Two Distinct Conformations of a Rinderpest Virus Epitope Presented by Bovine Major Histocompatibility Complex Class I N*01801: a Host Strategy To Present Featured Peptides

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Viral diseases and other infections caused by intracellular pathogens of cattle result in large global economic losses every year. In the past several decades, rinderpest virus (RPV) (23, 40), foot-and-mouth disease virus (FMDV) (7), and bovine tuberculosis infection (62) have brought disastrous consequences, including social panic and even a threat to human health. Nearly 6 decades ago, most countries were committed to developing more-effective vaccines (46), and remarkable achievements were made in controlling threatening bovine viral infectious diseases and other zoonoses (23, 46). In the 1960s, British virologist Walter Plowright developed a live attenuated vaccine against rinderpest virus, which was widely used in rinderpest eradication efforts (40). However, it still remains extremely challenging for humans to completely eradicate a virus. The questions of how the bovine attenuated vaccine induces immune responses and how the viral peptides are presented and recognized by host immune molecules are still largely unanswered. Therefore, studies on bovine immunity against the rinderpest virus may provide clues for our battle against other viral infections.

Generally speaking, effective vaccines have common characteristics: they are rich in T-cell and B-cell epitopes and can induce the immune system to generate protective immune responses (3, 36, 54). During this process, major histocompatibility complex (MHC) molecules play a pivotal role in the host. MHC class I (MHC I) molecules form a heterotrimeric complex composed of the heavy chain of MHC, the antigenic peptide, and 2-microglobulin (β2m). Generally, MHC I loads endogenous peptides, including the virus-derived peptides. Briefly, at the initial stage of virus infection, antigenic proteins are processed in a proteasome-dependent or -independent manner. The resultant short peptides are then transported to the endoplasmic reticulum and are loaded onto the peptide-loading MHC I complexes translocated to the cell surface and are recognized by cytotoxic T lymphocytes (CTLs) with specific T-cell receptors (TCRs) (49, 63). This immune recognition induces an MHC-restricted CD8 T-cell response that is characterized by the killing and elimination of the infected cells by effector T cells. The majority of virus-derived peptides are processed and presented through this pathway...
and consequently trigger the CTL-specific immune responses. These virus-derived peptides presented by MHC I molecules are known as antigenic CTL-specific epitopes.

Based on structural studies, the antigenicity of a peptide is partially dependent on the characteristics of the peptide related to its presentation by MHC molecules (31, 33, 34, 53, 55). Generally, peptides are divided into featured, featureless, and bulged peptides according to their presentation properties (25, 57). Featured peptides are those with solvent-exposed, prominent side chains or harmonious bulged conformations, which always correspond to a diverse repertoire of TCRs (32, 37, 38, 58). Featureless peptides have fewer or no solvent-exposed residues with prominent side chains. Bulged peptides are usually long peptides (>12 amino acids) that contain an extreme bend in the middle of the peptide chain binding to the MHC I molecule. The structural landscapes of featureless and bulged peptides result in an immune T-cell repertoire of limited diversity (58–60). Therefore, investigation of the structural characteristics of a peptide presented by MHC I molecules can improve the understanding of the antigenicity of that peptide and aid in the rational development and modification of peptide-based vaccines (33).

Bovine MHC I, which is called BoLA-I, has a critical role in presenting virus-derived antigen peptides and dominating bovine antiviral-specific CTL immune responses. In the 1970s, BoLA-I protein was first identified and genotyped by use of a specific serum antibody. BoLA-I was first cloned in 1988 (20), and since then, BoLA-I related research has entered the gene era. Thus far, hundreds of BoLA-I cDNA sequences have been registered at the National Center for Biotechnology Information (NCBI) (http://www.ncbi.nlm.nih.gov/), and 58 complete BoLA-A sequences have been uploaded to the Immune Polymorphism Database (IPD) (http://www.ebi.ac.uk/ipd/) (5). BoLA is located on the 23rd bovine chromosome (22), including at least six expressed gene loci (29). Further research has demonstrated that there are one to three gene loci that can be transcribed and expressed in one BoLA haplotype (16, 17). The fact that a breed of BoLA haplotypes expresses only one BoLA-A gene locus (6, 15) provides an important experimental model for controlling bovine viral diseases in cattle with a single MHC I gene locus.

