Modification of Cysteine Residues In Vitro and In Vivo Affects the Immunogenicity and Antigenicity of Major Histocompatibility Complex Class I-restricted Viral Determinants

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Summary

In studying the subdominant status of two cysteine-containing influenza virus nuclear protein (NP) determinants (NP<sub>39–47</sub> and NP<sub>218–226</sub>) restricted by H-2K<sup>d</sup>, we found that the antigenicity of synthetic peptides was enhanced 10–100-fold by treatment with reducing agents, despite the fact that the affinity for K<sup>d</sup> was not enhanced. Reducing agents also markedly enhanced the immunogenicity of cysteine-containing peptides, as measured by propagation of long-term T cell lines in vitro. Similar enhancing effects were obtained by substituting cysteine with alanine or serine in the synthetic peptides, demonstrating that sulfhydryl modification of cysteine is responsible for the impaired antigenicity and immunogenicity of NP<sub>39–47</sub> and NP<sub>218–226</sub>. We found similar effects for two widely studied, cysteine-containing peptides from lymphocytic choriomeningitis virus. The major modifications of cysteine-containing synthetic peptides are cysteinylation and dimerization occurring through cysteine residues. We demonstrate that both of these modifications occur in cells synthesizing a cytosolic NP<sub>218–226</sub> minigene product, and, further, that T cells specific for cysteinylated NP<sub>218–226</sub> are induced by influenza virus infection in mice, demonstrating that this modification occurs in vivo. These findings demonstrate that posttranslational modifications affect the immunogenicity and antigenicity of cysteine-containing viral peptides and that this must be considered in studying the status of such peptides in immunodominance hierarchies.

Key words: antigen processing • cysteine/immunology • major histocompatibility complex/immunology • viral vaccines/immunology • peptides/immunology

C<sup>D8+</sup>T cells (T<sub>CD8+</sub>) recognize peptides, usually of 8–10 residues, complexed to MHC class I molecules. The peptide binding site is formed by class I heavy chains, which are encoded by genes that are highly polymorphic in most species. Most of the variability between heavy chain alleles resides in residues located in the peptide binding site. As a result, each class I allele presents a unique spectrum of peptides to the immune system (1). Although the rules that govern peptide binding to different alleles are far from simple, most peptides recovered from a given allele conform to a simple motif dictated largely by the nature of two to three "pockets" present at the bottom of the binding groove (2). Using these motifs, it is possible to identify upwards of 90% of peptides in a given sequence that could potentially bind to a given allele with biologically significant affinity. Synthetic peptides corresponding to these sequences can be tested for immunogenicity, antigenicity, and binding to class I molecules.

Although this method is capable of identifying antigenic regions of proteins, it remains a considerable challenge to establish the precise structure of the naturally processed determinant. As a first step, the chromatographic properties of synthetic and naturally processed peptides recovered from class I molecules can be compared by HPLC. Coelution indicates a close structural relationship, but the exact structure of the naturally processed peptide can only be established by mass spectroscopy (3). This is sufficiently difficult to preclude publication of mass spectroscopic analysis of viral peptides to date.

The possibilities for surprises in the structure of naturally processed peptides have been amply documented by Meadows et al. (4), who have identified peptides with a number of posttranslational modifications. Of direct relevance to the present study, naturally processed peptides with cysteinylated cysteine residue were identified, as were T<sub>CD8+</sub> that required this modification for activation (4).
Due to its free SH group, cysteine is the most chemically reactive of the 20 common amino acids under physiological conditions. Cysteine is the second least frequent residue in proteins, representing 1.7% of the residues present in eukaryotic proteins. Its occurrence in antigenic peptides is consistent with random usage, being present in 14% of 384 class I ligands (1) (predicted frequency in nonamers is 14%). The frequency of cysteine-containing determinants is expected to be even higher in cysteine-rich peptides, such as Her2/Neu, a promising target for tumor immunotherapy (5). In this study, we show that modification of cysteine-containing determinants by the Biologic Resource Branch, National Institute of Allergy and Infectious Diseases. Peptides were dissolved at 1 mM in DMSO and stored at −20°C unless otherwise indicated. For analytical purposes, peptide masses were determined by matrix-assisted laser desorption ionization with time of flight detection (MALDI-TOF) using a Hewlett Packard mass spectrometer (model G2025A) and cyano-4-hydroxycinnamic acid as the matrix. Peptide binding to live cells was determined by protection of class I molecules to melting (6). In brief, T2-Kd cells were cultured for 14−16 h at 26°C. Synthetic peptides, diluted in FBS-free DMEM in the presence or absence of tris (2-carboxyethyl) phosphine hydrochloride (TCEP; Pierce Chemical Co.) as indicated, were added to cells, which were incubated at 37°C for 2 h to denature Kd molecules not stabilized by peptide binding. Cells were then washed and stained with fluorescent-conjugated SFI-1.1 (PharMingen). Live cells were gated based on scattering and exclusion of ethidium homodimer (Molecular Probes, Inc.) present at 5 µg/ml for 5 min before the last wash. For each histogram, 10,000−20,000 cells were counted on a Becton Dickinson FACScan™, and live cells were analyzed using CELLQuest™ software (Becton Dickinson).

