A Naturally Selected Dimorphism within the HLA-B44 Supertype Alters Class I Structure, Peptide Repertoire, and T Cell Recognition

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Abstract
HLA-B*4402 and B*4403 are naturally occurring MHC class I alleles that are both found at a high frequency in all human populations, and yet they only differ by one residue on the α2 helix (B*4402 Asp156→B*4403 Leu156). CTLs discriminate between HLA-B*4402 and B*4403, and these allotypes stimulate strong mutual allogeneic responses reflecting their known barrier to hematopoietic stem cell transplantation. Although HLA-B*4402 and B*4403 share >95% of their peptide repertoire, B*4403 presents more unique peptides than B*4402, consistent with the stronger T cell alloreactivity observed toward B*4403 compared with B*4402. Crystal structures of B*4402 and B*4403 show how the polymorphism at position 156 is completely buried and yet alters both the peptide and the heavy chain conformation, relaxing ligand selection by B*4403 compared with B*4402. Thus, the polymorphism between HLA-B*4402 and B*4403 modifies both peptide repertoire and T cell recognition, and is reflected in the paradoxically powerful alloreactivity that occurs across this “minimal” mismatch. The findings suggest that these closely related class I genes are maintained in diverse human populations through their differential impact on the selection of peptide ligands and the T cell repertoire.

Key words: class I histocompatibility molecules • antigen presentation • crystallography • X-ray diffraction • graft rejection • polymorphism

Introduction
HLA class I molecules are characterized by a high level of polymorphism presumed to reflect natural selection for protective immunity against microbes (1–3). HLA alleles can differ from each other by only a single amino acid (“micropolymorphism”) or by >30 amino acids (4). It has been suggested that there are nine major HLA class I “supertypes,” or clusters of alleles, that each possess a unique broad specificity for common anchor motifs in antigenic peptides (5). Alleles from each of these supertypic families are distributed in virtually all human populations and account

Abbreviations used in this paper: H-bond, hydrogen bond; MALDI-TOF, matrix-assisted laser desorption/ionization time-of-flight; MS, mass spectrometry; RP, reverse phase; v.d.w., van der Waals.
for the majority HLA-A and -B polymorphism, suggesting that their maintenance reflects an adaptable strategy for induction of immunity toward diverse microbial ligands. 

The widespread distribution of many low frequency HLA polymorphisms is probably an artifact of human migration and admixture of population groups that originally possessed a limited degree of HLA polymorphism (2). This might explain why most human populations often have only one or two alleles from each of the different HLA class I supertypes (5), such that a complete picture of HLA diversity is only appreciated when multiple racial groups are examined. An exception to this notion is the HLA-B44 family of alleles, in which both B*4402 and B*4403 are present in most human populations at combined phenotypic frequencies of up to 40% (6). HLA-B*4402 and B*4403 are the two most common members of the HLA-B44 supertype and participate in antiviral (7–10), antitumor (11–13), and minor Ag-specific responses (14, 15). HLA-B*4402 (Asp156) and B*4403 (Leu156) differ by only a single amino acid located on the α2 helix, essentially constituting a dimorphism within the HLA-B44 family because the other HLA-B44 alleles are much less common (generally, gene frequency <0.01; reference 6). These two B44 allotypes are also associated with different ancestral MHC haplotypes, suggesting an ancient origin for this di-allelism (16, 17).

Table I. Sequences of Individually Sequenced Peptides

| Peptide sequence | Source | Mass | Frac. | B*4402 | B*4403 |
|------------------|--------|------|-------|--------|--------|
| **Shared ligands** | | | | | |
| EEFGRAFSF | HLA DPA*0201 | 1,088.5 | 34 | ++++ | ++++ |
| AEDKENYKKF | Hsp90α (428–437) | 1,270.6 | 33 | ++ | + |
| AEMGKGSFKY | Translation elongation factor 1α (47-56) | 1,116.5 | 27 | ++ | + |
| EEYNGLVTV | Ig heavy chain (106-115) | 1,079.6 | 31 | +++ | + |
| FEQIERF | Human EST | 1,080.5 | 31 | ++ | + |
| EEPTVIKKY | Human EST | 1,105.6 | 28 | ++ | + |
| DEVGIVTKY | Unknown | 1,152.6 | 28 | ++ | + |
| EEVDLSKDIQHW | Ribonucleoside diphosphate reductase M2 chain | 1,497.7 | 32 | ++ | + |