Due to polymorphisms in BoLA-I, there are more than 50 serotypes of one BoLA-I allele locus (4, 13). The BoLA-A11 serotype specifically indicates that the BoLA-A11/pBoLA-I9 (D18.3) gene product N*01801 is expressed (18, 28, 47, 48). Hegde and colleagues obtained the N*01801 peptide binding motif by monoclonal antibody purification of MHC molecules and subsequent acid elution of peptides (28). The majority of the peptides that occupied the binding groove of N*01801 were nonamers. There was clear evidence that position 2 of the peptides was occupied by Pro, and the C-terminal amino acid anchor was Ile/Val. Among the alleles whose motifs have been reported, the peptide motif of N*01801 is similar to those of H-2Ld, HLA-B7, HLA-B*3501, HLA-B*5101, HLA-B*5102, HLA-B*5103, HLA-B*7801, and HLA-Cw*0401 (21, 45). Sequence comparison of the bovine MHC I allele N*01801 to these MHC I molecules demonstrates that key residues share similar characteristics (and are conserved in some cases), which are involved in accommodating the similar anchor residues in P2 and P9. This was the first report of a BoLA allele-specific peptide motif (ASPm; 28). To identify bovine-restricted CTL epitopes derived from pathogens, according to N*01801 ASPm, dozens of potential bovine CTL epitopes have been synthesized, and their antigenicity was preliminarily analyzed by CTL assays (27). In 2004, one of the CTL epitopes with a typical N*01801 ASPm (peptide IPA) was identified in rinderpest virus hemagglutinin (H) protein amino acids 408 to 416 (IPAYGVLTI) (51). Recently, Guzman et al. confirmed that there is an MHC-restricted CD8+ T-cell response after FMDV infection (26). However, the mechanism of bovine antiviral-specific T-cell immunity remains unclear, partially due to the lack of clues as to the binding and presentation of the antigenic peptides by BoLA-I.

To describe the landscape of peptide presentation by bovine MHC I N*01801 and to understand the immune basis of the eradication of rinderpest virus, we determined the crystal structures of N*01801 complexed to the rinderpest virus-derived peptide IPA. Analysis of the structures we determined suggests that compared to the structures of other MHC I molecules, including that of bovine N*01301, which was determined recently by Macdonald and colleagues (34), the structure of bovine MHC I N*01801 is distinctive. The structure of bovine N*01801 illuminates a novel presentation strategy for featured epitopes among mammalian MHCs and may lead to improved understanding of the structural basis of CTL-based immune responses (9).

**MATERIALS AND METHODS**

**Peptide synthesis.** The peptides used in this study (Table 1), including peptide IPA (IPAYGVLTI), derived from rinderpest virus attachment glycoproteins (51), were synthesized and purified by reverse-phase high-performance liquid chromatography (HPLC) (SciLight Biotechnology). The peptide purity was determined to be ~90% by analytical HPLC and mass spectrometry. These peptides were stored in lyophilized aliquots at −80°C after synthesis and were dissolved in dimethyl sulfoxide (DMSO) before use.

**Preparation of the bovine MHC I N*01801-β2m-IPA complexes.** Reverse transcription-PCR (RT-PCR) was used to amplify the full-length cDNA of N*01801 (A11) and bovine β2m from bovine kidney cells. Details of the primers used can be found in Table S1 in the supplemental material. The extracellular