**Materials and Methods**

**Cell Culture.** All media, including cystine-containing and cystine-methionine-free DMEM, were purchased from Biofluids, Inc. The mastocytoma cell line P815 (H-2d) and the thymoma cell line EL-4 (H-2d) were maintained in DMEM containing 10% fetal bovine serum (FBS), 15 × 10−6 M β-ME, antibiotics, and 2 mM glutamine (DE-ME-10). Kd−transfected T2 (T2-Kd) and RMA-S (RMA-S/Kd) cells were cultured in RPMI 1640 containing 10% FBS and the above supplements (R-P-10). CTL stimulation and maintenance were performed in R-P-10 medium containing 10 U/ml of recombinant human IL-2. In some assays, cells were incubated with IMDM supplemented with 10% FBS (I-10). CTL Priming in Vivo and A Assay. 8−10-wk-old female BALB/c mice or C57BL/6 mice were injected intraperitoneally with 1 ml of a 1:10 dilution of chicken egg alantoic fluid containing influenza A virus Puerto Rico/8/34 (PR8) or intravenously with 106 PFU of lymphocytic choriomeningitis virus (LCMV) WE strain. Splenocytes were stimulated with peptide-pulsed APCs for 7 d in vitro at least 3 wk after virus priming. Cells were stimulated in R-P-10 with 10 U/ml IL-2 in 6-well plates. In brief, 3 × 104 splenocytes were stimulated with 6 × 106 virus-infected or peptide-pulsed (1 nM) APC, which were irradiated with 200 Gy before addition to cultures. After 4 d, live cells were recovered via a Ficoll–Hypaque gradient and recultured with fresh IL-2-containing medium. CTL activities were tested after 7 d in standard Ficoll–Hypaque gradient and recultured with fresh IL-2–containing medium. CTL staining and maintenance were performed in R-P-10 medium containing 10 U/ml of recombinant human IL-2. In some assays, cells were incubated with IMDM supplemented with 10% FBS (I-10).

**Cysteine Modification of Viral Peptides**

In the present study, we show that modification of cysteine-containing determinants such as Her2/Neu, a promising target for tumor immunotherapy (5). The frequency of cysteine-containing determinants is expected to be even higher in cysteine-rich proteins, such as Her2/Neu, a promising target for tumor immunotherapy (5). In this study, we show that modification of cysteine-containing determinants by the Biologic Resource Branch, National Institute of Allergy and Infectious Diseases. Peptides were dissolved at 1 mM in DMSO and stored at −20°C unless otherwise indicated. For analytical purposes, peptide masses were determined by matrix-assisted laser desorption ionization with time of flight detection (MALDI-TOF) using a Hewlett Packard mass spectrometer (model G2025A) and cyano-4-hydroxycinnamic acid as the matrix. Peptide binding to live cells was determined by protection of class I molecules to melting (6). In brief, T2-Kd cells were cultured for 14−16 h at 26°C. Synthetic peptides, diluted in FBS-free DMEM in the presence or absence of tris (2-carboxyethyl) phosphine hydrochloride (TCEP; Pierce Chemical Co.) as indicated, were added to cells, which were incubated at 37°C for 2 h to denature Kd molecules not stabilized by peptide binding. Cells were then washed and stained with fluorescent-conjugated SFI-1.1 (PharMingen). Live cells were gated based on scattering and exclusion of ethidium homodimer (Molecular Probes, Inc.) present at 5 µg/ml for 5 min before the last wash. For each histogram, 10,000−20,000 cells were counted on a Becton Dickinson FACScan™, and live cells were analyzed using CELLQuest™ software (Becton Dickinson).

