Peptide binding to HLA-DP proteins at pH 5.0 and pH 7.0: a quantitative molecular docking study

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Abstract

**Background:** HLA-DPs are class II MHC proteins mediating immune responses to many diseases. Peptides bind MHC class II proteins in the acidic environment within endosomes. Acidic pH markedly elevates association rate constants but dissociation rates are almost unchanged in the pH range 5.0 – 7.0. This pH-driven effect can be explained by the protonation/deprotonation states of Histidine, whose imidazole has a pK\(_a\) of 6.0. At pH 5.0, imidazole ring is protonated, making Histidine positively charged and very hydrophilic, while at pH 7.0 imidazole is unprotonated, making Histidine less hydrophilic. We develop here a method to predict peptide binding to the four most frequent HLA-DP proteins: DP1, DP41, DP42 and DP5, using a molecular docking protocol. Dockings to virtual combinatorial peptide libraries were performed at pH 5.0 and pH 7.0.

**Results:** The X-ray structure of the peptide – HLA-DP2 protein complex was used as a starting template to model by homology the structure of the four DP proteins. The resulting models were used to produce virtual combinatorial peptide libraries constructed using the single amino acid substitution (SAAS) principle. Peptides were docked into the DP binding site using AutoDock at pH 5.0 and pH 7.0. The resulting scores were normalized and used to generate Docking Score-based Quantitative Matrices (DS-QMs). The predictive ability of these QMs was tested using an external test set of 484 known DP binders. They were also compared to existing servers for DP binding prediction. The models derived at pH 5.0 predict better than those derived at pH 7.0 and showed significantly improved predictions for three of the four DP proteins, when compared to the existing servers. They are able to recognize 50% of the known binders in the top 5% of predicted peptides.

**Conclusions:** The higher predictive ability of DS-QMs derived at pH 5.0 may be rationalized by the additional hydrogen bond formed between the backbone carbonyl oxygen belonging to the peptide position before p1 (p-1) and the protonated ε-nitrogen of His\(^{795}\). Additionally, protonated His residues are well accepted at most of the peptide binding core positions which is in a good agreement with the overall negatively charged peptide binding site of most MHC proteins.

**Background**

Major histocompatibility complex class II (MHC class II) proteins are normally found in B lymphocytes, dendritic cells, and macrophages; they are primarily involved in processing foreign, extracellular antigens, which are endocytosed and then enclosed in endosomes containing acid proteases. The pH in endosomes ranges from 4.5 to 6.0 [1]. In endosomes, antigens are degraded into oligopeptides.

MHC class II proteins are synthesized in the endoplasmic reticulum (ER) and bind to a protein known as the MHC class II-associated invariant chain (II). II facilitates the export from the ER of MHC class II proteins and prevents binding to peptides resident in the ER. The complex MHC-II enters a specific endocytic compartment, called MIIC (MHC class II compartment), which fuses with endosomes. II is cleaved initially to the so-called CLIP fragment, with CLIP later being displaced by high-affinity peptides. The peptide – MHC class II protein complex is translocated to the cell surface, where it is recognized by CD4+ T cells.
Peptides binding to MHC class II vary in length from 12 to 25 amino acids, yet the binding site accepts only nine peptide residues, the rest extending from both ends as the cleft is open-ended. The side chains of bound peptides project into several binding pockets while a system of hydrogen bonds forms between the peptide backbone and the side chain atoms of the MHC [2].

Human MHCs, known as HLA (Human Leukocyte Antigens), are extremely polymorphic and polygenic. The IMGT/HLA database lists over 1,600 HLA class II proteins [3]. HLAs class II contain three loci: DR, DQ and DP. DR and DQ proteins are well studied, while DP was initially considered of lesser importance in immune responses. However, it is now clear that HLA-DP proteins have important roles in mediating the immune response to many diseases, such as graft-versus-host (GVH) disease [4], sarcoidosis [5], juvenile chronic arthritis [6], Graves’ disease [7], hard metal lung disease [8] and especially, chronic beryllium disease [9]. Recently, a favorable DPβ side chain contacts the binding peptide in the vicinity of peptide position 2. Recently, a favorable π–π stacking between the aromatic rings of His30β and His2peptide was identified [15]. The other His residues are remote from the binding site and do not make contact with the bound peptide.