| **Ligands preferentially bound to B*4403** | | | | | |
| AETPDIKLF | Ribosomal protein s5 | 1,032.5 | 33 | + | +++ |
| KEIKDILIQY | Ribosomal protein s12 | 1,261.7 | 32 | – | +++ |
| AEVDKVTGRF | Ribosomal protein s21 | 1,120.6 | 26 | – | +++ |
| NELNIIHKF | Human EST | 1,126.6 | 32 | + | + |

| **Ligands preferentially bound to B*4402** | | | | | |
| EEVHDLERKY | Human EST | 1,316.6 | 24 | +++ | + |
| YEGLLDYW | Ig heavy chain (86-93) | 1,057.6 | 31 | +++ | + |

Members of the B44 supertype share a preference for peptides with negatively charged residues at P2 (Glu) and hydrophobic residues at the COOH terminus (18, 19). Previous comparisons of ligands eluted from B*4402 and B*4403 molecules have not revealed any difference in their peptide repertoire, raising the question of what functional differences between these alleles support their high prevalence in all human populations.

Despite the similarity of their peptide repertoires, mismatching of B*4402 and B*4403 in hemopoietic stem cell transplantation has been associated with transplant rejection (19, 20) and acute graft versus host disease (21), indicating that they represent a significant barrier to clinical transplantation. Moreover, CTL can discriminate between B*4402 and B*4403 allotypes, indicating structural differences that are highly relevant to T cells (22–24). The similarity in peptide repertoires of B*4402 and B*4403 suggests that the strong alloresponses between these alleles do not reflect presentation of different peptides (18, 19), but may reflect the accessibility of the α2 domain residue 156 for direct interaction with the TcR (24). Moreover, analyses of HLA-A2 mutants have confirmed the strong influence of position 156 on T cell recognition (25) and suggest a limited influence for this residue on peptide binding (26). On the other hand, residue 156 has also been predicted to influ-
ence the D/E pockets (24) that interact with the central core of the bound peptide (P4-P7; reference 27).

To help resolve these issues, we have evaluated the structural and functional properties of HLA-B*4402 and B*4403.

**Materials and Methods**

**Cell Lines and Methods.** The B-lymphoblastoid cell line Hmy2.C1R (28) is class I–deficient, having no detectable HLA-A, very low levels of HLA-B35, normal HLA-Cw4, and intact antigen processing and presentation pathways (29). Transfection of HLA-B*4402 and B*4403 into C1R (C1R.B*4402 and C1R.B*4403, respectively) and cell culture conditions have been described previously (29). LC-13 and DD1 CTL clones were maintained in RPMI 1640 with supplements and restimulated every 10–14 d in the presence of recombinant IL–2 and HLA-matched lymphoblastoid cell lines or PBMCs pulsed with the appropriate peptide antigen. Derivation of CD8+ CTL clone 5101.1999.23 has been described previously (30).

**Allogeneic MLR and Intracellular Cytokine Staining.** MLRs were performed using PBMCs from healthy blood donors (n = 13) matched for HLA-A, -B, and sometimes -C, but mismatched for HLA-B*4402 and B*4403 (and HLA class II). The HLA class I typing was as follows: B*4402 group (n = 10: A*02011/9, 03011; B*44021, 5701; and C*0501/02/03, 0602/03/07) and B*4403 group (n = 3: A*02011/9, 3011; B*44031, 5701; and C*0602/03/07, 1601).

In brief, 106 responder and 5 × 105 irradiated (3,000 rad) stimulator cells were cultured in RPMI 1640 plus 10% fetal calf serum, supplemented with 10 U/ml of recombinant IL–2 (Cetus Corporation) for 13 d at 37°C. On day 13, 2 × 105 responder T cells were harvested and restimulated with a panel of APCs (C1R, C1R.B*4402, and C1R.B*4403) at a cell concentration of 105 for 2 h at 37°C, 5% CO2. 10 µg/ml brefeldin was added for an additional 4 h, and responder T cells were stained with anti-CD4 PE (clone SK3; Becton Dickinson) and anti-CD8 CyChrome (BD Biosciences). Cells were fixed with 0.3% saponin (Sigma-Aldrich), permeabilized with 0.3% saponin (Sigma-Aldrich), and intracellular IFN-γ was detected with an anti–IFN-γ mAb (clone 25723.11; Becton Dickinson). The percentage of CD8+ T cells producing IFN-γ was determined by flow cytometry using Flowjo software (Tree Star Inc.).

