C virus, HIV, and Friend virus (19, 26, 28, 53). Punikosy et al. showed that Treg cells expanded depend on viral chronicity (54). Since replicating IAV is cleared within the first 10 days postinfection (55–57), one possibility for the significantly enhanced numbers of Treg cells in the lungs and mLN of IAV-immune mice is the response to self-antigens presented as a result of the damaged lung tissue during acute IAV infection (5). In this case, constant self-specific antigen and T cells would still be around after IAV is cleared. This might explain why we found significantly increased Treg cell numbers in IAV-immune mice even up to several months post-IAV infection. However, it is also possible that Treg cells were maintained at significantly higher numbers due to persistent IAV-derived antigen presentation, which has been described by others (58). Jelley-Gibbs et al. showed that antigen-specific CD4+ T cell responses could be generated up to 3 to 4 weeks post-IAV clearance. These late primed IAV-specific CD4+ T cells have an intermediate phenotype with decreased contraction capacity. Interestingly, these IAV-specific CD4+ T cells are Vβ 4, 7, and 14 (58). Treg cells in the lungs of our IAV-immune mice showed enhanced usage of Vβ 4 and 7 and significantly enhanced Vβ 14 (Fig. 1d). This is an indication that late IAV antigen presentation has an influence on the enhanced IAV-expanded Treg cell numbers in lungs of IAV-immune mice. The results from the recently reported IAV-induced Treg studies suggest that these cells are activated in an antigen-specific manner, raising the question of whether cross-reactive LCMV-specific antigens could activate these IAV-expanded Treg cells. Our unpublished results (M. F. Wlodarczyk, A. R. M. Kraft, L. L. Kenney, E. Carter, and L. K. Selin) suggest that there are cross-reactive CD4+ T-cell responses between LCMV and IAV, much as we have reported cross-reactive CD8+ T-cell responses (15).

The presence of Treg cells is usually associated with decreased immunopathology in models of autoimmune diseases, such as inflammatory bowel or celiac disease (5), and in some pathogen infections, such as respiratory syncytial virus (RSV) (16, 38) and IAV (51, 52), suggesting that depletion of Treg cells in IAV-immune mice resulted in increased lung pathology compared to nondepleted mice or even death after LCMV infection. Depletion of Treg cells by PC61 treatment prior to acute viral infection with, for instance, corneal HSV-1, neonatal HSV-1, or i.n. RSV, significantly enhances severe pathology mediated by increased virus-specific T-cell responses (16, 33, 38). These studies support the concept that the presence of Treg cells results in attenuation of effector T-cell activation. In IAV-immune mice prior to and throughout the subsequent LCMV infection, we found significantly increased numbers of IAV-expanded Treg cells and the suppressive cytokine IL-10, which has been shown to decrease pathology (19, 59, 60) or contribute to induction of chronic infection with high-dose LCMV clone 13 by induction of CD8+ T-cell exhaustion (61–63). Thus, it was possible that depletion of Treg cell function prior to LCMV infection further increased the severity of lung pathology upon LCMV infection in IAV-immune mice. Surprisingly, depletion of Treg cells in IAV-immune mice prior to LCMV infection resulted in decreased lung pathology, with no differences in viral titers and significantly decreased LCMV-specific CD8+ T-cell responses in the spleen but not the mLN. These results suggest that there is a delay in effector T cells trafficking out of the lymph node in the absence of Treg cells in both the LCMV-infected IAV-immune and the naive mice, which is consistent with previous reports demonstrating that Treg cells play a role in controlling egress of effector T cells from the lymph node (39).

However, Treg depletion with PC61 treatment did not alter the severity of immunopathology or viral load in naive mice infected with LCMV. In contrast to findings of Treg depletion via PC61 treatment during acute RSV or IAV infection, depletion of the IAV-expanded Treg cells during LCMV infection did not significantly enhance virus-specific T-cell responses (16, 25, 38). In stark contrast, the significant reduction in the severity of lung pathology appears to be mediated by overactivation and subsequent partial exhaustion of the LCMV-specific CD8+ T-cell response in LCMV-infected, Treg cell-depleted, IAV-immune mice.