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**TABLE 1. Peptides used in this study**

| Name | Derived protein | Position | Sequence | Reference |
|------|----------------|----------|----------|-----------|
| IPA  | Rinderpest virus H protein | 408–416 | IPAYGVLTI | 51 |
| APA  | BHV-F BICP0 protein | 482–490 | APAPSTM | 27 |
| NPM  | BHV-I glycoprotein B precursor | 780–788 | NPMKALYP | 27 |
| TPG  | BHV-I glycoprotein C precursor | 18–26 | TPGATTPY | 27 |

a Underlined boldface residues are the typical primary anchors of the peptides presented by N*01801.
b Peptide IPA was used in the structural determination of the bovine MHC class I N*01801.
c BHV-I, bovine herpesvirus 1.
region of N*01801 (amino acids 2 to 275) was amplified from the cDNA according to its nucleotide sequence (GenBank accession no. BC151402) using primers 1 and 2. Bovine β₂m (amino acids 1 to 98) was cloned from bovine kidney cell cDNA according to its nucleotide sequence (GenBank accession no. BC118352) with primers 3 and 4. Mouse β₂m (amino acids 1 to 99) was cloned from BALB/c mouse spleen cDNA according to its nucleotide sequence (GenBank accession no. M84364.1) with primers 5 and 6. The amplified products were ligated into a pET21a vector (Novagen) and were transformed into Escherichia coli strain BL21(DE3). The recombinant proteins were expressed as inclusion bodies and were then purified as described previously (8).

The N*01801-β₂m-peptide complexes were prepared essentially with refolding assays as described by Zhou et al. in 2004 (64). Briefly, the N*01801 heavy chain and bovine or murine β₂m inclusion bodies were separately dissolved in a solution of 10 mM Tris-HCl (pH 8.0) and 8 M urea. The N*01801 heavy chain, β₂m, and the peptide in a 1:1:3 molar ratio were refolded by gradual dilution. After 48 h of incubation at 4°C, the soluble portion of the complexes was concentrated and then purified by size exclusion chromatography on a Superdex 200 16/60 column. The comparison of the absorbance peaks of the refolded complexes using different peptides is presented in Fig. 1. If the complex was prepared for crystallization, it was further purified by Resource-Q anion-exchange chromatography (GE Healthcare).

**Crystallization and data collection.** The BoLA complexes were ultimately concentrated to 12 mg/ml in crystallization buffer (4.0 mM sodium formate), mixed with reservoir buffer at a 1:1 ratio, and crystallized by the hanging-drop vapor diffusion technique at 291 K. A Crystal Screen kit (Hampton Research, Riverside, CA) was used to screen for optimal crystal growth conditions. After several days, crystals of N*01801 complexed with the IPA peptide and bovine β₂m (N*01801) were obtained with Crystal Screen solution 33 (4.0 mM sodium formate). Diffraction data were collected to a resolution of 2.7 Å using an in-house X-ray source (Rigaku MicroMax007 desktop rotating anode X-ray generator with a Cu target operated at 40 kV and 30 mA) and an R-AXIS IV++ imaging-plate detector at a wavelength of 1.5418 Å. Similarly, the crystals of N*01801 complexed with the IPA peptide and murine β₂m (N*01801-Mβ) were grown in 0.2 M ammonium sulfate, 0.1 M Tris (pH 8.5), and 25% (wt/vol) polyethylene glycol 3350 (PEG 3350) at a concentration of 12 mg/ml, and the resolution of the diffraction data was 1.9 Å. The crystals were first soaked in reservoir solution containing 25% glycerol as a cryoprotectant and were then flash-cooled in a stream of gaseous nitrogen at 100 K (42). The collected intensities were indexed, integrated, corrected for absorption, scaled, and merged using the HKL2000 package (41).

**Structure determination and refinement.** The structures of the BoLA complexes were solved by molecular replacement using the MOLREP program with HLA-B*5101 (Protein Data Bank [PDB] code 1E27) as a search model. Extensive model building was performed by hand with COOT (19), and restrained refinement was performed using REFMAC5. Additional rounds of refinement were performed using the phenix refine program implemented in the PHENIX package (2) with isotropic atomic displacement parameter (ADP) refinement and bulk solvent modeling. The stereochemical quality of the final model was assessed with the PROCHECK program (30).

**Protein structure accession numbers.** The coordinates and structure factors of N*01801 and N01801-Mβ have been deposited in the Protein Data Bank with accession numbers 3PWV and 3PWU, respectively.