**Extraction of Cellular Peptides and Fractionation by Reversed-phase HPLC (R-P-HPLC).** Natural peptides were recovered and analyzed as previously described (7). In brief, cultures of P815 cells were expanded in roller bottles. 5 × 10−106 cells were infected as described above and incubated for 6 h at 37°C before being pelleted, washed twice in PBS, lysed with ice-cold 0.33% TFA/ H2O, and further disrupted using a Tenbroek tissue homogenizer on ice. At this stage, synthetic peptides were added to uninfected P815 cells as control. Lysates were sonicated and centrifuged at 10,000 g for 30 min, and the supernatants were passed through a 3K cutoff filter (Millipore™) and fractionated on a C8 column (Deltapack; Waters) at 1 ml/min on TFA/acetonitrile gradient (7). Either 0.25- or 1-ml fractions were collected.

**Microcystotoxicity Assay.** Generally, 104 target cells were labeled with 100 µCi of Na125I (Dupont) in minimum volume of medium at 37°C for 60 min. For some experiments, RMA-S/Kd cells that had been cultured for 12–14 h at 26°C were labeled at 26°C for the same time period. After two washes, 104 cells were aliquoted into round-bottom, 96-well plates containing serial dilutions of effector TCD8+. For testing HPLC fractions, target cells in 50 µl of either PBS or FCS-free medium were exposed to 5 µl of fractions for 30 min at 26°C before TCD8+ were added. In some experiments, TCEP was freshly dissolved in H2O and used at 200 µM, both at peptide-pulsing and microcystotoxicity assay stages. The radioactivity in supernatants collected after 4−6-h incubation at 37°C was determined using a gamma counter. The percent specific release was then determined as % specific release = (CTL−induced release − spontaneous release)/ (release by detergent− spontaneous release) × 100.

**Results**

**TCD8+.** Specific for Cysteine-containing Determinants Are Specifically Required for Peptide-C class I Complexes. We previously reported that in Kd−restricted responses to PR8 influenza virus nuclear protein (NP), NP39-47,155 is the immunodominant determinant, with NP39-47 and NP218-226 exhibiting subdominant status (8). This hierarchy is not accounted for by peptide affinity, as NP39-47 binds to Kd with the lowest efficiency, as determined by a Kd− "melting" assay performed either with RMA-S cells expressing Kd from a
transfected gene (data not shown) or T2 cells (8). We initially focused on T<sub>CD8</sub> avidity to explain the immunodominance of NP<sub>147–155</sub>, as 10-fold less synthetic NP<sub>147–155</sub> was usually required to sensitize target cells for lysis by T<sub>CD8</sub> lines raised to the individual peptides under conditions similar to those used for the peptide binding assay (Fig. 1 B). This was observed using either short- or long-term lines stimulated in vitro by synthetic peptides derived from animals immunized either with PR8 or rVV expressing NP or cytosolic or endoplasmic reticulum (ER)-targeted minigene product versions of the determinants. Taking into account the lower efficiency of NP<sub>147–155</sub> binding to K<sup>d</sup>, the data in Fig. 1 suggest that <10% of K<sup>d</sup>-NP<sub>147–155</sub> complexes are required for T<sub>CD8</sub> triggering relative to K<sup>d</sup> complex to either of the subdominant determinants.

Several findings, however, suggested that matters might be a bit more complicated. Unlike NP<sub>147–155</sub>, the dose–response curves of NP<sub>39–47</sub> and NP<sub>218–226</sub> varied considerably between experiments, depending in part on the manner in which the assay was executed. We also experienced difficulties in stimulating and maintaining T<sub>CD8</sub> lines to these subdominant determinants, often observing slower growth after restimulation and morphological abnormalities of the cells, which were frequently larger than T<sub>CD8</sub> specific for NP<sub>147–155</sub>. This was not strictly related to the subdominant status of these determinants, as T<sub>CD8</sub> specific for other subdominant determinants behaved similarly to NP<sub>147–155</sub>-specific T<sub>CD8</sub>.