Molecular docking is a key structure-based method with significant utility in drug design, bioinformatics, and immunoinformatics. In contrast to sequence-based approaches, virtual docking experiments do not require extensive pre-existing experimental data. The only information necessary is a reliable model of the peptide – MHC protein complex, as provided by X-ray crystallography. Docking methodology allows the development of predictive models where the training and test data are fully independent, thus, eliminating any possibility of over-fitting. We use rigid docking to identify optimised bound peptide conformations; since even for a nonamer, a fully unconstrained peptide docking would be of a prohibitively extended duration. However, since the number of distinct peptide conformations observed within currently-known X-ray structures remains very small, we make the parsimonious and wholly-reasonable assumption that peptides will bind in a similar conformation. Molecular docking has been extensively and rigorously tested on both peptide-MHC class I and peptide-MHC class II complexes. As an approach to evaluating peptide binding to MHCs, it has proved to be rapid, accurate, and reliable [15-17].

Recently, we applied a molecular docking protocol to a library of modeled peptide-DP2 complexes to assess the contribution of each of the 20 naturally occurred amino acids at each of the nine binding core positions and four flanking residues (two at both ends) [15]. The normalized binding scores formed a quantitative matrix (QM), known also as a position-specific scoring matrix (PSSM). PSSMs are a commonly used representation of motifs or patterns within biological sequences. The predictive ability of the derived QM was assessed using an external test set of known binders to DP2. A comparison to predictions made by existing servers for DP2 binding prediction indicated an improvement in performance offered by our docking score-based QM (DS-QM) [15].

In the present study, we modelled by homology four of the most frequent HLA-DR proteins [18]: DP1 (DPA1*0201/DPB1*0101), DP41 (DPA1*0103/ DPB1*0401), DP42 (DPA1*0103/DPB1*0402) and DP5 (DPA1*0201/DPB1*0501). We applied a similar docking protocol to
derive DS-QMs for peptide binding prediction [15]. To investigate the influence of pH on the predictive ability, different QMs were derived at two pH values: 5.0 and 7.0. The QMs were validated using external test sets and compared to other servers for DP binding prediction. Additionally, in order to analyze the peptide-MHC protein interaction interface, a single docking of HLA-DP2 (DPA*0103, DPB1*0201) in complex with a self-peptide derived from the HLA-DR α-chain (pdb code: 3lqz) was analyzed using Rosetta Dock [19]. Our analysis affords a

Table 1 Alignment of HLA-DPA1 (α chain) and HLA-DPB1 (β chain) for the five most frequent DP proteins

| AA Pos. | 10 | 20 | 30 | 40 | 50 | 60 |
|---------|----|----|----|----|----|----|
| DPA1*01:03 | IKADHVSTYA | AFVQTHRPTG | EFMEFFDE | P6 | P6 | P6 |
| DPA1*02:01 | ANAILNNNL | LTLQRSNHT | |
| AA Pos. | 70 | 80 |
| DPA1*01:03 | ANAILLN | LTLQR | |
| DPA1*02:01 | ------- | ------- | |
| AA Pos. | 10 | 20 | 30 | 40 | 50 | 60 |
| DPB1*01:01 | RATPENFYVQ | GRQECYAFNG | TQRFLERY | NREEYARFDS | DVGEFRAVTE | LGRPAAEYWN |
| DPB1*02:01 | ------- | ------- | ------- | ------- | ------- | ------- |
| DPB1*04:01 | ------- | ------- | ------- | ------- | ------- | ------- |
| DPB1*04:02 | ------- | ------- | ------- | ------- | ------- | ------- |
| DPB1*05:01 | ------- | ------- | ------- | ------- | ------- | ------- |
| AA Pos. | 70 | 80 | 90 |
| DPB1*01:01 | SQKIDLEEKV | AVPDVCRH | YELDEAVTLQ |
| DPB1*02:01 | ------- | ------- | ------- | ------- | ------- | ------- |
| DPB1*04:01 | ------- | ------- | ------- | ------- | ------- | ------- |
| DPB1*04:02 | ------- | ------- | ------- | ------- | ------- | ------- |
| DPB1*05:01 | ------- | ------- | ------- | ------- | ------- | ------- |

Table 2 The most sensitive models for HLA-DP peptide binding prediction at threshold of top 5%

| DP protein | pH 7.0 | pH 5.0 |
|------------|--------|--------|
| model | sensitivity | AUC | model | sensitivity | AUC |
| DP1 | p2p7p8 | 0.426 | 0.865 | p3p7p8 | 0.455 | 0.860 |
| DP41 | p1p2 | 0.490 | 0.886 | p1 | 0.497 | 0.864 |
| DP42 | p1p3p5p5p7p7 | 0.471 | 0.883 | p1p2p3p8p9 | 0.504 | 0.900 |
| DP5 | p5p6 | 0.514 | 0.880 | p1p2p5p7p9 | 0.523 | 0.883 |
deep and detailed analysis of the different amino acid preferences at each position of peptides binding DP proteins.