**Purification of Cell Surface–Associated HLA–B44 Complexes and Peptide Analysis.** Purification of HLA-B*4402 and B*4403 was performed from 5 × 106 C1R.B4402 and C1R.B4403 cells grown in roller bottles as described previously (31). Peptides were recovered as described previously (31). Peptides were separated by reverse phase (RP)-HPLC using a SMART system HPLC (Amersham Biosciences) with a µRPC C2 /C18 column (2.1 mm [inside diameter] × 10 cm). Eluted peptides were resolved from contaminating detergent polymers by using a rapid gradient from 0 to 60% acetonitrile in 0.1% aqueous TFA (12% increase in buffer B (organic)/min, 200 µl/min). This material was subjected to pool Edman sequencing and matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF MS).

MS. MALDI-TOF MS was performed using a Reflex mass spectrometer (Bruker) as described previously (31). Care was taken to ensure reproducibility of MS results on HPLC fractions (Figs. S1–S3 available at http://www.jem.org/cgi/content/full/jem.20030066/DC1). Peptide sequencing by Q-TOF electron-
son of the fragmentation spectra using the Mascot algorithm (33) followed by manual assignment of expected fragments from the highest score sequences (Table I).

**Crystallization, Data Collection, and Structure Determination.** Recombinant B44 complexes were prepared from *Escherichia coli*, refolded, and crystallized as described previously (34). The data were processed and scaled using the HKL package (35) and CCP4 suite (36), Table II. The HLA-B*4402 structure was solved first using the molecular replacement method, using programs from CCP4 suite (37). The progress of refinement was monitored by the $R_{	ext{free}}$ value (3% of the data) with neither a $R_{	ext{free}}$ nor a low $R_{	ext{free}}$ value (3% of the data) with neither a $R_{	ext{free}}$ nor a low $R_{	ext{free}}$ value (3% of the data) with neither a $R_{	ext{free}}$ nor a low $R_{	ext{free}}$.

Figure 1. Differential T cell recognition of HLA-B*4402 and HLA-B*4403 by CD8+ T cells. CTL clones, LC13 (A), and DD1 (B) lyse HLA-B*4402 but not HLA-B*4403 targets; however, transfect C1R (closed triangles); and C1R-transfected APCs expressing either B*4402 (open squares), B*4403 (open circles), B*4405 (closed circles), or B*4407 (closed squares). (C) The CTL clone 5101.1999.23 is alloreactive with HLA-B*4402 and HLA-B*4403. IFN-γ production was assayed after 4 h. APCs: HERLUFF, B*4403 + LCL; P1OTUK, B*4402 + LCL; and auto HSV-2, autologous cells infected with HSV-2. (D) PBMCs from donors mismatched for B*4402 and B*4403 were cocultured in a 13-d MLR supplemented with IL-2. The specificity of the responding cells was evaluated by restimulation for 6 h with defined APCs. Vertical axis, IFN-γ staining; horizontal axis, CD8 staining. Gates are shown for IFN-γ-producing CD8+ T cells. The percentage of positive cells is shown in each histogram. (E) The percentage of allospecific CD8+ T cells was assayed as in Fig. 4 D for 28 independent MLR reactions between B*4403 responders and B*4402 stimulators and 13 independent MLR reactions between B*4402 responders and B*4403 stimulators.

**Results**

T Cell Discrimination between HLA-B*4402 and B*4403. Despite only a single amino acid difference in the primary structure of HLA-B*4402 (156Asp) and B*4403 (156Leu), previous papers have shown that CD8+ T cells can readily distinguish between these two molecules (22, 40, 41). For example, allospecific CD8+ CTL clones, LC13 and DD1 (42), kill C1R transfecants expressing HLA-B*4402 and B*4405 but not transfecants expressing HLA-B*4403 (Fig. 1, A and B). On the other hand, an allospecific, anti-B*4402 CTL clone (30, 43) makes IFN-γ in response to C1R-transfectants expressing HLA-B*4402, B*4403, and B*4407 but not B*4405 (Fig. 1 C). Accordingly, allogenic T cell reactivity can be used to probe structural differences in MHC allotypes that may influence thymic selection (44) and peripheral diversity of T cells in immunity (3).