A property shared by NP<sub>218–226</sub> and NP<sub>39–47</sub> is the presence of cysteine (Table I). The report by Meadows et al. (4) demonstrating the dramatic effects of sulfhydryl modification of cysteine-containing residues on T<sub>CD8</sub> recognition prompted us to examine possible effects of cysteine modification on NP<sub>39–47</sub> and NP<sub>218–226</sub> binding and antigenicity. We first studied the properties of synthetic peptides in which cysteine is replaced by serine or alanine (the most conservative substitutions). For NP<sub>39–47</sub>, the cysteine→serine substitution had no significant effect on peptide binding, whereas the cysteine→alanine substitution increased peptide potency by ~10-fold (Fig. 2 A). For NP<sub>218–226</sub>, either substitution reduced peptide potency in stabilizing K<sup>d</sup> molecules by ~10-fold (Fig. 2 B). Each of the substitutions resulted in large increases in antigenicity relative to the wild-type peptides (Fig. 2, C and D). For NP<sub>218–226</sub>, the substituted peptides were 1,000–10,000-fold more antigenic on a per-complex basis (assuming that complex formation

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**Figure 1.** Antigenicity of synthetic peptides corresponding to dominant and subdominant determinants. (A) T2-<sup>K<sup>d</sup></sup> cells were cultured for 14 h at 26°C and added to the wells containing synthetic peptides at the indicated concentrations. The samples were immediately shifted to 37°C and incubated for 2 h to denature K<sup>d</sup> molecules lacking peptides and then stained with a fluorescent-conjugated anti-K<sup>d</sup> mAb. The mean channel fluorescence (MCF) of viable cells was determined by flow cytometry. (B) Splenocytes from PR8-primed animals stimulated in vitro for 7 d with synthetic peptides corresponding to NP<sub>39–47</sub>, NP<sub>147–155</sub>, or NP<sub>218–226</sub> were tested in a microcytotoxicity assay for their ability to lyse <sup>51</sup>Cr-labeled P815 target cells incubated in I-10 with synthetic peptides at the indicated concentrations.

**Figure 2.** Binding and antigenicity of synthetic peptides A and B show data from the K<sup>d</sup>-melting experiment displayed in Fig. 1 A. In this case, data for wild-type NP<sub>39–47</sub> and NP<sub>218–226</sub> are replotted with the substituted peptides. In C and D, T<sub>CD8</sub> specific for NP<sub>39–47</sub> or NP<sub>218–226</sub> were tested in a microcytotoxicity assay for their ability to lyse <sup>51</sup>Cr-labeled P815 target cells incubated in I-10 in the presence of the synthetic peptides at the indicated concentrations.
at the endpoint of peptide titrations is proportional to peptide binding efficiency determined by the melting assay.

These findings prompted us to study the effects of sulfhydryl modification on the antigenicity of NP<sub>39-47</sub> and NP<sub>218-226</sub>. In most experiments described below, the two peptides were studied in parallel. Because the results were highly similar, only results with NP<sub>218-226</sub> are shown for the sake of clarity and simplicity.

Reduction of cysteine enhances the antigenicity of cysteine-containing viral peptides. Cysteine readily forms disulfide bonds at neutral or slightly basic pH in the presence of O<sub>2</sub>. Cysteine forms can be reduced to the original thiol form by exposure to reducing agents. To determine whether disulfide formation affected peptide antigenicity, synthetic NP<sub>218-226</sub> was added to cells in the presence of dithiothreitol or TCEP, and cells were tested for lysis by NP<sub>218-226</sub>-specific T<sub>CDB+</sub>. Either of these reducing agents increased peptide potency by ~10-fold (Fig. 3 A). Reducing agents did not affect the potency of noncysteine-containing peptides, including NP<sub>147-155</sub> (Fig. 3 B), an LCMV peptide (described below), or the cysteine→serine- or cysteine→alanine-substituted NP<sub>39-47</sub> peptides (not shown). Enhancement of cysteine peptide recognition is, as expected, dependent on the concentration of reducing agent, with TCEP being more effective on a molar basis than dithiothreitol (not shown). The optimal concentration for TCEP was 200 μM (used in additional experiments), as higher concentrations (1 mM) sometimes increased spontaneous release values in <sup>51</sup>Cr-release assays. The effect of TCEP on NP<sub>218-226</sub> antigenicity is particularly impressive when considered in view of the 100–1,000-fold decrease in peptide binding to K<sup>+</sup> in the presence of TCEP (described below).

In additional experiments, we found that inclusion of reducing agents in the media used to stimulate and propagate T<sub>CDB+</sub> specific for NP<sub>218-226</sub> or NP<sub>39-47</sub> greatly enhanced their growth and altered their appearance, to the extent that these T<sub>CDB+</sub> were indistinguishable from T<sub>CDB+</sub> raised to NP<sub>147-155</sub> or other immunodominant determinants.