**Methods**

**Input data**
The X-ray structure of the HLA-DP2 (DPA*0103, DPB1*0201) protein, in complex with a self-peptide derived from the HLA-DR α-chain, was used as the starting structure for homology modelling [10]. The covalently bound peptide was separated and defined as chain C. It consists of nine binding core positions (FHYLPLFLPS) and six flanking residues (RK at the N terminus and TGGS at the C terminus). The conformation of the protein was used to model by homology the four HLA-DP proteins. The conformation of the bound peptide was used as a template for the modelling of four virtual combinatorial peptide libraries.

**Homology modelling**
Models of four HLA-DP proteins were built using the X-ray structure of HLA-DP2 protein (pdb code: 3lqz) as the template for homology modelling. HLA-DP proteins used were: DP1 (DPA1*0201/DPB1*0101), DP41 (DPA1*0103/DPB1*0401), DP42 (DPA1*0103/DPB1*0402), and DP5 (DPA1*0201/DPB1*0501). The polymorphic amino acids among the first 80 amino acids from chain α (DPA1) and the first 90 amino acids from chain β (DPB1) were mutated accordingly. The resulting structure, in complex with the native peptide from the starting X-ray structure, was subjected to energy minimization by simulated annealing using the AMBER force field [20]. Each peptide-DP protein complex was used as a starting structure for generating the corresponding virtual peptide library.
Combinatorial peptide library

The nine positions forming the peptide binding core were examined. Four peptide libraries, each consisted of 172 peptides (19 amino acids × 9 positions + 1 original ligand), were built using PyMOL [21]. The SAAS (single amino acid substitution) approach was used to model the conformations of each altered side chains: after substitution, the peptide was minimized while keeping the MHC protein rigid. The protonation state of ionisable protein side chains was assigned to a standard ionisable state: neutral for His; positively charged for Arg and Lys; and negatively charged for Asp and Glu [22]. In the case of docking at pH 5, His was considered to be positively charged.

AutoDock protocol

A parallelized version of AutoDock 4.2 [23], employing an implementation of the Lamarckian genetic algorithm (GA), was used to model the peptide binding to HLA-DPs. All simulations were run on the IBM Blue Gene – P of the Bulgarian Supercomputing Centre. The input ligands for AutoDock 4.2 were prepared by using tools developed in-house using C# and .NET. The output data were mined by python scripts using the MGL Tools 1.5.4 package [24]. All retained poses considered in the study had an RMSD below 2.0 Å. To limit the computational burden of calculating peptide–MHC interactions at positions not involved in the static docking, all coordinates were kept fixed apart from the peptide residues of interest. These were left flexible. All GA settings were kept to their default values, apart from the number of energy evaluations and the number of generations which were set to 2 500 000 and 27 000, respectively. The docking grid was defined as a cuboid with respective dimensions of: 68 Å × 80 Å × 80 Å for DP1, 72 Å × 80 Å × 82 Å for DP41, 72 Å × 80 Å × 82 Å for DP42 and 72 Å × 80 Å × 82 Å for DP5 which encompassed the entire peptide binding site on DP. The output from ten independent GA runs for each ligand was processed and the pose (binding conformation) with the lowest Free Energy of Binding (FEB) was considered. FEB values represent the direct output from the AutoDock 4.2 scoring function which takes into consideration weighted terms for van der Waals dispersion/repulsion, hydrogen bonding, electrostatics, desolvation interactions, and the change in torsional free energy when the ligand goes from an unbound to a bound state.

Table 3 Pair energies in peptide binding pocket 1

| DP chain | position | aa | peptide position | aa | \(E_{\text{total}}\) | \(E_{\text{attractive}}\) | \(E_{\text{repulsive}}\) | \(E_{\text{sol}}\) | \(E_{\text{Hbnd}}\) | \(E_{\text{pair}}\) |
|----------|----------|----|-----------------|----|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| A        | 9        | Tyr | 1               | Phe | 0.02            | −0.4            | 0.41            | 0.41            | 0               | 0               |
| A        | 24       | Phe | 1               | Phe | −0.46           | −0.45           | 0.01            | 0.01            | 0               | 0               |
| A        | 32       | Phe | 1               | Phe | −1.69           | −1.64           | 0.05            | 0.05            | 0               | 0               |
| A        | 43       | Trp | 1               | Phe | −0.01           | −0.13           | 0.12            | 0.12            | 0               | 0               |
| A        | 52       | Phe | 1               | Phe | −0.39           | −0.42           | 0.04            | 0.04            | 0               | 0               |
| A        | 53       | Ser | 1               | Phe | −0.88           | −0.50           | 0.33            | 0.33            | −0.71           | 0               |
| A        | 54       | Phe | 1               | Phe | −0.75           | −1.50           | 0.41            | 0.34            | 0               | 0               |
| A        | 55       | Glu | 1               | Phe | 0.01            | −0.01           | 0.02            | 0.02            | 0               | 0               |
| B        | 80       | Asn | 1               | Phe | −0.12           | −1.52           | 0.03            | 1.38            | 0               | 0               |
| B        | 83       | Leu | 1               | Phe | −0.11           | −0.15           | 0.04            | 0.04            | 0               | 0               |
| B        | 84       | Gly | 1               | Phe | −0.17           | −0.21           | 0.04            | 0.04            | 0               | 0               |