Therefore, we evaluated the in vitro alloresponse between individuals mismatched for B*4402 and B*4403, but matched at both HLA-A loci and for the HLA-B trans al-
lele. After reciprocal allogeneic MLRs, CD8+ T lymphocytes were restimulated with defined APCs to determine their specificity, and T cells making IFN-γ were enumerated. Less than 1% of CD8+ T cells responded after the allogeneic MLR when either syngeneic PBMCs or untransfected C1R cells were used as APCs (Fig. 1 D, left). However, ~20% of CD8+ T cells from B*4402 responders were activated by C1R.B*4403 APCs (Fig. 1 D, bottom). Similarly, restimulation of B*4403+ responders with C1R.B*4402 APCs provoked IFN-γ production in ~10% of CD8+ T cells (Fig. 1 D, top).

The median percentage of alloreactive CD8+ T cells in the B*4402 anti-B*4403 vector was 11.3% (n = 28; independent donor pairs) compared with a median of 2% B*4403 anti-B*4402 CD8+ T cells (n = 13; independent donor pairs). Hence, there were nearly sixfold more responding T cells identified in MLRs from B*4402 anti-B*4403 mismatches than vice versa, indicating an asymmetry in the magnitude of the alloresponse stimulated between these donor pairs (Fig. 2).

Isolation of HLA-B44-bound Peptides and Analysis of Ligand Specificity. Differential T cell recognition of B*4402 and B*4403 could result from differences in either peptide repertoire or HLA heavy chain conformation (45, 46). Alternatively, identical peptide repertoires could be presented in an altered manner due to structural changes at the interface between the peptide loaded class I molecule and the TcR (47, 48). Therefore, peptide repertoires of HLA-B*4402 and B*4403 were examined using pool Edman sequencing and high-resolution MALDI-TOF MS. Apart from the previously reported P2Glu and P4Tyrr/Phe anchor residues (12, 18, 19), minor differences were noted between subdominant anchor residues with more pronounced yields of valine at P3 for B*4403 and tryptophan at P9 for B*4402 (Fig. 2).

When peptide pools were examined by MALDI-TOF MS, a high degree of overlap was detected in the ligands from HLA-B*4402 and B*4403 (Fig. 2), consistent with previous work (12, 18, 19). Each peptide pool exhibited a complex ionization pattern, with a common prominent signal at (M + H)+ = 1089.6, subsequently shown to be self-peptide derived from HLA DPB1a (34). Differences in ligand repertoire were evident between HLA-B*4402 and B*4403, such as the species of m/z 1033.6, 1121.5, and 1145.6 that were all more prevalent in HLA-B*4403 relative to B*4402.

Additional fractionation of the total eluates was performed using a second dimension of RP-HPLC followed by MALDI-TOF MS of each fraction. Fig. 3 A shows the complexity of ligands identified in B*4402 as a function of RP-HPLC retention time (right) and mass spectra (left). The individual species identified in ~60 RP-HPLC fractions exhibited >95% overlap between HLA-B*4402 and B*4403 (Fig. 3, B–G and unpublished data). Fig. 3 B shows the mass spectra of fraction “B” (Fig. 3 A, right), derived from HLA-B*4402 (MALDI-TOF, positive polarity), and HLA-B*4403 (negative polarity). Here, the species at m/z 1121.3 appears to be specific to the HLA-B*4403 eluate and was not detected in neighboring fractions from either HLA-B*4402 or B*4403. This peptide was identified as amino acids 37–46 of human ribosomal protein S21, a natural ligand of HLA-B*4403 (Table I; reference 18). On the other hand, the species of mass 1180.4 was preferentially recovered from B*4402 eluates. Fractions “C” and “F” (Fig. 3 A, right) revealed greater complexity of peptides bound to HLA-B*4403 versus B*4402 (Fig. 3, C and F), whereas spectra of equal complexity but distinct composition were observed in other fractions (Fig. 3, D and E). Some fractions contained ligands that were mostly identical in B*4402 and B*4403 (Fig. 3 G).

Approximately 5% of all ligands were either preferentially presented by B*4403 or B*4402, or were uniquely recovered from one or other allotype. A systematic examination of all fractions identified 116 such peptides, of which 80% were “biased” toward presentation by the B*4403 molecule. Only 20% of the biased ligands were preferentially presented by B*4402. This bias was even more marked (>90% B*4403) when the analysis was confined to
ligands with the strongest ionization signals. Sequencing of shared or unique peptides associated with HLA-B*4402 and B*4403 verified the observations made by MS (Table I). Tryptophan and valine were identified at the COOH terminus of some eluted peptides. There was no characteristic motif in the peptides preferentially associated with B*4403.