These findings indicated that sulfhydryl modification can have major effects on the antigenicity and immunogenicity of synthetic peptides, effects that can lead to erroneous conclusions regarding the nature of their interactions with class I molecules, the affinity of T<sub>CDB+</sub> specific for the peptide, and the growth characteristics of the cells. To broaden these findings, we examined the effect of reducing agents on the in vitro antigenicity of three other LCMV-specific, D<sup>+</sup>-restricted determinants and the T<sub>CDB+</sub> they induce.

Nature of Thiol-modifying Agents Affects NP<sub>218-226</sub>. We turned our attention to why reducing agents enhance the antigenicity of synthetic NP<sub>218-226</sub>. The major possibilities were reduction of disulfide-linked peptide dimers and reduction of disulfide-bound species derived from culture media. The most abundant sulfhydryl-containing compound present in DMEM is cysteine (cysteine-cysteine dimers). To determine whether NP<sub>218-226</sub> becomes cysteinylated in DMEM, freshly dissolved NP<sub>218-226</sub> was incubated in normal or cysteine-free DMEM for 2 h. Peptides present in media were then separated by RPHPLC, and the masses in peptide-containing fractions were determined by mass spectroscopy (Fig. 5). After incubation in cysteine-free media, the only modification detected was a small amount of dimerization (Fig. 5 B). By contrast, in cysteine-containing media, most of the monomer was converted to a separate eluting form representing cysteinylated peptide (Fig. 5 C). This fraction also contained a minor species with an additional mass of 16 daltons that probably represents oxidation of the neighboring methionine residue in the peptide.

The effect of cysteinylation on NP<sub>218-226</sub> antigenicity was
examined by measuring the Kd binding and antigenicity of an unmodified preparation of NP218–226 incubated for 2 h in cystine-containing DMEM with or without TCEP before addition to cells. As seen in Fig. 6 A, the resulting cysteinylation was associated with an ~10-fold increase in rescuing Kd molecules from melting and an ~10,000-fold decrease in capacity to sensitize target cells for lysis by TCD8+ induced by APCs pulsed with unmodified NP218–226.

We also examined the effect of NP218–226 dimerization on antigenicity and Kd binding. Analysis of various stocks by RP-HPLC in conjunction with mass spectrometry revealed that a 1-yr-old stock of peptide in DMSO was ~95% dimerized (Fig. 5 D). Using this stock as an NP218–226 dimer source, we investigated the effect of 2-h incubation at room temperature in cystine-free or cystine-containing DMEM. NP218–226 dimers were stable under these conditions (Fig. 5, E and F).

Having identified a source of dimers and demonstrated the stability of dimers in DMEM, we could examine the Kd binding and antigenicity of dimers (Fig. 6), which revealed that dimers behaved similarly to cysteinylated NP218–226.

We draw two conclusions from these findings. First, cysteinylation and dimerization of NP218–226 is associated with enhanced Kd binding yet greatly reduced antigenicity, using TCD8+ restimulated by the reduced peptide. Second, normal DMEM, cysteinylation occurs preferentially to peptide dimerization, even when NP218–226 is present at relatively high concentrations. As a second-order reaction, dimerization should be greatly disfavored at decreasing peptide concentrations, whereas cysteinylation continues at a first-order reaction rate. Therefore, cysteinylation is probably the major process for modifying cysteine-containing peptides at the concentrations used in Kd-binding and 3HCr-release assays.

Cysteinylated NP218–226 is produced by virus-infected Cells. Given the potential for cysteine modification in vitro, we examined whether NP218–226 produced by PR8-infected cells was modified in vivo. Low Mr peptides present in acid extracts from whole cells were fractionated by RP-HPLC and tested for their abilities to sensitize target cells for lysis by TCD8+ raised to reduced NP218–226. TCEP was added to the fractions to reveal the presence of SH-modified forms of peptides rendered nonantigenic by the modification. As shown in Fig.
7 A, antigenic peptides were recovered in fractions eluting from 25–27 min, matching the elution times of cysteinylated (25 min) and unmodified (27 min) NP218–226. No activity was present in the 29-min fraction, where dimeric NP218–226 elutes. The amounts of peptide recovered were well below that required for saturation (peak lysis of column fraction of 40% versus 70% with a saturating amount of synthetic peptide; Fig. 7 A, top right), suggesting that a considerable fraction of NP218–226 recovered from PR8-infected cells is cysteinylated. Cysteinylination of NP218–226 might have occurred artefactually during the extraction process. To examine this possibility, cell homogenates were doped with synthetic, unmodified NP218–226 and then processed identically to virus-infected cells (note that in this and subsequent experiments, to increase the chromatographic resolution, fraction size was reduced from 1 ml to 0.25 ml) (Fig. 7 B). In this case, 1% of the antigenic activity (as determined by titrating fractions; data not shown) was recovered in the cysteinylated form. In this experiment, peptides were tested in the presence or absence of TCEP. Even unmodified NP218–226 required TCEP treatment, which we attribute to rapid peptide cysteinylation during target cell sensitization.