Polymorphic residues are given in bold. \(E_{\text{sol}}\) corresponds to the sum of all energies between the pair residues; \(E_{\text{attractive}}\) and \(E_{\text{repulsive}}\) are the Lennard–Jones attractive and repulsive energies, respectively; \(E_{\text{Hbnd}}\) – the solvation energy; \(E_{\text{Hbnd}}\) – energy of hydrogen bonding per residue; \(E_{\text{pair}}\) – statistics-based pair term.

Figure 4 At pH 5.0 an additional hydrogen bond between the backbone carbonyl oxygen from the peptide position (−1) (here LYS-2) and the imidazole e-nitrogen of protein His79β is formed.
Docking score-based quantitative matrices (DS-QMs)
The FEBs derived from the docking experiments had negative and positive values. Negative FEBs correspond to binding peptides, while positive FEBs correspond to non-binding peptides. Only negative FEBs were considered; non-binding amino acids were assigned the penalty score of $-10$. The FEBs were normalized position per position using the following formula:

$$FEB_{i,\text{norm}} = \frac{FEB_i - FEB}{FEB_{\text{max}} - FEB_{\text{min}}},$$

Where $FEB_i$ is the binding energy of the $i$-th peptide, $FEB$ is the average for a given position, $FEB_{\text{max}}$ and $FEB_{\text{min}}$ are the maximum and minimum FEBs for a given position. Normalized FEBs were multiplied by $(−1)$ before being entered into the quantitative matrices (QMs) for ease of presentation. Thus, the positive FEBs correspond to preferred amino acids, and negative FEBs to non-preferred residues. Eight QMs were derived: two for each HLA-DP protein at pH 5.0 and pH 7.0, respectively.

Test set
Four test sets of peptides known to bind HLA-DP1, HLA-DP41, HLA-DP42, and HLA-DP5, respectively, were collected from the Immune Epitope Database [25] (June 2011 release). The test set of DP1 binders contained 102 peptides originating from 60 proteins. The DP41 test set contained 152 binding peptides from 71 proteins. The DP42 test set contained 122 binding peptides from 66 proteins. The DP5 test set contained 108 binding peptides from 66 proteins. The peptides had different lengths. No multiple binders were used. Each protein was represented as a set of overlapping nonamers. The nonameric subsequence of any known binder with the highest score was considered a binder; all other protein nonamers were considered as non-binders. The binding score of each nonamer was calculated as a sum of the weights of all nine positions or of different combinations thereof.

The tests were performed under conditions similar to those which an experimental immunologist might use: proteins were cleaved into overlapping nonamers, the binding score of each nonamer was predicted. Nonamers were then ranked according to their binding score and the top 5% of the predicted nonamers was selected. The selected peptides were then compared to the known binders. If the nonamer sequence was part of the known binder sequence, the predicted peptide was considered as a true predicted binder. The ratio of all true predicted binders to all binders in the corresponding test set defined the sensitivity of prediction at the top 5% cutoff. The test sets used in the present study are given as Additional file 1.

| Table 4 Pair energies in peptide position 2 |
|-------------------------------------------|
| DP chain | position | aa | peptide position | aa | $E_{\text{total}}$ | $E_{\text{attr}}$ | $E_{\text{rep}}$ | $E_{\text{sol}}$ | $E_{\text{bond}}$ | $E_{\text{pair}}$ |
| A | 9 | Tyr | 2 | His | $-0.49$ | $-0.69$ | 0 | 0.59 | $-0.39$ | 0 |
| A | 54 | Phe | 2 | His | $-0.16$ | $-0.25$ | 0 | 0.09 | 0 | 0 |
| B | 76 | Met | 2 | His | $-0.57$ | $-0.63$ | 0 | 0.06 | 0 | 0 |
| B | 79 | His | 2 | His | $-1.45$ | $-2.45$ | 0.03 | 1.13 | 0 | $-0.16$ |
| B | 80 | Asn | 2 | His | $-1.94$ | $-1.62$ | 0 | 1.73 | $-1.95$ | $-0.10$ |
| B | 83 | Leu | 2 | His | $-0.14$ | $-0.19$ | 0 | 0.05 | 0 | 0 |
| sum | | | | | $-4.75$ | $-5.83$ | 0.03 | 3.65 | $-2.34$ | $-0.26$ |

