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A pentapeptide as minimal antigenic determinant for MHC class I-restricted T lymphocytes. — Source link <a> □

Matthias J. Reddehase, Jonathan B. Rothbard, Ulrich H. Koszinowski

Institutions: Lincoln's Inn, University of Ulm

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Table 3 Effect of DIDS and chloride absence on acid-loading rate (J_{load}) following a 10% CO₂ prepulse

		J	load (µM S-1)	
	Control	DIDS	No chloride	DIDS-sensitive (HCO ₃ transport)
-AVP	630 ± 22	62 ± 3	51 ± 5	568
+ AVP	$1,504 \pm 76$	119±9	80 ± 3	1,385
AVP dep.	874 ± 35	57 ± 8	31 ± 7	717
% Stimulation	140 ± 11			143

Fluxes calculated as in Table 1, but at pH 7.7. β_{HCO_1} was 114.9 mM per pH unit at pH, 7.7.

crucial mitogenic signal. It could be advantageous for an activated cell to maintain a steady-state pH; close to the unactivated level given the sensitivity to pH of nearly all cellular processes. Such a steady-state pH; is achieved when processes that acidload the cell (for example Na+-independent Cl-/HCO3 exchange) balance those that alkali-load it (for example, Na⁺/H⁺) and Na⁺-dependent Cl⁻/HCO₃ exchange). It would also be advantageous however, for the activated cell to be able to recover more rapidly from acid and alkali loads. The requirements for both a near-normal steady-state pH; and enhanced pH; regulation can only be met if increases in acid loading and acid extrusion by the growth factor are comparable. This is precisely the response of the mesangial cell to AVP: application of the growth factor causes only a small decrease in steady-state pHi, but enormous increases in the rate at which the cell recovers from both acid (Fig. 1a) and alkali loads (Fig. 2c). This effect of AVP on pH; does not seem to be an isolated event, as we have found eight other growth factors that also produce only a small pH_i decrease in the presence of HCO₃ (data not shown). Thus, an important physiological effect of the growth factor is to maintain a near-normal pH; while stimulating several acidbase transporters.

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A pentapeptide as minimal antigenic determinant for MHC class I-restricted T lymphocytes

Matthias J. Reddehase*, Jonathan B. Rothbard† & Ulrich H. Koszinowski‡

- * Federal Research Centre for Virus Diseases of Animals, POB 1149, 7400 Tübingen, FRG
- † Laboratory of Molecular Immunology, Imperial Cancer Research Fund, Lincoln's Inn Fields, London WC2A 3PX, UK
- Department of Virology, Institute for Microbiology,
- University of Ulm, POB 4066, 7900 Ulm, FRG

Peptides that are antigenic for T lymphocytes are ligands for two receptors, the class I or II glycoproteins that are encoded by genes in the major histocompatibility complex, and the idiotypic α/β chain T-cell antigen receptor 1-9. That a peptide must bind to an MHC molecule to interact with a T-cell antigen receptor is the molecular basis of the MHC restriction of antigen-recognition by T lymphocytes 10,11. In such a trimolecular interaction the aminoacid sequence of the peptide must specify the contact with both receptors: agretope residues bind to the MHC receptor and epitope residues bind to the T-cell antigen receptor 12,13. From a compilation of known antigenic peptides, two algorithms have been proposed to predict antigenic sites in proteins. One algorithm uses linear motifs in the sequence14, whereas the other considers peptide conformation and predicts antigenicity for amphipathic α helices^{15,16}. We report here that a systematic delimitation of an antigenic site precisely identifies a predicted pentapeptide motif as the minimal antigenic determinant presented by a class I MHC molecule and recognized by a cytolytic T lymphocyte clone.