To examine the maximal potential for posttranslational modification of NP218–226, we infected cells with an rVV (VV-NP218–226) that expresses the peptide in the cytosol as a minigene product (with NH$_2$-terminal methionine to enable translation initiation). As reported previously (7–9), this greatly enhances the number of peptide-class I complexes generated by cells. R-P-HPLC fractionation from minigene-expressing cells revealed the presence of material coeluting with unmodified NP218–226, as well as cysteinylated peptide (Fig. 7 C). For the first time, dimeric peptide was also detected. These activities cannot be attributed to methionine-extended NP218–226, which, for each form, elutes slightly later than NP218–226 (data not shown). Titration of the antigenic activities in the fractions (not shown) revealed a ratio of unmodified/cysteinylated/dimeric forms of ~6:3:1.

In an additional experiment (data not shown), we examined the K$^d$ dependence of NP218–226 recovery in HPLC fractions after VV-NP218–226 by using L929 cells and L929 cells expressing K$^d$ from a transgene. Peptides correspond-
ing to cysteinylated and unmodified NP<sub>218–226</sub> were recovered from L929-K<sup>d</sup> cells but not L929 cells. Dilution of peak fractions revealed that expression of K<sup>d</sup> resulted in at least a 25-fold increase in the recovery of NP<sub>218–226</sub>. This confirms numerous prior studies demonstrating the MHC dependence of antigenic peptide recovery (1).

Cysteinylated NP<sub>218–226</sub> Is Presented In Vivo. The ultimate demonstration of the biological relevance of cysteinylated NP<sub>218–226</sub> is that specific T<sub>CD8+</sub>, are elicited in PR 8-infected mice. We could show this by stimulating splenocytes derived from PR 8-infected mice with RP-HPLC-purified, cysteinylated peptide. After three to four rounds of stimulation, we obtained T<sub>CD8+</sub> that preferentially recognize cysteinylated peptide (Fig. 7 D). In the same assay, the noncysteinylated peptide is preferentially recognized by T<sub>CD8+</sub>, induced in the standard manner (Fig. 8 B). The recovery of T<sub>CD8+</sub> specific for the cysteinylated peptide could not be attributed to in vitro stimulation of naïve T<sub>CD8+</sub>, as we failed to obtain any activity using splenocytes from nonimmunized mice (data not shown).

Using T<sub>CD8+</sub> stimulated by the cysteinylated NP<sub>218–226</sub>, it was possible to formally demonstrate that the 25-min fraction derived from minigene-expressing cells contained cysteinylated peptide (Fig. 7 D). Indeed, now antigenicity was destroyed by TCEP exposure, in contrast to the enhancing effect derived from minigene-expressing cells contained cysteinylated peptide. After three to four rounds of stimulation, we obtained T<sub>CD8+</sub> that preferentially recognize cysteinylated peptide (Fig. 8 A). In the same assay, the noncysteinylated peptide is preferentially recognized by T<sub>CD8+</sub>, induced in the standard manner (Fig. 8 B). The recovery of T<sub>CD8+</sub> specific for the cysteinylated peptide could not be attributed to in vitro stimulation of naïve T<sub>CD8+</sub>, as we failed to obtain any activity using splenocytes from nonimmunized mice (data not shown).

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Discussion

In this paper, we confirm and extend the findings of Meadows et al. (4) that modification of the SH group of cysteine-containing peptides has important positive and negative effects on their antigenicity and immunogenicity in vitro and in vivo. Failure to consider these effects can have disastrous consequences for the accurate interpretation of several different types of experiments.