Polymorphic residues are given in bold.
Additionally, the models were compared in terms of the area under the receiver operating characteristics curve (AUC). Two variables - sensitivity and 1-specificity - were calculated at different thresholds. AUC is a quantitative measure of predictive ability and varies from 0.5 for random prediction to 1.0 for a perfect prediction.

Rosetta Dock protocol

The Rosetta Dock server (http://rosettadock.graylab.jhu.edu) was used to generate the pair interaction energies across the peptide-DP2 protein binding interface. The X-ray structure of the peptide-HLA-DP2 (DPA*0103, DPB1*0201) complex (pdb code: 3lqz) was used as input. The RosettaDock output file contains a table of pair energies. Several energy terms are generated: $E_{\text{tot}}$ is the sum of all energies between the pair residues; $E_{\text{atr}}$ and $E_{\text{rep}}$ are the Lennard–Jones attractive and repulsive energies, respectively; $E_{\text{solv}}$ is the solvation energy according to the Lazaridis–Karplus solvation model [26], which penalizes buried polar groups; $E_{\text{hbnd}}$ is the hydrogen bonding energy per residue; $E_{\text{pair}}$ is a statistically-based pair term derived from the PDB database, which favours salt bridges.

Results

Pair energies across peptide – HLA-DP2 protein binding interface

The peptide-DP protein binding interface was analysed using the RosettaDock server [19]. It consists of 39 residues: 21 residues belong to α-chain (9, 11, 22, 24, 32, 43, 52, 53, 54, 55, 57, 58, 62, 63, 65, 66, 68, 69, 70, 72 and 73) and 18 residues are from β-chain (9, 11, 12, 13, 24, 26, 28, 45, 55, 59, 65, 69, 72, 76, 79, 80, 83 and 84) [10] (Figure 1). Only five of the residues are polymorphic among the five most frequent DP proteins (Table 1). These are Tyr/Phe$^\beta_9$, Ala/Asp/Glu$^\beta_{55}$, Lys/Glu$^\beta_{69}$, Val/Met$^\beta_{76}$ and Asp/Gly$^\beta_{84}$. Asp$^\beta_{55}$ is involved in a salt-bridge with peptide Ser$^9$, while the other polymorphic residues do not form either an H-bond or a salt bridge with the bound peptide.

Docking score-based quantitative matrices (DS-QMs) for DP1, DP41, DP42 and DP5

Four libraries, each consisting of 172 peptides (19 amino acids × 9 positions + 1 original ligand), were built. Each peptide was docked separately into the corresponding DP rigid binding site. DS-QMs were derived based on normalized FEB scores, as described in Data and Methods. Dockings were performed at two pH values: 5.0 and 7.0. Over this pH range, only His undergoes protonation/deprotonation. At pH 5.0, His is protonated and very hydrophilic, yet at pH 7.0 His is neutral and less hydrophilic. The eight DS-QMs (four at pH 5.0 and four at pH 7.0) derived here are given in Additional file 2.

| Table 5 Pair energies in peptide position 3 |
|-------------------------------------------|
| **DP chain** | **position** | **aa** | **peptide position** | **aa** | $E_{\text{total}}$ | $E_{\text{atr}}$ | $E_{\text{rep}}$ | $E_{\text{solv}}$ | $E_{\text{hbnd}}$ | $E_{\text{pair}}$ |
| A | 9 | Tyr | 3 | Tyr | 0.53 | 0.67 | 0 | 0.14 | 0 | 0 |
| A | 22 | Phe | 3 | Tyr | 0.03 | 0.03 | 0 | 0 | 0 | 0 |
| A | 54 | Phe | 3 | Tyr | 0.75 | 0.77 | 0.03 | 0 | 0 | 0 |
| A | 55 | Glu | 3 | Tyr | 0.77 | 0.93 | 0 | 0.79 | 0.63 | 0 |
| A | 57 | Gln | 3 | Tyr | 0 | 0.01 | 0 | 0.01 | 0 | 0 |
| A | 58 | Gly | 3 | Tyr | 0.80 | 1.11 | 0 | 0.32 | 0 | 0 |
| A | 62 | Asn | 3 | Tyr | 0.01 | 0.18 | 0 | 0.18 | 0 | 0 |
| B | 76 | Met | 3 | Tyr | 0.28 | 0.40 | 0 | 0.11 | 0 | 0 |
| B | 80 | Asn | 3 | Tyr | 0.01 | 0.07 | 0 | 0.08 | 0 | 0 |