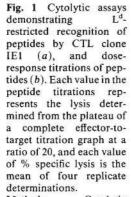
Synthetic peptides have been derived from the amino-acid sequence of pp89, an immediate-early (IE) phase regulatory protein of murine cytomegalovirus¹⁷⁻²⁰. The 19-mer P(161-179) contained within its 595 residues¹⁸ is an antigenic sequence²¹. sequence H₂N-¹⁶¹GRLMYDMYPHFMPTNLGPS¹⁷⁹-

Delimitation of the antigenic motif for CTL clone IE1

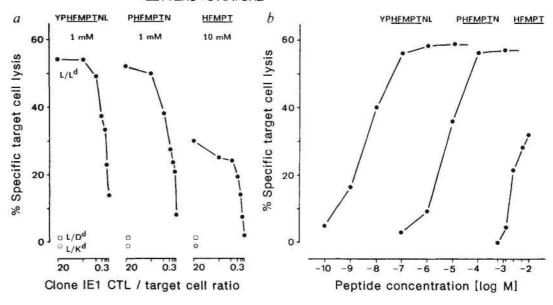
			Peptide concentration [log M]	
Peptide		Recognition (competition)	detection limit	detection saturation
11-mer:	MYPHFMPTNLG	+	-7 to -6	-4 to -3
10-mers:	MYPHFMPTNL	+	-9 to -8	-7 to -6
	YPHFMPTNLG	+	-8 to -7	-6 to -5
9-mers:	MYPHFMPTN	-(-)		
	YPHFMPTNL	+	-12 to -10	−9 to −7
	P <u>HFMPT</u> NLG	+(-)	-4 to -3	-2
8-mers:	YPHFMPTN	-(-)		
	P <u>HFMPT</u> NL	+(-)	-4 to -3	
7-mers:	YPHFMPT	-		
	P <u>HFMPT</u> N	+	-8 to -7	-4
	HFMPT NL	+	−5 to −4	
6-mers:	YPHFMP	-		
	PHFMPTN	+	-2 to -3	
	HFMPTN	+	-2	
	FMPT NL	-		
5-mer:	HFMPT	+	-3	
4-mers:	HFMP	-		
	FMPT	-		

Dose-response titrations of peptides give the peptide molarities in solution required for detectable target formation (detection limit) and for optimal target formation (detection saturation). The concentration ranges are compiled from at least 3 and up to 12 independent experiments.

COOH (one-letter code) contains the predicted motif HFMPT¹⁴. By screening a series of related peptides that had been reduced in length from both terminals (not all shown), we found that the nonapeptide YPHFMPTNL represents the optimal antigenic peptide (Fig. 1 and Table 1) for the cytolytic T lymphocyte (CTL) IE1 clone^{22,23}. Even though binding of peptides to class I molecules has not yet been demonstrated directly, the selective recognition on L/Ld cells implies that the Ld molecule is the



Methods. Cytolytic effector cells: The murine (BALB/c strain; MHC^d) CTL line IE1.13-IL was derived from clone IE1 by recloning^{22,23}. Clone IE1 expresses an α/β (V β 6) TCR/CD3 complex and displays the surface



phenotype CD4⁻CD5⁻CD8⁺Thy-1⁺. L fibroblast transfectants: L/L^d cells were derived from L cells by transfection with the L^d gene, and L/D^d and L/K^d transfectants were provided by M. Cochet and J. P. Abastado (Institut Pasteur). Surface expression of the L^d, D^d and K^d molecules was confirmed by cytofluorography with monoclonal antibodies. Synthetic peptides: Peptides were synthesized by using the Barany-Merrifield solid-phase technique on an Applied Biosystems Peptide Synthesizer, purified preparatively and analysed as described previously¹³. Preparation of target cells and cytolytic assay: Target cells were labelled at 37 °C for 1 h with Na₂[⁵¹Cr]O₄. The radioactive cells were then distributed in aliquots of 10⁵ cells and incubated in 0.2 ml for a further 1 h at 27 °C with peptides at defined molarities (1 M corresponds to 10¹⁵ peptide molecules per cell) dissolved in RPMI 1640 culture medium supplemented with 2% of FCS. The highest peptide concentration tested was 10⁻² M. This limit was imposed by the solubility of the peptides. After washing to remove excess peptide and ⁵¹Cr, a standard 3-h cytolytic assay was performed at 37 °C with 1,000 target cells and graded numbers of clone IE1 CTL. Competition assays: The antigenic peptides to compete with the optimal antigenic nonapeptide YPHFMPTNL was tested in two ways. Either the concentration of the antigenic peptide was kept constant at 10⁻⁴ M and the antigenic peptide was titrated from 10⁻⁵ to 10⁻¹⁰ M. In both cases the L/L^d target cells were pre-incubated for 30 min with the competitor peptide before the antigenic peptide was added.

