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Disabling an integral CTL epitope allows suppression of autoimmune diabetes by intranasal proinsulin peptide

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Insulin is a major target of the autoimmune response associated with destruction of pancreatic β cells in type 1 diabetes. A peptide that spans the junction of the insulin B chain and the connecting (C) peptide in proinsulin has been reported to stimulate T cells from humans at risk for type 1 diabetes and autoimmune diabetes–prone NOD mice. Here we show that proinsulin B24–C36 peptide binds to I-A^q, the MHC class II molecule of the NOD mouse, and, after intranasal administration, induces regulatory CD4+ T cells that, in the absence of CD8+ T cells, block the adoptive transfer of diabetes. Curiously, however, intranasal B24–C36 did not inhibit development of spontaneous diabetes in treated mice. We then determined that B24–C36, and its core sequence B25–C34, bind to K^d, the NOD mouse MHC class I molecule, and elicit CD8+ CTLs. When the CD8+ T lymphocyte epitope was truncated at the C34 valine anchor residue for binding to K^d, the residual CD4+ T cell epitope, B24–C32/33, significantly inhibited diabetes development after a single intranasal dose. This study identifies a novel CTL epitope in proinsulin and demonstrates that the therapeutic potential of a “tolerogenic” autoantigen peptide can be compromised by the presence of an integral CTL epitope.


Introduction

In type 1 diabetes (T1D), T cells target and destroy insulin-producing β cells in the islets of the pancreas. Several lines of evidence indicate that proinsulin plays a key role as an autoantigen in driving this process. For example, susceptibility of humans to T1D maps to a variable number of tandem repeats 5’ of the insulin gene, the length of which correlates with thymic expression of proinsulin mRNA and inversely with disease risk (1, 2). In the NOD mouse, a model of spontaneous autoimmune diabetes, transgenic expression of proinsulin in APCs prevents mononuclear cell infiltration of the islets (insulitis) and diabetes (3). A peptide in human proinsulin that spans the cleavage site between the B chain of insulin and the connecting (C) peptide in proinsulin, B24–C36, was shown to stimulate peripheral blood T cells from humans at risk for T1D (4). Recently, a peptide spanning the B–C chain junction in mouse proinsulin II was shown to contain a naturally processed epitope for autoreactive T cells in young NOD mice (5).

Peptide autoepitopes are candidate therapeutic agents for preventing autoimmune disease (6). Mucosal administration of a peptide antigen can suppress subsequent systemic immune priming by the antigen (mucosal tolerance) and has been used to inhibit a range of experimental autoimmune diseases (6–8). We therefore administered proinsulin B24–C36 peptide intranasally to NOD mice in the expectation that it would inhibit the development of diabetes. However, although intranasal B24–C36 elicited antidiabetogenic regulatory T cells, it did not protect mice from diabetes. In exploring the reason for this, we determined that B24–C36, and its core sequence B25–C34, bind to K^d, the NOD mouse MHC class I molecule, and elicit CD8+ CTLs. When the CD8+ T lymphocyte epitope was truncated at the C34 valine anchor residue for binding to K^d, the residual CD4+ T cell epitope, B24–C32/33, significantly inhibited diabetes development after a single intranasal dose. This study identifies a novel CTL epitope in proinsulin and demonstrates that the therapeutic potential of a “tolerogenic” autoantigen peptide can be compromised by the presence of an integral CTL epitope.

Methods

Mice. NOD mice (Lt/Jax) were bred under specific pathogen-free conditions at The Walter and Eliza Hall Institute of Medical Research. Diabetes was diagnosed if two sequential measurements of retro-orbital venous blood glucose exceeded 11 mM.

Protein and peptides. Mouse proinsulin II peptides B24–C36 (FFYTPMSRRREVED), B24–C32, B24–C33,
B25–C33, B26–C34, and B25–C34, hen-egg lysozyme (HEL) 10–23, and *Listeria monocytogenes* listeriolysin O (LLO) 91–99 (GYKDGNEYI) were synthesized and purified to greater than 95% homogeneity, determined by HPLC-MS, by Mimotopes (Melbourne, Australia). OVA (Sigma-Aldrich, St. Louis, Missouri, USA) and peptides were resuspended in PBS.