| sum | **−3.14** | **−4.17** | **0.03** | **1.63** | **−0.63** | **0** |

Polymorphic residues are given in bold.
External validation

A test set comprising 484 peptides known to bind HLA-DP1, HLA-DP41, HLA-DP42 or HLA-DP5, originating from 263 proteins, was used for external validation of the derived DS-QMs. Initially, the sensitivity of the top 5% of the best scored peptides for each position was assessed using DS-QMs calculated at pH 7.0 and pH 5.0. Next, all possible combinations of different positions were evaluated. The most predictive models among all possible combinations between the nine peptide positions are shown in Table 2. It is evident that almost all positions are involved in these highly predictive models, indicating that no peptide positions have a negligible effect on binding. The results also indicate that the models derived at pH 5.0 seem to predict better than those derived at pH 7.0. Moreover, different peptide positions are important for binding at different pH values.

Comparison to existing servers for HLA-DP binding prediction

The best performing models derived here were compared to two state-of-the-art servers for MHC class II binding prediction: NetMHCII [27] and IEDB [28]. Both use sequence-based models powered by artificial neural networks (ANN). NetMHCII identifies nonamers, while IEDB works only with 15mers. The tests were performed as follows: protein sequences were converted into sets of overlapping peptides (9mers for NetMHCII and 15mers for IEDB), and the binding score of each peptide was predicted; peptides were ranked according to their binding score, and the top 5% of the ranked peptides were selected and compared to known binders. If the predicted peptide was included in the known binder sequence, it was considered a true predicted binder. The ratio of all true predicted binders to all binders in the corresponding test set defined the sensitivity of prediction at the top 5% cut-off. The sensitivities were recorded and compared to our best predicted models at pH 5.0 (Figure 2a). Additionally, servers were compared in terms of AUC (Figure 2b). It is evident that our DS-QM models out-performed state-of-the-art servers for DP1, DP42 and DP5 proteins.

Effect of pH on peptide and protein His residues

As peptides typically bind to class II MHC proteins in an acidic environment, with a pH between 4.5 and 5.5, dockings were performed at pH 5.0 and pH 5.0, and compared in terms of their predictive ability. Better prediction was found for the DS-QMs derived at pH 5.0.

Table 6 Pair energies in peptide binding pocket 4

| DP chain position | aa | peptide position | aa | E_total | E_atr | E_rep | E_sol | E_bind | E_pair |
|-------------------|----|------------------|----|---------|-------|-------|-------|--------|--------|
| A 9               | Tyr | 4                | Leu| −0.02   | −0.04 | 0     | 0.03  | 0      | 0      |
| A 62              | Asn | 4                | Leu| −1.05   | −0.72 | 0     | 0.81  | −1.13  | 0      |
| B 13              | Gln | 4                | Leu| 0.18    | −1.78 | 0.71  | 1.90  | −0.66  | 0      |
| B 24              | Phe | 4                | Leu| −0.37   | −0.40 | 0.11  | −0.08 | 0      | 0      |
| B 26              | Glu | 4                | Leu| 0.08    | −0.10 | 0     | 0.19  | 0      | 0      |
| B 72              | Val | 4                | Leu| −0.76   | −0.62 | 0.11  | −0.25 | 0      | 0      |
| B 76              | Met | 4                | Leu| −0.86   | −0.88 | 0.16  | −0.14 | 0      | 0      |
| sum               |     |                  |    | −2.80   | −4.54 | 1.09  | 2.46  | −1.79  | 0      |

Polymorphic residues are given in bold.
The only amino acid sensitive to pH in the range 5.0 to 7.0 is Histidine. The pKₐ of the His imidazole is 6.0, thus making His protonated and very hydrophilic at pH 5.0 and unprotonated and less hydrophilic at pH 7.0. The influence of pH on the affinity of peptide binding to HLA-DP proteins has two potential aspects: influence on peptide protonation/deprotonation and influence on protein binding site protonation/deprotonation. Figure 3 summarizes the normalized FEB values for protonated and nonprotonated His residues at each of the nine peptide binding core positions. It is clear that protonated His residues are preferred in most peptide positions (p3 to p9). As the peptide binding site on DP proteins is predominantly negatively charged [29], preference for positively charged His were expected.