receptor involved in the interaction with the T-cell antigen receptor (TCR) of clone IE1 for the family of peptides tested (Fig. 1a). Titration of the nonapeptide YPHFMPTNL, the heptapeptide PHFMPTN, and the pentapeptide HFMPT resulted in dose-response saturation curves showing thousandfold differences in antigenic potency (Fig. 1b). The data refer to a one-hour incubation of target cells with peptide. Differences in antigenic potency are also reflected by the time needed for optimal target formation; five minutes is long enough to prepare an optimal nonapeptide target, which is in line with data for a high-affinity peptide24, whereas plateau lysis could be achieved with the pentapeptide only when the peptide pulse was prolonged to two hours or more (not shown). Because clone IE1 does not discriminate between nonapeptide and pentapeptide target once the target is formed, the limiting step is apparently the association between the peptide ligand and its MHC receptor on the target-cell surface. We therefore conclude that HFMPT forms the antigenic core of the peptides by comprising both the complete epitope for the TCR of clone IE1 and an agretope that is adequate, but not optimal for specification of the interaction with Ld

The deletion of Tyr1 (Y) from peptide YPHFMPTNL diminished and the deletion of Leu 9 (L) destroyed the antigenicity, and the resulting octapeptides PHFMPTNL and YPHFMPTN both also failed to compete with YPHFMPTNL (Table 1). According to the currently used theorem for classifying residues as T-cell contact residues or as MHC-molecule contact residues¹², Tyr 1 and Leu 9 would have been interpreted as residues contacting the MHC molecule. This conclusion, however, was proven incorrect by the findings that PHFMPTN, which lacks both residues, was >10³-fold more antigenic than PHFMPTNL (Table 1) and could not be competed by a 10³-fold excess of YPHFMPTN (not shown), although both octapeptides include the heptapeptide sequence entirely. Results compiled

in Table 1 follow a consistent pattern from the pentameric core motif up to the nonapeptide size, in that symmetric additions of residues improved the antigenicity, whereas addition of residues at either end had a negative effect.

This communication adds new aspects to the current understanding of the MHC molecule-peptide-TCR interaction. The first aspect concerns the minimal length of antigenic peptides. Until now, the shortest peptides reported as antigenic were heptapeptides presented by MHC class II molecules^{25,26}. A recent study described a nonapeptide presented by an MHC class I molecule as minimal antigenic peptide of the glycoprotein of LCMV²⁷. It should be emphasized that in our example systematic shortening from both termini was critical for the identification of the pentapeptide as the minimal antigenic peptide. Since the sequence of this pentapeptide is a predicted pentameric motif, and as most motifs are tetrameric¹⁴, recognition even of tetrapeptides may be possible.

The second aspect concerns the classification of epitope and agretope residues. For the hen egg-white lysozyme peptide (52-61), Allen et al.12 defined residues as MHC-antigen contact residues when a substitution led to the loss of both antigenicity and the ability to compete with the unmodified antigenic peptide. With this approach and proposing a helical conformation, agretope residues were segregated from epitope residues. From a set of overlapping peptides, Sette et al.28 predicted a heptameric core in a planar conformation for the ovalbumin peptide (323-339). In contrast to these examples, we have positively identified the motif HFMPT as an antigenic core in pp89 peptides by demonstrating a direct antigenicity of the pentapeptide HFMPT. The important implication is that residues whose deletion causes loss of both antigenic potency and competitive ability are not necessarily agretope residues in the classical sense of MHCreceptor binding sites. We propose that residues flanking an antigenic core motif can affect the antigenic potency positively

or negatively by their influence on peptide conformation.

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Class I cross-restricted T cells reveal low responder allele due to processing of viral antigen

Helen C. Bodmer, Frances M. Gotch & Andrew J. McMichael

Institute for Molecular Medicine, John Radcliffe Hospital, Oxford OX3 9DU, UK

Cytotoxic T lymphocytes (CTL) recognize protein antigens which have been processed by the target cell and then presented in association with the relevant class I molecule of the major histocompatibility complex (MHC)1. Short synthetic peptides, which are able to associate directly with target cells1, may substitute for these processed fragments in stimulating antigen-specific CTL responses. Using this approach, a dominant HLA-A2-restricted epitope has previously been mapped to residues 58-68 of influenza A virus matrix protein². Here we report HLA-A2-restricted CTL which are also able to recognize this short synthetic peptide in association with HLA-Aw69, but which fail to recognize HLA-Aw69 expressing cells infected with influenza A virus. Furthermore, individuals possessing HLA-Aw69 who respond to influenza A virus, do not respond to M58-68. These results imply that the low response to this epitope on infection of HLA-Aw69 individuals with influenza A is due to failure of the naturally processed product of matrix protein to associate with Aw69.