**RESULTS**

**Typical peptides bound to bovine MHC I molecule N*01801 revealed by in vitro refolding.** A series of bovine MHC I allele N*01801-restricted CD8⁺ T-cell epitopes derived from different pathogens have recently been screened and characterized (14, 27, 51); from these epitopes, the peptide motif of this MHC I molecule was defined. However, no rapid and efficient methods to evaluate the ability of a peptide with a given motif to bind to bovine MHC I molecules in vitro have been developed yet. As indicated in previous studies (31–33, 55), the refolding experiments can semiquantitatively reflect the binding capability of peptides for MHCs. Four previously identified N*01801-restricted CD8⁺ T-cell epitopes (Table 1) were synthesized and used in refolding assays to evaluate their abilities to bind to N*01801 molecules and to define the candidates used for determination of the N*01801 structure. These four N*01801-restricted peptides helped the heavy chain of N*01801 and bovine β₂m refold in vitro, as evidenced by the absorbance peaks of the complexes (Fig. 1). All four peptides contain a proline at position P2, while the Pc (C terminus of the peptide) positions were occupied by either isoleucine or...
N*01801 consists of the characteristic with a resolution of 2.7 Å (Table 2). The overall structure of N*01801-Mₐ (82.9%).

Between the intraspecies alleles of cattle: N*01801 and sequentially the most similar to N*01801, displaying 80.29% identity (Fig. 3). Notably, this is at the same level as the identity with the corresponding portions of N*01801, with RMSDs of 0.337 and 0.345 Å to molecule 1 and molecule 2, respectively (Fig. 4). This may indicate that the structure of the peptide loaded in the groove formed by the α₁ and α₂ domains of N*01801 was not affected by the substitution of the βₐm subunit. In the consideration of a higher resolution of N*01801-Mₐ (with murine βₐm), we use this structure together with the structure of N*01801 (with bovine βₐm) to analyze peptide presentation by bovine MHC I.

Two distinct conformations of peptide IPA presented by bovine MHC I N*01801. In the BoLA binding grooves of molecule 1 and molecule 2, the electron densities around the IPA peptides are well defined and clearly reveal two distinct

**Table 2. X-ray diffraction data processing and refinement statistics**

| Statistic                      | N*01801 | N*01801-Mₐ |
|--------------------------------|---------|------------|
| Data processing                |         |            |
| Space group                    | P₄₁₂₂   | P₂₁₂₁      |
| Cell parameters                | a (Å)   | 83.9       | 47.9       |
|                               | b (Å)   | 83.9       | 70.0       |
|                               | c (Å)   | 153.0      | 120.8      |
|                               | α (°)   | 90.0       | 90.0       |
|                               | β (°)   | 90.0       | 90.0       |
|                               | γ (°)   | 90.0       | 90.0       |
| Resolution range (Å)           | 50.0–2.7 (2.8–2.7)* | 50.0–1.9 (1.97–1.9) |
| Total reflections              | 201,829 | 228,256    |
| Unique reflections             | 28,833  | 15,065     |
| Completeness (%)               | 99.4 (98.9) | 99.9 (100.0) |
| Rmerge (%)                     | 14.2 (54.8) | 6.0 (53.5) |
| I/σ                            | 16.2 (4.3) | 29.2 (3.1) |
| Refinement                     |         |            |
| Rwork (%)                      | 18.8    | 19.2       |
| Rfree (%)                      | 23.2    | 23.0       |
| RMSD                           |         |            |
| Bond lengths (Å)               | 0.006   | 0.006      |
| Bond angles (°)                | 0.892   | 0.909      |
| Average B factor               | 44.6    | 28.7       |
| Ramachandran plot quality      |         |            |
| Most favored (%)               | 88.9    | 90.2       |
| Disallowed (%)                 | 0       | 0          |

*Values in parentheses are given for the highest-resolution shell.

Overall structure of bovine MHC I N*01801-peptide complexes. The bovine MHC I N*01801 complexed with peptide IPA was crystallized in the P₂₁₂₁₂₁ orthorhombic space group with a resolution of 2.7 Å (Table 2). The overall structure of N*01801 consists of the characteristic α₁ and α₂ domains of a heavy chain underpinned by α₃ domains and βₐm (Fig. 2A).