Five His residues are present in the HLA-DP binding site: four belong to the α-chain (positions 5, 16, 44 and 79) and one belongs to the β-chain (position 79). All are conserved among the studied DPs. Only His⁷⁹β contacts the binding peptide in the vicinity of peptide position 2; the other His residues are distant from the binding site. The protonation of His⁷⁹β allows an additional H-bond to be formed between the backbone carbonyl oxygen belonging to peptide position –1 (the position before p1) and the imidazole ε-nitrogen of His⁷⁹β (Figure 4). The estimated N-H…O=C bond energy for polypeptides in water environment lies within the range: 1.5 – 2 kcal/mol [30]. This means that the formation of this additional H-bond can increase the binding affinity constant of the peptide-protein complex by over an order of magnitude in the absence of other effects. This may explain the enhanced experimentally-observed equilibrium binding level seen at pH 5.0 [13].

Discussion

In the present study, molecular docking procedures developed recently for peptide binding prediction to HLA-DP2 protein [15] were significantly extended to include the four most frequent DP proteins [18]: DP1 (DPA1*0201/DPB1*0101), DP41 (DPA1*0103/ DPB1*0401), DP42 (DPA1*0103/DPB1*0402) and DP5 (DPA1*0201/ DPB1*0501). The X-ray structure of the peptide – HLA-DP2 protein complex was used as a starting template to model by homology the structure of the four DP proteins. In turn, these were used to generate combinatorial peptide libraries built using the SAAS principle. Peptides were docked into the DP binding site using AutoDock at pH 5.0 and pH 7.0. The resulting scores were recorded, normalized, and used to generate DS-QMs. The predictive ability of these QMs was tested using an external test set and compared to existing servers for DP binding prediction. The models derived at pH 5.0 predict better than those derived at pH 7.0, showing significantly improved predictions for three of the four DP proteins, when compared to current state-of-the-art servers. DS-QMs can recognize 50% of the known binders in the top 5% of predicted peptides. Moreover, a single docking

Table 7 Pair energies in peptide position 5

| DP chain | position | aa | peptide position | aa | E_total | E_str | E_rep | E_sol | E_hbond | E_pair |
|----------|----------|----|------------------|----|---------|-------|-------|-------|---------|--------|
| A        | 62       | Asn | 5                | Pro| 0.38    | −0.86 | 0.25  | 0.99  | 0       | 0      |
| A        | 65       | Ile | 5                | Pro| −0.29   | −0.21 | 0     | −0.08 | 0       | 0      |
| B        | 13       | Gln | 5                | Pro| −0.06   | −0.39 | 0     | 0.36  | −0.02   | 0      |
| B        | 26       | Glu | 5                | Pro| 0.04    | −0.04 | 0     | 0.08  | 0       | 0      |

No polymorphic residues exist here.
of HLA-DP2 (DPA*0103, DPB1*0201) in complex with a self-peptide derived from the HLA-DR α-chain (pdb code: 3lqz) was analysed using RosettaDock. This characterised more fully the interacting amino acids across the peptide – MHC binding interface, helping identify amino acid preferences at each position of the peptide binding core.

Peptide binding pocket 1 (p1) consists of 11 residues (Table 3). Ten of them are conserved and only Asp/Glyβ84 is dimorphic (Figure 5). DP1 and DP5 contain Aspβ84, while DP41 and DP42 contain Glyβ84 as does DP2. Aromatic amino acids such as Phe, Tyr, Trp and His, as well as aliphatic Ile and Leu are able to bind into this pocket. Additionally, the Aspβ84-containing proteins DP1 and DP5 accept positively charged Lys, Arg and His (when is charged at pH 5.0). A hydrogen bond is formed between Ser53α and NH of peptide position 1 (p1) (Table 1).

The peptide position 2 (p2) makes contacts with 6 residues of the binding site (Table 4), 5 of them are conserved, one (Met/Val76β) is dimorphic (Figure 6). Only DP1 contains Val76β, the remaining DPs have Met76β. The p2 side chain protrudes up the binding site close to the β chain and a variety of amino acids are well situated here. His at p2 makes H-bonds with Tyr9α and Asn80β, and salt bridges with His79β and Asn80β (Table 4). A π-π stacking of aromatic rings explains the preference of aromatic residues here [15]. Protonated His is not favored here.