**Structural Basis of HLA-B*4402 and B*4403 Binding to a Shared Peptide Ligand.** The crystal structures of B*4402 and B*4403 were determined in complex with a shared natural, high affinity ligand derived from HLA-DPα (DPA*0201, residues 46–54 EEFGRAFSF) (Fig. 2 and Fig. 3 G, (M + H)+ = 1089.6 ± 0.2). The data collection and refinement statistics are shown in Table II. The resolution of the structures was 1.6 and 1.7 Å, respectively. The peptide was bound in an extended conformation in both HLA-B*4402 and B*4403, with a small bulge around Gly 4 (Fig. 4 A–D). There were 13 water-mediated hydrogen bonds (H-bonds) and 15 direct H-bonds in the HLA–B*4402 complex. Fewer H-bonds were observed in the HLA–B*4403 complex (9 water mediated H-bonds and 15 direct H-bonds). The peptide also participates in three conserved salt bridges in both complexes (Table S1 available at http://www.jem.org/cgi/content/full/jem.20030066/DC1). The dominant solvent-accessible residues of the peptide ligand were Glu1, Gly4, Arg5, and Ser8, whereas Phe3 and Phe7 are only partially solvent accessible (Fig. 4 D). In both structures, Arg5 was the most mobile residue.
(B factors 29–35, Å²), reflecting the greater solvent accessibility of the polar head group of this residue.

Structural Basis for Selection of Shared Anchor Residues by the B and F pockets. The antigen-binding cleft of class I HLA molecules consists of a series of pockets designated A–F (27, 49). In different alleles, these pockets vary in their depth, electrostatic potential, and hydrophobicity, thus determining their ligand specificity. Specificity for P2-Glu occurs in HLA-B*4402, HLA-A*29, and H-2Kk. However, because the structural basis for this specificity has not yet been described, these interactions are described in some detail here and compared with the B pockets of B*2705 (50) and B*3501 (51, 52) that possess alternative specificities (Fig. 5). The B pocket is formed by residues at positions 9, 24, 45, 66, 67, and 99 of the class I heavy chain; the P2-Glu interacts with all these residues except for Ile66 in our structures and makes an additional interaction (position 70). In B*4402 and B*4403, the peptide anchor Glu2 forms a salt bridge with Lys45, which is situated at the bottom of the B pocket and presumably provides substantial energetic con-

Figure 4. Structures of bound DPα ligand complexed to two HLA-B44 alleles, highlighting polymorphic amino acids and the electron density of the peptide. The structures of the DPα(EEFGRASF) complexed to (A) HLA-B*4402 and (B) B*4403 highlighting the polymorphism at position 156. (C) Superposition of the two antigen-binding clefts B*4402 (green) and B*4403(yellow) showing the shift in the α1 helix around residues 70–77. (D) Superposition of the two peptide ligands (side view) highlighting the solvent-accessible amino acids.

Figure 5. Novel structure of the B pocket of HLA-B*4403 and B*4402 determines the selection of dominant anchor residues P2 Asp/Glu. The structure of the B pocket of HLA-B*4403 is compared with previously reported structures for HLA-B*2705 and HLA-B*3501. The upper panels are ball and stick representations of B pocket amino acids 9, 24, 45, 66, 67, 70, and 99 (gray) and the P2 amino acid of the ligand (green); B*4403 (P2 Glu, left), HLA-B*2705 (P2 Arg, middle), and B*3501 (P2 Pro, right). The lower panels represent electrostatic surfaces of the area bounding the B pocket using the program GRASP (70) and showing the bound peptide ligand for the structures in ball and stick format (blue, electropositive; red, electronegative). The B pocket of B*4402 is virtually identical to B*4403.
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distribution to the specificity of the B pocket (Fig. 5). Furthermore, the charged P2-Glu carboxylate forms direct H-bonds to Tyr9 and Tyr99. Additionally, there is a buried, well-ordered water molecule at the base of the pocket that mediates H-bonds to Ser67 and Thr24 in the HLA-B*4402/03 structures. In B*4403, this conserved water molecule forms a H-bond to Asn70, a residue within the α1-helix that flexes as a result of the polymorphism at position 156. The long aliphatic moiety of Glu2 packs predominantly against the aromatic ring of Tyr7 and is flanked by Ile66, although it doesn’t interact with the latter residue.