The peptide lies along the peptide binding groove formed by the α₁ and α₂ domains (Fig. 2B). Within one asymmetric unit, there are two N*01801 molecules (referred to below as molecule 1 and molecule 2) that have similar overall conformations, with a root mean square difference (RMSD) of 0.300 Å (0.267 and 0.137 Å for the heavy chain and βₐm, respectively) (Fig. 2C). The overall structure of N*01801 is similar to those of other class I MHC molecules. Structural alignment of molecule 1 of N*01801 with the structures of the bovine MHC I molecule N*01301 and human MHC HLA-B*5101 yielded RMSD values of 0.563 and 0.533 Å, respectively (Fig. 2D and E). Among all human MHC I molecules, HLA-B*5101 is sequentially the most similar to N*01801, displaying 80.29% identity (Fig. 3). Notably, this is at the same level as the identity between the intraspecies alleles of cattle: N*01801 and N*01301 (82.9%).

Due to the high cost of cattle for experimental use, it is necessary to investigate the possibility of utilizing mice as a model (e.g., MHC heavy-chain-transgenic mice) with which to study bovine-specific cellular immunity. To elucidate whether the peptide-presenting features of a bovine MHC I molecule can be influenced by binding to murine βₐm, we solved the structure of the N*01801 heavy chain complexed with murine βₐm and peptide IPA (referred to as N*01801-Mₐ below to distinguish it from the above-mentioned bovine βₐm complex) at a resolution of 1.9 Å. There is only one molecule in an asymmetric unit. The α₁ and α₂ domains, which form the peptide binding groove of N*01801-Mₐ, are quite similar to the corresponding portions of N*01801, with RMSDs of 0.337 and 0.345 Å to molecule 1 and molecule 2, respectively (Fig. 4). This may indicate that the structure of the peptide loaded in the groove formed by the α₁ and α₂ domains of N*01801 was not affected by the substitution of the βₐm subunit. In the consideration of a higher resolution of N*01801-Mₐ (with murine βₐm), we use this structure together with the structure of N*01801 (with bovine βₐm) to analyze peptide presentation by bovine MHC I.

Two distinct conformations of peptide IPA presented by bovine MHC I N*01801. In the BoLA binding grooves of molecule 1 and molecule 2, the electron densities around the IPA peptides are well defined and clearly reveal two distinct

![Fig. 2. Overview of the structure of bovine MHC I N*01801.](http://jvi.asm.org/Downloaded from)
conformations of the peptide. For simplicity, the peptides presented by molecule 1 and molecule 2 are referred to as IPA-M1 and IPA-M2 below; they are shown in Fig. 5A and B, respectively. IPA-M1 and IPA-M2 have an RMSD of 1.284 Å. However, the conformation of IPA-M2 is quite similar to that of the IPA peptide in the N*01801-M/H9252 structure, with an RMSD of 0.474. Furthermore, the interactions, including the hydrogen bonds and the nonpolar contacts, of IPA-M2 in the structure of N*01801 are also similar to those of the IPA peptide in the N*01801-M* structure. The IPA peptide pre-

FIG. 3. Structure-based sequence alignment of N*01801 and eight different types of MHC I molecules. Cylinders indicate α-helices, and black arrows indicate β-strands. Residues highlighted in red are completely conserved, and boxed residues are highly (>80%) conserved. Residues that play a critical role in peptide presentation are asterisked (residue 73, highlighted in yellow, and residue 167, highlighted in cyan). The sequence alignment was generated with Clustal X (56) and ESPript (24).
between the peptide conformations of molecule 1 and that of N*01801. Thus, we describe only the differences between the two distinct conformations of the IPA peptide. Indeed, residues at position 73 that locate in the flanking side of the peptide binding groove are a key factor for the peptide conformation in other MHC I molecules (11, 34). For different MHC molecules, residues have diverse polymorphism in position 73 (Fig. 3). Therefore, the residues in this position have different impacts on peptide presentation. In a recently solved structure of the bovine MHC I N*01301 molecule, peptide Tp1214–224 displays an unusual bulged C-terminal conformation, which is affected byTp73 beneath the peptide (34). Another factor may also influence the flexibility of the IPA peptide. Both of the bovine MHCs possess E97 in the bottom of the peptide binding groove. While the secondary anchor residue of K5 of Tp1214–224 forms a stable salt bridge with E97 (34), IPA does not interact with E97 due to the hydrophobicity of the middle residues of the peptide (Gly5, Val6, and Leu7). Thus, the lack of a salt bridge may partially contribute to the flexibility of IPA.