In our own case, the use of synthetic peptides in the absence of reducing agents led us to erroneously favor the idea that the subdominant status of two cysteine-containing peptides was due to a greater number of complexes required for T<sub>CD8+</sub> recognition and correlated with the atypical growth of T<sub>CD8+</sub> in vitro. In other experiments (our unpublished results), it also led us to the incorrect quantitation of peptides recovered from virus-infected cells. A further potential methodological pitfall is that autooxidation of reducing agents can cause additional artifacts. This can be minimized by inclusion of a chelating agent, such as DTPA (diethylenetriaminepentaacetic acid), with the reducing agent.

These errors are probably widespread. We demonstrate that two cysteine-containing LCMV determinants restricted by a different class I molecule are also modified in vitro through their sulfhydryl groups, with a resulting 10–100-fold loss in antigenicity. These determinants were the subject of a recent study (10) focused on the factors involved in the immunodominance hierarchy of the determinants. The failure to add reducing agents during peptide titration probably led to erroneous calculations of peptides present in virus-infected cells, particularly because the peptide titration curves were nearly identical to those we obtained in the absence of reducing agents. Results obtained in this study with in vivo transfer of T<sub>CD8+</sub> lines must also be questioned, as the cells were propagated in vitro with synthetic peptides in the absence of reducing agents. In another recent study of the fine specificity of a T<sub>CD8+</sub> clone for a cysteine-containing peptide, amino acids were substituted for cysteine, many of which enhanced the antigenicity of the peptide (11). Based on our findings, we would predict that simple reduction of the wild-type peptide would have a similar (or greater) effect.

In addition to reducing antigenicity, modification of cysteine can result in the generation of T<sub>CD8+</sub> specific for the modified determinant. We show that PR 8-infected mice generate T<sub>CD8+</sub> that prefer cysteinylated NP<sub>218–226</sub>. We also demonstrate that PR 8-infected cells generate an SH-modified peptide that coelutes with cysteinylated NP<sub>218–226</sub> and is recognized by T<sub>CD8+</sub> specific for cysteinylated NP<sub>218–226</sub>. This species almost certainly represents cysteinylated NP<sub>218–226</sub>, although definitive evidence requires mass spectrometry. The recognition of posttranslationally modified peptides by T<sub>CD8+</sub> adds to the already formidable challenge of understanding in vivo T<sub>CD8+</sub> responses but can be ignored only at the peril of the investigator.

We can only speculate where NP<sub>218–226</sub> is cysteinylated during its processing and presentation by virus-infected cells. It is theoretically possible that the cystine derives naturally from a disulfide bond present in NP. It is difficult, though admittedly not impossible, to imagine the proteolytic liberation of cysteinylated NP<sub>218–226</sub>. Given the highly reducing environment of the cytosol and nucleus, it also seems unlikely that cysteinylation would occur before peptide translocation into the ER. The ER provides a much more oxidizing environment and possesses resident proteins that catalyze thiol-disulfide interchange, including protein disulfide isomerase (12). There is evidence that exogenous homocysteine is added to HLA class I molecules.
in an early secretory compartment (13). If cysteinylilation occurs in the ER, it may occur before peptide loading onto class I molecules. Alternatively, cysteinylilation could occur after peptide binding, particularly in the case of NP218–226, as the SH must be directed away from the groove (so as to accommodate dimer binding). In this case, it could occur anywhere from the ER to the cell surface. Cysteine is thought to be the major reductant in the endosomal pathway, and alternatively, cysteinylation could occur in the ER, it may occur before peptide loading onto class I molecules. Cysteine is thought to be the major reductant in the endosomal pathway, and it may occur in the vicinity of TAP, which is required for transport of NP218–226 into the ER (our unpublished results).

These findings have important clinical implications. First, for synthetic peptide vaccines (or other exogenous antigen vaccine preparations with vulnerable cysteine residues in antigenic peptides), modification of the cysteine in vitro or in vivo can obviously have major negative effects on immunogenicity. This can be avoided by modifying the side chain to a nonreactive form. For NP218–226 and NP39–47, this is achieved simply by substitution with alanine or serine, which did not detrimentally affect peptide binding or T CD8+ triggering. This strategy will probably work for most peptides. For others, it is possible that chemical modification of the SH group (e.g., treatment with an alkylating agent or a heavy metal) will do the job. Second, if increased cysteinylilation is associated with a disease process, this could lead to autoimmune recognition of cysteinylated self peptides.

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