Individuals who express the common HLA-A2 variant, HLA-A2.1 (HLA-A*0202 as defined in ref. 3), and who can mount a cytotoxic response to influenza infected target cells, following in vitro stimulation with virus have been shown, in all cases tested to date, to possess CTL that specifically recognize synthetic peptides representing residues 56-68 of the matrix protein⁴. Individuals who have other variants of A2, or the closely

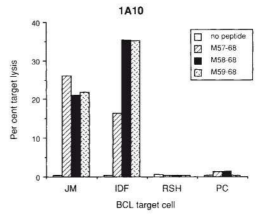


Fig. 1 CTL are cross-restricted, recognizing peptide on both autologous and Aw69-expressing target cells. Recognition by CTL clone 1A10 from the donor JM (A2, 2; B15, 51; DR4, 4) of peptides M57-58, M58-68, and M59-68 of influenza matrix protein on autologous BCL target cells (JM), three HLA-mismatched BCL, IDF (A26, w69; B18, 38; DR11, 11), RSH (A68, 68; B42, 42; DR3, 3) and PC (A3, 24; B37, 62; DR6, 6). K/T = 1.5, peptides present throughout assay 10 µM.

Methods. CTL clone 1A10 was derived from a parent line from the donor, JM. Peripheral blood lymphocytes were separated from whole blood, stimulated initially with A/X31 virus (grown and stored as previously described¹⁸) and maintained in culture by repeated stimulations with peptide M57-68, as previously described4. CTL were cloned by selection of CD8 positive cells (labelled with B941 antibody from Dr C. Mawas, and fluoresceinlabelled anti-mouse immunoglobulin) using an Ortho Cytofluorograf cell sorter, into 96-well plates containing 104 irradiated BCL (pretreated with 50 µM M57-68) and IL-2 10 U ml⁻¹ (Cetus Corp.) in RPMI containing 10% fetal calf serum. Monoclonality was indicated by the presence of a single β -chain gene rearrangement on Southern blotting (data not shown). Cytotoxic activity was assessed by incubating EBV-transformed lymphoblastoid cells (BCL), labelled with 51Cr, with CTL and diluted peptide (where appropriate) for 4 h in round-bottomed 96-well plates. Per cent target lysis was calculated from the formula $(E - M/D - M) \times 100$ where $E = \text{experimental}^{51}\text{Cr}$ release, M = release in presence of culture medium (always <18%) and D = release by 5% Triton X-100. For details of the peptides see legend to Fig. 3.

related allele HLA-Aw69, which differs by six amino acids from A2*0202 (listed in Table 1)5, did not respond to this region of matrix protein4.

We made several CTL lines and clones from a single HLA-A2 expressing individual, JM, some of which were able to recognize matrix peptide 56-68 in association with HLA-Aw69 as well as A2, although they were otherwise HLA-restricted (Fig. 2). CTL clone 1A10 was able to recognize the short synthetic peptides M57-68, M58-68 and M59-68 in association with both the autologous (JM) HLA-A2-expressing EBV-transformed B lymphoblastoid cell line (BCL), and a BCL expressing Aw69 which did not share any HLA alleles with JM (Fig. 1), but did not recognize two other HLA mismatched BCL in the presence of these peptides.

Although these matrix-specific A2-restricted CTL were able to recognize Aw69-expressing target cells in the presence of synthetic peptide, a discrepancy between recognition of A2 and Aw69 BCL was seen following A/X31 virus infection of the target cells (Fig. 2). Both autologous and A2-matched BCL were recognized following pretreatment with the peptide, or infection with influenza virus (Fig. 2a, e). In contrast Aw69-expressing BCL were very poorly recognized following virus infection, although they were well recognized after peptide treatment (Fig. 2b, c, f). In addition, virus-specific polyclonal CTL from an Aw69-expressing donor were unable to recognize the synthetic peptide M58-68 (Fig. 2d). Even with repeated stimulation of these latter CTL with M58-68 in vitro, we were unable to obtain a response from either of two Aw69-expressing (HLA identical)