High doses of LCMV clone 13 given intravenously leads to chronic infection that is thought to occur because the high antigen load drives LCMV-specific CD8+ T-cell exhaustion (45–48). These antigen-specific cells can be in various phases of partial to complete exhaustion thought to correlate with antigen load (45). Exhaustion includes disruption of predictable immunodominance hierarchies of the LCMV-specific CD8+ T-cell responses (loss of the dominant NP396-specific CD8+ T-cell response), impairment of functional T-cell responses, including loss of TNF-α production, and upregulation of PD-1 expression on exhausted virus-specific T cells (45). All of these parameters of exhaustion were detected on LCMV-specific CD8+ T cells in the spleens of Treg-depleted, IAV-immune mice but not in Treg-depleted, naive mice after LCMV infection.

Why would Treg depletion with PC61 treatment prior to LCMV infection in IAV-immune mice but not Treg depletion in naive mice result in partial exhaustion? There was no evidence to suggest that effector CD4+ T-cell activation and function in the IAV-immune or naive mice infected with LCMV was directly affected by the PC61 treatment. In naive mice infected with LCMV, the only effect that PC61 treatment had was to slightly decrease the size of the LCMV-specific CD8+ T-cell effector response in the spleen while it increased in the mLN, consistent with a delay in egress from the lymph node, a finding consistent with Treg depletion in Foxp3-DTR mice (39). A similar delay in LCMV-specific CD8+ effector T-cell expansion was observed in the Treg-depleted IAV-immune mice. There was no effect on the viral load or immunopathology in Treg-depleted (PC61Tx) naive mice. However, only in the IAV-immune mice did the PC61 treatment result in partial exhaustion of the non-cross-reactive NP396-specific CD8+ T-cell response and decreased immunopathology, as summarized in Table 1. The unique features of the LCMV response in IAV-immune mice are the presence of increased numbers of IAV-expanded Treg cells, increased IL-10 level, weakly activated cross-reactive IAV-specific memory CD8+ T cells with increased levels.
of IFN-γ (15), and greater viral load than that in naive mice (5) (Table 1). This contrasts with sequential infection with Pichinde virus in LCMV-immune mice, where no evidence for LCMV-expanded Treg cells existed (Fig. 1) and where a strong cross-reactive memory CD8+ T-cell response rapidly expanded and clears PV after infection with no significant immunopathology (40). PC61 treatment in these LCMV-immune mice did not affect activation and immunodomination of the cross-reactive NP205-specific CD8+ T-cell response upon PV infection (Fig. 6), suggesting that PC61 treatment did not affect cross-reactive memory cell activation and severity of lung pathology in LCMV-immune mice infected with LCMV.

It is unlikely that the IAV-expanded Treg cells had a significant effect on the activation of the cross-reactive memory T cells but instead were able to attenuate the priming of the naive, non-cross-reactive LCMV-specific CD8+ T-cell responses. Depletion of these activated IAV-expanded Treg cells in this setting would allow a strong activation of these non-cross-reactive naive NP396-specific CD8+ T-cell responses in the presence of activated cross-reactive IAV-specific memory CD8+ T cells and the increased viral load observed in IAV-immune mice (5). This overactivation could lead to partial exhaustion of the LCMV-specific responses, decreased IFN-γ production, and less pathology. This is only partial exhaustion, as the LCMV infection was ultimately cleared and a persistent LCMV infection was not established (45). Based on our studies, it is interesting to speculate about studies done using PC61 treatment (anti-CD25) to prevent rejection following human kidney transplantation or in clinical trials to treat patients with multiple sclerosis (MS) (64). It has been postulated that PC61 treatment was effective, as it led to an impairment of CD8+ T cells by blocking priming or depleting activated CD8+ T cells. Our findings suggest another hypothesis in these cases in which large amounts of donor organ antigen or self-antigen in MS patients is present: PC61 treatment leads to depletion of induced Treg cells, and unwanted antigen-specific CD8+ T-cell responses become exhausted.

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