**Intranasal administration of peptides.** In a series of preliminary experiments, unanesthetized female NOD mice (12 per group) were given either carrier PBS or 0.4, 4, 40, or 80 µg of mouse proinsulin B24–C36 in 10 µl PBS intranasally on three alternating days from 8 weeks of age, and their incidence of diabetes was monitored out to 26 weeks of age. The 0.4 and 4 µg of mouse proinsulin had no effect, whereas both the 40 and 80 µg of mouse proinsulin had a small (11%–24%) effect to decrease diabetes incidence. Subsequently, 40 µg of test or control peptide was given on three alternating days. In diabetes incidence studies, blood glucose was measured every 4 weeks from 100 days of age.

**Adoptive transfer of diabetes.** Male 6- to 9-week-old NOD mice (8 or 12 per group) were irradiated (8 Gy) from a cobalt source and 3–6 hours later received 107 pooled diabetogenic splenocytes from recently NOD female mice, together with 107 splenocytes or cells fractionated from this number from either proinsulin peptide- or OVA-treated mice, in 200 µl via the tail vein. The onset of diabetes was monitored every 2 weeks by measuring blood glucose beginning 2 weeks after transfer.

**Fractionation of splen cells.** Splenic CD4+ and CD8+ cells were either selected or depleted by magnetic separation with monoclonal antibodies bound to MACS microbeads (Miltenyi Biotec GmbH, Bergisch Gladbach, Germany). The yield of CD4+ and CD8+ cells was 70%–80% and 20%–30%, respectively, and their depletion or purity by flow cytometry more than 95% and more than 85%, respectively.

**I-A\(^{\beta}\) binding assay.** Binding of peptides to purified, soluble I-A\(^{\beta}\) was measured in an ELISA as previously described (9). Competition dose-response binding curves were generated, and peptide binding affinity for I-A\(^{\beta}\) was expressed as an IC\(_{50}\); the molar concentration of peptide that inhibited by 50% binding of the indicator peptide, N-terminal biotinylated HEL 10–23.

**Prediction of peptides that bind to MHC class I.** Peptides within mouse proinsulin B24–C36 that could potentially bind to the NOD mouse MHC class I molecules, K\(^{\beta}\) and D\(^{\beta}\), were identified in the Web-based databases SYFPEITHI (10) and BIMAS (11).

**Modeling of peptide binding to MHC molecules.** When homology modeling was performed, no crystal structures of K\(^{\beta}\) or I-A\(^{\beta}\) were available. The two crystal structures of I-A\(^{\beta}\) published subsequently (12, 13) agree in nearly all respects with the modeled structure used here (14). Modeling of I-A\(^{\beta}\) was based on the coordinates of the I-A\(^{\beta}\) /HEL 50-62 peptide complex (15) and was performed with the program Discover II of Accelrys (San Diego, California, USA). Modeling of K\(^{\beta}\) was based on the coordinates of HLA-A2 hepatitis B nucleocapsid 18–27 peptide complex (16) (Protein Data Bank access code: 1hhh.pdb [ref. 15]). We found that HLA-A2 is a better base molecule for modeling K\(^{\beta}\) than the more homologous mouse K\(^{\beta}\), giving better agreement with binding and T cell proliferation data for variants of the insulin B15–23 CTL epitope (17). The modeling procedure for K\(^{\beta}\) was identical to that for I-A\(^{\beta}\), at an ambient pH of 7.0. The models are shown in solid-surface representation with colors according to the surface electrostatic potential (gray, neutral; blue, positive; red, negative). The peptide residues are in van der Waals space-filling form, whereas selected MHC heavy chain residues shown are in stick representation (oxygen, red; nitrogen, blue; carbon, green; hydrogen, white). Secondary structure elements of the MHC molecule are also shown for orientation purposes: \(\alpha\)-helix in red, \(\beta\)-pleated sheet in turquoise, and random coil in gray. A transparency function has been included to allow secondary structural elements of the MHC molecule and the peptide residues buried in it to be slightly visible.