Some of the structural framework of the B pocket is conserved between B*4403, B*2705, and B*3501 (Fig. 5). Thus, the B pockets of B*2705 (50) and B*4402/B*4403 are deep to accommodate the long aliphatic side chains of arginine or glutamate. However, in contrast to B*4402 and B*4403, the deep electronegative B pocket of B*2705 (50) accommodates a positively charged canonical P2-Arg residue involved in charge complementation with the glutamic acid residue at position 45 of the heavy chain. In the B pocket of HLA-B*3501, the projecting Phe67 creates a hydrophobic, sterically occluded pocket that preferentially accommodates proline residues at P2 of bound peptide ligands (51, 52).

The residues that surround the F pocket are very similar in both the B*4402 and B*4403 structures, and only the structure of B*4402 is shown in Fig. 6. The P9 anchor residue phenylalanine projects into a hydrophobic F pocket, where it stacks between the aromatic ring of Tyr123 and the aliphatic moiety of Asn77. The main chain of this COOH-terminal residue is tethered by H-bonds to Asn77, Tyr84, and Thr143. In addition, the walls of the F pocket are bound by Trp147, Tyr74, and Ile95. The large cluster of aromatic residues within the F pocket provides a structural basis for the preference of aromatic residues at the P9 position. Asp116 forms the base of this pocket, where the carboxylate group of Asp116 is oriented away from the F pocket, leaving the aliphatic moiety of Asp116 to interact with the hydrophobic Phe side chain (Table S1 and Fig. 6) and to form a direct H-bond with Tyr74 that is lost in B*4403.

Impact of α2 Helix Position 156Asp→Leu Polymorphism on HLA-B*4402 and B*4403 Structures. In the structures of both B*4402 and B*4403, the side chain of amino acid 156 was buried and, therefore, not directly accessible for TcR interaction (Fig. 4, A and B). Superposition of the Cα backbones of peptides bound to B*4402 and B*4403 revealed a root mean square deviation of 0.41 Å with a 1.0-Å shift around Ala6 being the most significant movement between the two structures (Fig. 4 C). In HLA-B*4402, Asp156 forms a salt bridge with Arg97 on the floor of the cleft and participates in a number of H-bond and van der Waals (v.d.w.) interactions. These interactions include water-mediated H-bonds to the Arg97 guanadinium head group and the C5 carbonyl of the peptide ligand backbone, and an unusual direct H-bond to Asp114 (Fig. 7 A). The H-bond between Asp156 and Asp114 orients the

Figure 6. The F pocket of B*4402. The main chain of the peptide carboxy-terminal residue is tethered by H-bonds to Asn77, Tyr84, and Thr143. The P9 anchor residue phenylalanine projects into a hydrophobic F pocket where it stacks between the aromatic ring of Tyr123 and the aliphatic moiety of Asn77. The walls of the pocket are bound by Trp147, Tyr74, and Ile95. Peptide residues in green, HLA heavy chain in gray and H-bonds shown as dashed lines.

Figure 7. The structural interactions around residue 156 constrain the antigen-binding cleft of HLA-B*4402 (156 Asp) relative to HLA-B*4403 (156 Leu) and alter the orientation of peptide side chains. The side chain of residue 156 is shown in yellow and forms a unique H-bond with Asp114 in B*4402 (A) that is missing in B*4403 (B), where instead Asp114 is reoriented to maximize its H-bond network with Arg97, creating a wider cavity in this region of the cleft. Peptide, green; HLA heavy chain, gray; H-bonds, dotted lines; W, water.
Asp114 side chain, allowing a salt bridge to the guanidinium head group of Arg97 as well as a water-mediated H-bond to Asp116. The pairing of Asp156 and Asp114 has not been seen in any other crystal structures of HLA class I molecules, and is presumably stabilized by salt bridging between both these residues and Arg97, allowing an O–H⋯H interaction between carboxylates. However, the potential for H-bond formation between Asp156 and Asp114 exists in several other HLA-B molecules, including other members of the HLA-B44 family. In addition Asp156 forms a v.d.w. interaction with Trp133 and Leu126, as well as with Phe3 of the bound peptide ligand. Here, the peptide Phe3–Asp156 interaction is facilitated by the side-on orientation adopted by the Phe3 side chain.