Different exposed areas and buried residues formed by the two IPA conformations. The two distinct conformations of IPA in the two molecules of the asymmetric unit of the bovine MHC I N*01801 structures may contribute to the exposure of different surfaces of the peptides to the solvent (Fig. 6). With the “double-M” conformation, IPA-Mβ protrudes from the side chains of all of the even-numbered residues (Tyr4, Val6, and Thr8) of the C-terminal portion of the peptide (P3 to P9), which consecutively presents their side chains for potential TCR docking (Fig. 6A). The side chain of Ile73 protrudes into the solvent, leaving enough space for the exposure of the middle region of IPA-Mβ (Fig. 6C). Calculation of the accessible areas shows that the exposed surface of the three residues occupies most (87.7%) of the total exposed area of IPA-M1. The odd-numbered residues in this portion of the peptide (Ala3, Gly5, Leu7, and Ile9) bury themselves deeply into the groove. In the structure of N*01801-Mβ, P4 Tyr, P7 Leu, and P8 Thr are the most prominent residues and are likely important for TCR interaction (Fig. 6B). The exposed surface of these three residues (Tyr4, Leu7, and Thr8) accounts for 81.0% of the total exposed surface of IPA-Mβ (Fig. 6D). In addition, the main chains of P5 Gly and P6 Val also partially contribute to the exposed area of IPA-Mβ. However, the side chain of Ile73 extends over the middle region of IPA-Mβ, which may affect the direct docking of the TCR to the main chain of P5 Gly and P6 Val. These distinct characteristics of the exposed area of the peptide presented by N*01801 may be involved in recognition by a greater range of the TCR repertoire.
Peptide presentation characteristics of bovine N*01801 compared to those of other MHC I molecules. Superposition of the peptide presented by bovine MHC I N*01801 with N*01301 and MHC I molecules from other species revealed key characteristics of N*01801-restricted peptide presentation. Figure 7A shows the Cα backbone of IPA-M1 with other MHC I-presented nonameric peptides (excluding the 13-mer peptide presented by N*01301). Compared to the nonamer presented by HLA-B*5101, the main chain of the N-terminal portion (P1 to P4) of the peptide IPA has a similar conformation. In P5 of the nonamer, valine acts as a secondary anchor residue with its side chain inserted deep into the groove of HLA-B*5101. In N*01801, P5 is a glycine and thus lacks a side chain. However, the highly buried area of Gly5 indicates that this residue also acts as a secondary anchor for the peptide N*01801. Together with Leu7, whose side chain protrudes into the E pocket of the groove, we can conclude that peptide IPA contains two secondary anchor residues including Gly5 in the C-terminal portion of the IPA-M1 peptide (P4 to Pc). This “double-M” conformation is seldom observed among the nonameric peptides presented by MHC I molecules. The IPA-M2 (Fig. 7B) and IPA-Mβ (Fig. 7C) peptides have similar “M-shaped” conformations, which are common among peptides presented by other MHC molecules. The peptides within other MHC grooves possess either the M-shaped conformation with one secondary anchor residue (HLA-B*5101, HLA-A*0201, Mamu-A*01, and H-2Kb) or a bulged conformation without any secondary anchor in the P5-to-Pc region (for HLA-A*2402 and N*01301).