**K\(^{\beta}\)-peptide binding stabilization assay.** K\(^{\beta}\)-transfected RMA-S cells (18), 10^4 cells per well, were incubated overnight at 27°C in RPMI-1640 medium with 100 µM test peptide, heat-shocked for 2 minutes at 56°C, washed, and incubated with anti-K\(^{\beta}\) antibody (34-1-2S) for 30 minutes at 4°C, followed by FITC-conjugated sheep anti-mouse immunoglobulin (Silenus Laboratories, Victoria, Australia). LLO 91–99, known to bind K\(^{\beta}\) with high affinity (19), was used as a reference peptide. Cells were examined with a FACScan flow cytometer (Becton Dickinson Immunocytometry Systems, San Jose, California, USA) that used CellQuest software. Binding of peptide to K\(^{\beta}\) is observed as a temperature-dependent stabilization of surface K\(^{\beta}\) expression.

![Figure 1](image-url)

**Figure 1** Intranasal proinsulin B24–C36 induces CD4+ T cells that block adoptive transfer of diabetes. Female 8-week-old NOD mice \((n = 4 \text{ per group})\) were given 40 µg of proinsulin B24–C36 peptide or OVA protein in 10 µl PBS, intranasally on three alternating days. Two weeks after the last treatment, the mice were killed and 10^7 pooled splenocytes from mice treated with proinsulin B24–C36 (filled circles) or OVA (open circles), or 10^6 pooled splenocytes from mice treated with proinsulin B24–C36 either selected for CD4+ cells (filled triangles) or depleted of either CD4+ (filled inverted triangles) or CD8+ (filled squares) cells, were cotransferred with 10^7 pooled splenocytes from recently NOD female mice into irradiated young NOD male recipients \((n = 8 \text{ or } 12 \text{ per group})\). The development of diabetes in recipients after transfer was monitored by measuring blood glucose every 2 weeks. Results are representative of five similar experiments.
Chromium $[^{51}Cr]$-release CTL assay. Effector cells were generated in female NOD mice by subcutaneous injection of 50 µg peptide in CFA (Difco Laboratories, Detroit, Michigan, USA). After 14 days, splenocytes were harvested and restimulated in vitro for 6 days in HEPES Eagle’s Medium (HEM)/2.5% FCS at 37°C with peptide-coated (10 µg/ml) irradiated (15 Gy) splenocytes (20). Restimulated effector cells (100 µl) were then seeded in duplicate with $^{51}$Cr-labeled RMA-S-Kd target cells (100 µl) pulsed with 100 µM peptide. RMA-S-Kd targets were prepared by incubating $1 \times 10^6$ cells with 100 µM peptide and 100 µCi $^{51}$Cr in 200 µl for 2 hours at 27°C. Plates were incubated for 6 hours at 37°C in 5% CO$_2$/air. Percent specific lysis (cpm in 100 µl supernatant) was determined as $(\text{experimental release} - \text{medium release})/(\text{maximum release} - \text{medium release}) \times 100$.

Statistical analysis. Group differences were analyzed with Fisher exact test (two-tailed) and differences between Kaplan-Meier survival curves were analyzed by the log-rank test, with the use of GraphPad Prism version 3.0a for Macintosh (GraphPad Software Inc., San Diego, California, USA).

Results

Intranasal proinsulin B24–C36 induces CD4$^+$ regulatory T cells but fails to suppress spontaneous diabetes. Female 8-week-old NOD mice were treated with three intranasal doses of proinsulin B24–C36 peptide. Two weeks later, their unfractionated splenocytes or splenocytes from which CD4$^+$ or CD8$^+$ cells had been depleted or selected, were cotransferred with splenocytes from other recently diabetic NOD mice into young irradiated NOD males. Cotransfer of splenocytes from NOD mice given intranasal proinsulin peptide B24–C36 with diabeticogenic splenocytes modestly decreased the incidence of diabetes in recipients ($P = 0.05$) (Figure 1). Diabetes incidence was significantly reduced by cotransfer of either purified splenic CD4$^+$ cells or splenocytes depleted of CD8$^+$ cells from mice given intranasal proinsulin B24–C36 ($P = 0.008$ for both compared with splenocytes depleted of CD4$^+$ cells) (Figure 1). Intriguingly, however, intranasal proinsulin peptide given in a single dose, or in three doses either on alternating days from 8 weeks of age or monthly from 4 weeks of age, had only a small effect to suppress spontaneous diabetes development in female NOD mice monitored for 26 weeks or longer (data not shown). These results suggested to us that intranasal B24–C36 could have induced not only regulatory CD4$^+$ but also pathogenic CD8$^+$ T cells, with a net neutral effect of the whole splenocyte population on disease transfer.