In contrast to B*4402, the nonpolar nature of Leu156 in B*4403 results in loss of a water-mediated H-bond to the peptide backbone, giving rise to a significant repositioning of the peptide between the B*4403 α-helices. On the other hand, the side chain of Leu156 in HLA-B*4403 is rotated by ~90° (Fig. 7 B), enhancing v.d.w. contacts within the hydrophobic pocket formed by Leu126, Trp133, and Val152. In addition, to avoid unfavorable interactions with Leu156, the orientation of the Asp114 side chain in B*4403 changes by ~90°, simultaneously maximizing interactions with Arg97 and maintaining a water-mediated H-bond to the C5 carbonyl of the peptide backbone. Hence, despite the loss of some H-bonds in B*4403, including the Asp116–Tyr74 interaction, the stability of the EEFGRAFSF peptide does not seem to be significantly impaired in the B*4403 complex consistent with experiments that show little difference in the thermostability of B*4402 versus B*4403 in complex with the EEFGRAFSF peptide (unpublished data).

However, the altered interactions between residue 156 and neighboring heavy chain residues, and the peptide ligand, significantly change the dimensions of the Ag-binding cleft in B*4402 and B*4403 (Fig. 4, C and D). A "rigid body shift" in the α1 helix, particularly around positions 70–77, results in a 0.6-Å displacement of the α-helices between the antigen-binding cleft of HLA-B*4402 (green) versus B*4403 (Fig. 4 C, yellow). Amino acid residues 70–77 lie adjacent to the central region of the peptide that has undergone a shift of 1.0 Å in the Cα backbone around Ala6. This conformational adjustment is directly associated with the loss of a direct H-bond between Asp116 and Tyr74 of the B*4403 heavy chain (Fig. 7 A and B). Thus, in the HLA-B*4402 structure the direct H-bond between Asp116 and Tyr74 results in a narrower antigen-binding cleft (Figs. 6 and 7 A), whereas in the B*4403 structure loss of this interaction is associated with distortion and relative "breathing" of the binding cleft (Fig. 7 B). Instead of making a direct H-bond with Asp116, Tyr74 participates in a water-mediated H-bond with this residue in HLA-B*4403; this is attributable to changes in the peptide orientation, resulting from the substitution at amino acid 156. Consequently, the slightly larger cavity of the HLA-B*4403 cleft is occupied by an acetate ion in this structure, reflecting potentially greater accommodation of peptide side chains by the B*4403 allotype compared with B*4402 (Fig. 7 B). Any thermodynamic penalty for “breathing” of the B*4403 cleft may not be apparent with some ligands because of compensatory peptide interactions arising from the altered cleft structure, as appears to be the case for B*4403 binding of the EEFGRAFSF peptide.

Discussion

The impact of point mutations and other micropolymorphisms on the structure and function of class I molecules has been studied extensively by in vitro mutagenesis (53, 54) and through exploitation of the experimentally derived Kb mutant system (3, 44, 47, 48). These systems have been enormously important for unraveling key details of structure–function relationships in MHC molecules. However, it is equally important to study the structural and functional consequences of naturally selected microvariation in HLA allotypes found in outbred human populations. To this end, we have determined the structural and functional properties of two closely related, naturally occurring human class I alleles, HLA-B*4402 and B*4403. These were studied because of their evolutionary success as common alleles in diverse human populations, notwithstanding their marked structural similarity and shared peptide binding motif.

Structural differences in B*4402 and B*4403 are distinguishable by many T cells and these allotypes provoke strong, mutual allogeneic T cell responses in vitro. Indeed, up to 25% of CD8+ T cells are specifically alloreactive in extended MLRs between B*4402/B*4403 mismatched individuals. These observations indicate that B*4402 and B*4403 molecules select both shared and unique T cells during thymic development.

The structures of B*4402 and B*4403 show that amino acid 156 is neither solvent accessible, nor directly involved in forming major specificity pockets. Nonetheless, the 156Asp→Leu polymorphism is a nonconservative substitution and markedly alters the characteristics of the P3 specificity (E pocket) as well as the secondary effects of the antigen-binding cleft. Hence, the 156Asp→Leu substitution alters the repertoire of peptide ligands bound by both molecules and directly alters the conformation of the antigen-binding cleft. Subtle differences were observed in preferred residues at minor anchor sites in B*4403 versus HLA-B*4402, however, these differences may be more apparent than real. For instance, a Trp can be modeled at P0 in B*4402 and B*4403 (unpublished data) and a tryptophan is found at PΩ in the EEVILSDKIQHW peptide eluted from both B*4402 and 4403 (Table I). Thus, any preference for Trp at PΩ in B*4402 is probably quantitative (statistical) or context-dependent.