Another characteristic resulting from the “double-M” conformation of peptide IPA-M1 is that the main chain of the peptide is deeply located in the groove of N*01801. The Cα backbone of the entire chain of the peptide lies deeper in the peptide binding groove than that in other MHC I molecules. Scrutinizing the available MHC I structures, we found that the main chain of the nonameric peptide presented by H-2Kb is also buried deep in the groove. However, as shown in Fig. 7A, the Cα atoms of the most exposed residues (Tyr4 and Val6) of IPA-M1 are still below the Cα of P5 of the nonamer presented by H-2Kb. In the other conformation of peptide IPA (IPA-M1 and IPA-Mβ), the entire peptide is also located deep in the groove (Fig. 7B).
Moreover, the highest Cα atom in the backbone of the peptide is still lower than the Cα atom of P5 of the nonamer presented by H-2Kb. The buried main chain and exposed side chains of the middle residues of IPA may dictate the specific TCR recognition of the peptide presented by N*01801.

Several factors may contribute to the low positioning of IPA in both of the conformations. First, the shorter residue Ile73 may allow the portion near the C terminus of the peptide to fall into the binding groove. In contrast, Trp73 of BoLA-A18, which forms a bulged ridge in the groove, pushes the peptide up into the solvent (34). Second, the large E pocket of N*01801 may allow the residues in the middle of the peptide to be inserted deeply into the groove (Val6 for IPA-M1; Leu7 for IPA-M2 and IPA-M3).

Pockets in the N*01801 peptide binding groove. Residues Tyr59, Asn63, Tyr159, Arg163, Gly167, and Tyr171 form the A pocket of the peptide binding groove of N*01801 (Fig. 8A). The P1 Ile of the peptide is located in the A pocket and points its side chain out of the groove. Considering MHC I molecules from different species (Fig. 3), Gly167 of N*01801 is rare among all of the MHC I molecules (i.e., it is found only in certain alleles, such as HLA-A*2402). In the N*01801 structure, the A pocket has the small residue Gly in position 167, making the space in the A pocket larger (Fig. 8B and C), which is also seen in the structure of HLA-A*2402 (Fig. 8D) (32). The side chain of Ile1 of the IPA-M1 peptide in molecule 1 points toward the N terminus (or toward the right side if observed from the 2 domain of the MHC). The different conformation of the side chain of Ile1 in molecule 2 compared to molecule 1 reveals the flexibility of Ile1 in the N*01801 structure (Fig. 5A, B, and D), which reflects the large space of the A pocket. The most common residue at position 167 in other MHC alleles (in human, monkey, mouse, chicken, and bovine BoLA-A18) is Trp (Fig. 3), which forms the right side of the A pocket (observed from the α2 domain of the MHC). Trp167 of these MHC I alleles reduces the size of the A pocket.
The structure of N*01801 (Fig. 6) reveals Pro2 as a primary anchor residue fitted into a small B pocket surrounded by Ile66, Tyr67, and Tyr8. These residues are similar to the corresponding residues (Ile66, Phe67, and Tyr8) of HLA-B*5101 (Fig. 3), which also accommodates Pro (or other small residues, such as Gly or Ala) as the P2 anchor. Structural comparison of these residues demonstrated very similar conformations, with an RMSD of 0.099 Å. This may lead to the common preference of N*01801 and HLA-B*5101 for Pro as the P2 anchor. In the F pocket of N*01801, the peptide Ile9 is surrounded by hydrophobic amino acids: Ala81, Tyr84, Phe95, Phe116, and Tyr147 (Fig. 6; see also Fig. S1 in the supplemental material). However, the space in the F pocket is larger than the side chain of Ile9, which indicates that the F pocket may be able to accommodate larger residues. Aside from small aliphatic residues (e.g., Val, Ile, and Leu), in previous studies on the peptide motif of N*01801, aromatic residues (e.g., Tyr) were also defined as the C-terminal residues of N*01801-restricted peptides. The structure of N*01801 complexed with IPA clearly confirmed the peptide motifs of N*01801 that were defined by pool sequencing of eluted peptides in previous reports (14, 27, 51).