Proinsulin B24–C36 comprises overlapping MHC class II- and class I-restricted epitopes. To determine whether proinsulin B24–C36 comprised epitopes for both CD4$^+$ and CD8$^+$ T cells, we first analyzed B24–C6 for MHC class II- and class I-binding peptides. B24–C36 and the shorter peptide B23–C33 bound to I-A$^b$, the MHC class II molecule of NOD mice, with relatively high affinity (IC$_{50} = 3–5$ µM) compared with the reference HEL 10–23 peptide (Figure 2a). Conversely, proinsulin B22–C31 failed to bind to I-A$^b$. Homology modeling (Figure 2b) revealed that the core nonamer sequence accommodated in I-A$^b$ was B24–C32 (FFYTPMSRR). However, the three acidic residues beyond position (p) 9 in B24–C36, that is, p10E (C33), p12E (C35), and p13D (C36), interact with the side chain carboxylate (and secondarily the terminal carboxylate) of p13D (C36), while α76R interacts with p10E (C33) and p12E (C35), and

![Figure 2](https://example.com/figure2.png)

(a) Binding of proinsulin B24–C36 peptides to purified, soluble NOD mouse MHC class II, I-A$^b$. Competition between biotinylated HEL 10–23 peptide and unlabeled HEL 10–23 (open squares), proinsulin B24–C36 (filled circles), B23–C33 (open triangles), or B22–C31 (filled diamonds) for binding to I-A$^b$ measured by ELISA. (b) Model of B24–C36 bound to I-A$^b$ at pH 7.0, viewed from the perspective of the T cell receptor (TCR). Anchor residues at position 1F (p1F) (B24) and p9R (C32) point into the groove of I-A$^b$ and are only partially visible. The p4T points into the groove at pocket 4 and is partially visible, as is p6M. By contrast, residues at p2F, p3Y, p5P, p7S, and p8R point upward and are accessible by the TCR. Note the interaction of α75K and α76R with the three acidic residues at the C-terminus of the peptide as shown in the main figure and in the bottom right-hand corner inset. The latter is generated by rotating this segment of the main figure by 40 degrees along the x axis (bottom up, top into the paper) and 20 degrees along the y axis (right up, left into the paper). α75K interacts with the side chain carboxylate (and secondarily the terminal carboxylate) of p13D (C36), while α76R interacts with p10E (C33) and p12E (C35).
p13D (C36), have the capacity to greatly enhance binding by forming salt bridges with I-A^\* chain 76R (C33, C35) and 75K (C36). There is also an interaction between R (C32) and E (C33), probably accommodated because of the wider opening of the p9 pocket in I-A^\* compared with other MHC class II alleles (12–14).

To identify peptides in B24–C36 that might bind to NOD MHC class I molecules (K^d and D^b), we initially scanned the Web-based peptide motif databases SYFPEITHI (10) and BIMAS (11). Three overlapping nonamers, B24–C32, B25–C33, and B26–C34, and one decamer, B25–C34, were identified as potential binders to K^d. Homology modeling (Figure 3a) revealed that the core binding sequence was likely to be the decamer, B25–C34, with anchor residues for K^d binding at p2 (Y) and p10 (V). To confirm this prediction, we performed direct binding studies of synthetic peptides to cell surface K^d, using the TAP-deficient cell line RMA-S-K^d. This treatment delayed diabetes onset and reduced the incidence of diabetes to 30% at 280 days, compared with 70% in mice similarly treated with either B24–C32 or HEL 10–23 (P = 0.03).

The K^d-binding B26–C34 and B25–C34 peptides were then tested to determine whether they could prime K^d-restricted CTLs. Female NOD mice were immunized by subcutaneous injection of 50 \(\mu\)g of peptide in CFA. Spleen cells were recovered after 14 days and restimulated with peptide in culture before assay against \(^{51}\text{Cr}\)-labeled RMS-K^d cells prepped with peptide. In accordance with their ability to bind K^d, B25–C34 and, to a lesser extent, B26–C34 elicited CTLs capable of lysing target RMS-S cells and releasing \(^{51}\text{Cr}\) (Figure 4). It is noteworthy that B24–C36 could prime CTLs able to recognize not only B24–C36 itself, but also B25–C34. Priming with the non–K^d-binding peptides, B24–C32, B24–C33, and B25–C33, failed to induce CTLs (data not shown). CTLs were consistently demonstrated after subcutaneous immunization with B24–C36 in adjuvant but could not be reliably detected after intranasal B24–C36.