The peptide repertoire of HLA-B*4402 and B*4403 showed >95% overlap but most of the unique peptides were associated with B*4403. These differences in peptide selection can be partly understood by an examination of the two structures. Comparison of the antigen-binding cleft of HLA-B*4402 and B*4403 revealed a relative narrowing of the B*4402 cleft between the α-helices, particularly around
amino acids 70–77. This distortion of the cleft shape appears to be the result of both indirect and ligand-mediated conformational changes associated with the 156Asp→Leu polymorphism. The Asp114–Asp156 interaction in B∗4402 is unusual, and being constrained, appears to limit the type of residue that can sit in the P3, and/or the P7 pocket. A large network of polar interactions at position 156 dissipates this unfavorable interaction and draws the helix 1 closer to the peptide, thereby sterically constraining the makeup of potential bound peptides in B∗4402. Thus, the relatively restricted range of ligands recovered from B∗4402 compared with B∗4403 probably results from this structural constraint that limits the size of side chains that can be accommodated in the narrowed region. In contrast, the 156Asp→Leu substitution in B∗4403 results in a loss of this bonding network, allowing the helices to breathe in this region. This breathing of the cleft is further reflected by the presence of an acetate moiety in the extra space around amino acids 70–77 in the B∗4403/EEFGRAFSF complex. Breathing, or opening of the binding cleft, is also observed in the structures of HLA-DQ8 (55) and I–Ag(C) (56) when these are compared with other class II molecules.

The orientation of side chains in the EEFGRAFSF peptide bound by HLA-B∗4402 and B∗4403 was also altered by the 156Asp→Leu polymorphism. Structural adjustments within identical ligands bound by related MHC allotypes can be sufficient to induce T cell alloseactivity (24). However, altered heavy chain conformation (57) and binding of unique ligands (58) can also contribute to T cell alloseactivity. Thus, T cell discrimination of B∗4402 and B∗4403, either during positive selection of the T cell repertoire, or during B∗4402 anti-B∗4403 alloseactivity, could involve any one, or all of the modes of recognition discussed here.

Ironically, the structural similarity between B∗4402 and B∗4403 is likely to engender strong reciprocal T cell alloseactivity due to thymic selection of a large repertoire of T cells with intrinsic “B44”–reactivity biased toward these allotypes. Hence, the number of T cells that can distinguish subtle structural differences in B∗4402 and B∗4403 is likely to be very significant. Thus, the magnitude of the alloseactivity is paradoxically as great, or even greater, than observed across more disparate HLA allelic differences (19–21).

The subtle differences in peptide repertoire between B∗4402 and B∗4403 contrast with the differences observed between many HLA class I allotypes, especially where substitutions occur in the dominant anchor residues located within the B and F pockets (59–61). For instance, HLA-B∗2705 and B∗2709 differ at position 116 (Asp→His) at the base of the F pocket (62, 63), resulting in peptide repertoires that overlap by ∼79 and ∼88%, respectively (64), and producing mutual alloseactivity (62, 65).

However, natural class I polymorphism outside the P2 and P6 peptide anchor sites appears to modulate peptide repertoire in a much more subtle, often imperceptible manner, as reported for subtypes of HLA-B7, B17 and B15 (66), B5 and B22 (67), and B27 (65–69). Thus, our findings in relation to B∗4402 and B∗4403 are likely to be of general relevance in understanding the impact of microvariation in HLA class I genes. For instance, among all known HLA class I molecules, there are five different amino acids observed at position 156 (Asp, Leu, Arg, Gln, and Trp). Dimorphism at this residue is found in virtually all the common class I subtypes, suggesting that this position exerts a strong influence on the structure and function of other class I allotypes as well as B44 (25).

The functional basis underlying the retention of the B∗4402 and B∗4403 alleles in diverse human populations presumably rests with their independent selective advantage in immunity. Selection could be related to differential ligand selection by B∗4402 and B∗4403 (1); or, alternatively, may reflect differences in either thymic positive selection of T cells (44) or clonal diversification of the peripheral immune response (3). In addition, selection could be influenced by other elements of conserved HLA-B44 haplotypes, although these are not well-preserved in all populations, making this explanation less likely.

We conclude that the relatively small changes in peptide repertoire and MHC–peptide conformation between naturally occurring HLA polymorphisms can be sufficient to maintain selection of these alleles in diverse human populations. These micropolymorphisms are potentially capable of enhancing protective immunity by expanding ligand selection but are also likely to diversify T cell repertoire during thymic positive selection and at the time of the immune response.

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