**DISCUSSION**

To determine the structural characteristics of peptide binding and presentation by bovine MHC I N*01801 (BoLA-A11), we crystallized N*01801 in complex with a CTL epitope from rinderpest virus. Additionally, we also determined the structure of N*01801 complexed with the same peptide but with murine H-2Kb instead of bovine H-2Kb. Interspecies MHC complexes have been widely exploited in structural studies of MHC molecules. The H-2Kb does not directly interact with the peptide and has no observable effect on peptide binding and presentation (1, 10, 43, 50). Here, the peptide binding grooves formed by the N*01801 α1α2 domains were similar in the two structures determined (Fig. 4). The interspecies structure may also veritably reflect the peptide binding of N*01801 (at a higher resolution) in addition to the structure of N*01801 complexed with bovine H-2Kb.

As revealed by peptide elution from an N*01801-transfected

![Diagram of peptide binding](http://jvi.asm.org/Downloaded from)
cell line, nonameric peptides account for most of the naturally separated peptides (28). Proline is found almost exclusively at the P2 position of these peptides, and as shown in the N*01801 structure, the relatively small, bowl-shaped, hydrophobic B pocket perfectly accommodates a proline at this position. The B pocket of N*01801 is formed by residues similar to those in the B pocket of HLA-B*5101, which may explain the same preference for proline as the prevalent anchor residue in the P2 position. In the C-terminal position of the IPA peptide, isoleucine and valine act as the primary anchor residues, inserting into the hydrophobic F pocket of N*01801, as indicated in the structure of N*01801 and the refolding assays. The space within the F pocket appears larger than the side chain of isoleucine, which may indicate that larger hydrophobic and even aromatic side chains can be accommodated. This is coincident with the peptide motif studies of N*01801, which indicated that Ile/Val and a small proportion of Leu/Tyr occupied the C termini of the naturally processed peptides of N*01801 (28).

Studies of CD8+ T-cell epitopes indicate that different peptides elicit diverse T-cell responses, corresponding to distinct T-cell repertoires. Recent studies have also demonstrated that even minor modification of an epitope can lead to a profound effect on the antigenicity of the peptide (37, 58). Peptides with a relatively rigid conformation may result in limited types of TCR docking, leading to a specific T-cell repertoire with limited diversity. Recently, this has been thoroughly investigated by Macdonald and colleagues in structural and function studies of bovine MHC I N*01301 (34). Hence, a peptide with a flexible structure presented in the binding groove of an MHC molecule may lead to a diverse TCR profile. In a number of reported cases, a broader T-cell repertoire for the pathogen-specific epitope may facilitate a more effective host immune response against the invading pathogen and may prevent the emergence of immune escape mutants (39, 44, 52). The IPA peptide in our structure unexpectedly appears to be presented by N*01801 in two distinct conformations. The completely different exposed areas in the middle of the peptide may indicate different manners of TCR docking. Further study should focus on whether the two distinct conformations of the IPA peptide correspond to different T-cell repertoires. The contribution of the uncommon peptide presentation strategy of N*01801 to the CTL-specific responses of N*01801 + cattle to pathogens also needs further exploration.

Generally, a featured peptide contains exposed residues protruding out of the landscape of the MHC complexed to peptide, and this can be implemented via two different strategies. First, epitopes with characteristic long side chain residues, which are solvent exposed, may act as featured peptides to elicit diverse T-cell repertoires (58). Second, moderately bulged peptides help the short side chains rise to a suitable level for TCR docking (32). In the structure of bovine MHC I N*01801 complexed with IPA, N*01801 presents the same peptide in two distinct conformations, which is associated with the flexibility of Ile73 in the peptide binding groove. Different exposed residues and secondary anchor residues result from these conformational changes. This adjustable strategy of host antigen presentation may represent a novel type of featured peptide and may expand our understanding of the crucial role of T-cell epitopes in antipathogen immunity.

BoLA-A11 is one of the most common global haplotypes in dairy cattle. Studies of T-cell-specific responses against pathogens have identified a large number of CD8+ T-cell epitopes with the BoLA-A11 restriction. Here, we demonstrated the distinctive characteristics of peptide presentation by BoLA-A11 through the structural determination of bovine MHC I N*01801 (one of the most common alleles of BoLA-A11 haplotypes) complexed with a rinderpest virus-derived IPA peptide. Our study may lead to further definition of featured T-cell epitopes in a structural manner and, moreover, may pave the way for rational CTL-based design of vaccines for cattle or other species.

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