**Treatment with the K^d-restricted CTL epitope, proinsulin B25–C34, reduces diabetes incidence.** Systemic administration of autoepitope peptides with or without incomplete Freund’s adjuvant (IFA) has been reported to trigger antigen-induced cell death and delete autoantigen-specific T cells (21–24). To obtain evidence that B25–C34-specific CTLs may be involved in \(\beta\) cell destruction, 50 \(\mu\)g of B25–C34 in IFA was administered intraperitoneally to female NOD mice (n = 10) at 18 days of age, before the onset of insulitis. This treatment delayed diabetes onset and reduced the incidence of diabetes to 30% at 280 days, compared with 70% in mice similarly treated with either B24–C32 or HEL 10–23 (P = 0.03).

**Disabling the B25–C34 CTL epitope permits diabetes prevention by intranasal peptide.** To determine whether the failure of intranasal B24–C36 to prevent spontaneous diabetes, despite induction of regulatory CD4^+ T cells, was due to the B25–C34 CTL epitope, B24–C36 was truncated at the C-terminus to eliminate the C34 valine anchor residue for binding to K^d. To confirm that intranasal administration of the truncated B24–C34 peptide could induce regulatory CD4^+ T cells, 40 \(\mu\)g of the peptide was administered intranasally to NOD mice on three alternating days from 8 weeks of age. Two weeks later, 10^5 splenocytes, 10^6 purified splenic CD4^+ cells, or 10^5 splenocytes depleted of CD4^+ cells, from treated mice, were cotransferred intravenously

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**Figure 3**

Proinsulin B24–C36 peptides bind to H-2K^d. (a) Model of proinsulin B25–C34 bound to H-2K^d at pH 7.0, viewed from the perspective of the TCR. Anchor residues at p2Y and p10V point into the groove of K^d and are only partially visible. The p3T points into the groove at pocket D and is not visible. The p7R points into the groove at pocket E and is partly visible. By contrast, residues at p1F, p4P, p5M, p6S, p8R, and p9E point upward and are thus accessible by the TCR. Note the tight packing of p1F into pocket A and against 167W, as well as the interaction of these aromatic residues with 171Y. Residue 152D is below p8R and forms a salt bridge with it (not shown), while 155Y has its aromatic plane nearly parallel to the plane of the paper. (b) Proinsulin B26–C34, B24–C36, and B25–C34 bind to H-2K^d. TAP-deficient RMS-K^d cells were incubated in medium alone (no peptide) or in the presence of the indicated proinsulin peptides, or reference LLO peptide, and the level of surface K^d expression was then measured by flow cytometry. Cells incubated in media alone were also stained with an isotype control antibody (dotted line).

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with 10^7 splenocytes from recently NOD females into irradiated 6-week-old NOD males (n = 12 per group). Splenic CD4+ cells from mice that had received intranasal B24–C33 prevented transfer of diabetes: 6 weeks after cotransfer, only one of eight recipients of whole splenocytes and none of the eight recipients of purified splenic CD4+ cells from B24–C33–treated mice were diabetic, in contrast to seven of the eight recipients of splenocytes depleted of CD4+ T cells (P = 0.001).

To determine the effect of intranasal B24–C33 and the core I-A^d–binding nonamer sequence, B24–C32, on spontaneous diabetes incidence, female NOD mice (n = 12 per group) were given a single 40-µg dose of each peptide, the parent B24–C36 peptide, or PBS carrier only at 8 weeks of age. B24–C36 had no effect on diabetes incidence: at 182 days of age 83% of mice had become diabetic compared with 92% of PBS controls. On the other hand, after treatment with B24–C33 or B24–C32, the incidence of diabetes (log-rank survival analysis) was significantly reduced, being 33% (P = 0.02) and 25% (P = 0.01), respectively, at 154 days and 58% (P = 0.03) and 25% (P = 0.004), respectively, at 184 days (Figure 5).

Discussion

A sequence in proinsulin, spanning the cleavage site between the B chain of insulin and the C-peptide of proinsulin, is recognized by T cells from humans at risk for T1D (4) and NOD mice (5). Here we show in the NOD mouse that proinsulin B24–C36 comprises overlapping CD4+ and CD8+ T cell epitopes. B25–C34 is a novel MHC class I–restricted (K^d-restricted) CTL epitope that overlaps the core MHC class II–binding (I-A^d–binding) nonamer, B24–C32, and limits the ability of the latter to prevent diabetes after intranasal administration. Although the B24–C36 peptide induced regulatory CD4+ T cells after intranasal administration, it did not alter the incidence of spontaneous diabetes. The CTL epitope was disabled by truncation of B24–C36 at its C-terminus to eliminate the C34 valine, an anchor residue for binding into the p9 pocket of K^d. Without the C34 valine, the peptide cannot bind K^d, as demonstrated. However, it can still bind to I-A^d and, when administered intranasally, not only induced regulatory T cells but suppressed the development of diabetes. Interestingly, CTLs could be elicited not only by B25–C34 but also by the B24–C36 peptide, which also bound to K^d. Although MHC class I molecules are generally considered to accommodate shorter nonamer or decamer peptides, the C-terminal binding pocket of some MHC class I molecules can allow C-terminal peptide extensions out of the binding cleft (25). The only other CTL epitope reported in the NOD mouse, insulin B chain 15–23 (26), also overlaps a CD4+ T cell epitope within the B chain 9–23 sequence. Contiguous or overlapping CD4+ and CD8+ T cell epitopes have been found as well in the two other islet autoantigens, tyrosine phosphatase–like insulinoma antigen-2 (27, 28) and glutamic acid decarboxylase (29). These findings suggest that a single combitope sequence may be processed by an APC and serve to efficiently elicit both helper CD4+ T cell and associated CD8+ CTL responses.

Although we detected B25–C34–specific CTLs in the NOD mouse spleen after subcutaneous priming, we were unable to detect their priming by intranasal B24–C36 administration. This may be due to the limited sensitivity of the ^51Cr-release assay. We have used this assay to detect OVA-specific CTLs after low-dose intranasal OVA (30), but because OVA is a non-self-protein, it would be recognized by a higher frequency
of CTL precursors. Even so, when CTLs are primed by mucosal OVA, they exhibit lower lytic activity than when they are primed subcutaneously in adjuvant (20). It is also relevant that the CTLs generated by subcutaneous priming with B24–C36 in adjuvant exhibited lower lytic activity than CTLs primed to non–self-OVA or LLO peptides. Thus, a relatively low frequency of CTLs or a relatively low avidity of CTLs for B25–C34 self-peptide is likely to mitigate against the detection of these CTLs ex vivo after intranasal treatment.

Two findings implicate the B25–C34 CTL epitope in the pathogenesis of NOD mouse diabetes. First, disabling the CTL epitope by truncation of the C-terminal anchor residue at C34 uncovered the ability of the N-terminal CD4+ T cell epitope, B24–C32/33, to prevent diabetes after intranasal administration. Second, when B25–C34 was administered systemically to young mice, before the onset of insulinitis, diabetes development was suppressed. Immunoprotection after systemic high-dose administration of a CTL epitope has previously been reported in a transgenic T cell receptor (TCR) model of autoimmune diabetes (24). This effect of B25–C34 administration just before the appearance of insulinitis is consistent with evidence that direct β cell recognition by MHC class I–restricted CD8+ T cells is a requirement for the onset of insulinitis (31).

Mucosal administration of autoantigen often confers only partial protection from experimental autoimmune disease and in some cases has been shown to exacerbate disease (32–34). In addition, trials in humans of oral myelin basic protein for multiple sclerosis (35), oral collagen type II for rheumatoid arthritis exacerbate disease (32–34). In addition, trials in autoimmune disease and in some cases has been shown to be mediated by CTLs or a relatively low avidity of CTLs for B25–C34 self-peptide. This may be achieved by mutat- ing or deleting CTL epitopes, as shown here, or by blocking costimulation-dependent CTL activation as shown previously in a transgenic OVA model (39).

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