The side chain of peptide position 3 (p3) protrudes up of the binding site close to the α chain. It contacts 7 α-chain residues and 2 β-chain residues, one of which is the dimorphic Met/Val76β (Table 5). Glu55α makes a hydrogen bond with Tyr OH-group (Figure 7). The amino acid preferences here are quite uniform for the four DPs: Tyr,

| DP chain | position | aa | peptide position | aa | E_total | E_str | E_rep | E_sol | E_bond | E_pair |
|----------|----------|----|-----------------|----|---------|-------|-------|-------|--------|--------|
| A        | 9        | Tyr | 6               | Phe | −0.01   | −0.04 | 0     | 0.03  | 0      | 0      |
| A        | 11       | Ala | 6               | Phe | −0.31   | −0.36 | 0.05  | 0     | 0      | 0      |
| A        | 22       | Phe | 6               | Phe | −0.17   | −0.16 | 0     | 0     | 0      | 0      |
| A        | 62       | Asn | 6               | Phe | −1.09   | −2.44 | 1.00  | 1.74  | −1.39  | 0      |
| A        | 63       | Ile | 6               | Phe | −0.01   | −0.01 | 0     | 0     | 0      | 0      |
| A        | 65       | Ile | 6               | Phe | −1.15   | −1.38 | 0.02  | 0.22  | 0      | 0      |
| A        | 66       | Leu | 6               | Phe | −0.76   | −1.33 | 0.77  | −0.21 | 0      | 0      |
| A        | 69       | Asn | 6               | Phe | 0.02    | −0.05 | 0     | 0.06  | 0      | 0      |
| B        | 11       | Gly | 6               | Phe | −0.66   | −0.82 | 0     | 0.17  | 0      | 0      |
| B        | 12       | Arg | 6               | Phe | −0.21   | −0.29 | 0     | 0.08  | 0      | 0      |
| B        | 13       | Gln | 6               | Phe | −0.31   | −1.56 | 0     | 1.25  | 0      | 0      |
| B        | 26       | Glu | 6               | Phe | −0.03   | −0.06 | 0     | 0.04  | 0      | 0      |
| B        | 28       | Tyr | 6               | Phe | −0.50   | −1.43 | 0.35  | 0.58  | 0      | 0      |

sum: −5.18 −9.93 2.14 4.01 −1.39 0

No polymorphic residues exist here.

Table 8 Pair energies in peptide binding pocket 6

Figure 9 Peptide position 5. The α chain residues are shown in light blue, the β chain residues – in dark salmon. No polymorphism exists here. Gln13β (given in magenta) make H-bonds with p5 residue Pro (given as PRO-7 in orange).
Trp, Phe, Pro and the positively charged Arg. Met<sup>76</sup>β-containing DPs (DP41, DP42 and DP5) accept a protonated His here.

Binding pocket 4 (p4) is large, shallow and negatively charged due to the presence there of Glu<sup>26</sup>β, Glu<sup>68</sup>β and Glu<sup>69</sup>β [10]. It strongly attracts positively charged amino acids such as Arg, Lys and protonated His. Leu, Tyr, Trp and Phe are also well accepted here. Asn<sup>62</sup>α and Gln<sup>13</sup>β make H-bonds with p4, as Leu does not fill the pocket [10]. Surprisingly, Glu, Gln and Asn also fit well into this pocket making H-bonds with Asn<sup>62</sup>α and Gln<sup>13</sup>β.

Position 5 (p5) protrudes from the binding cleft but it is still in close proximity to the negatively charged residues Glu<sup>26</sup>β, Glu<sup>68</sup>β and Glu<sup>69</sup>β. This explains the observed preferences for the positively charged Arg, Lys and protonated His and the disinclination for Asp and Glu. Pro at p5 hydrogen bonds to Gln<sup>13</sup>β and contacts Asn<sup>62</sup>α, Ile<sup>65</sup>α and Glu<sup>26</sup>β (Table 7 and Figure 9). Phe and Trp are also well accepted at p5. No polymorphism exists here (Table 2).

Binding pocket 6 (p6) is deep and formed by 8 residues from the α-chain and 5 residues from the β-chain (Table 8 and Figure 10). Asn<sup>62</sup>α makes an H-bond with the NH of Phe6. No polymorphism exists here (Table 2) and that makes the amino acid preferences at this pocket uniform for the five DPs. Phe, Tyr, Trp and His (protonated and nonprotonated) are well accepted here. Lys and Arg also fit well.

The side chain of position 7 (p7) lies tangentially to the binding site and is oriented towards the β-chain (Table 9). It is considered to be a secondary anchor position for some MHC class II proteins [31,32]. NH and CO of Leu7 make H-bonds with Tyr<sup>28</sup>β and Asn<sup>69</sup>α, respectively. The p7 side chain makes contacts with Ile<sup>65</sup>α, Glu<sup>26</sup>β, Phe<sup>45</sup>β, Trp<sup>59</sup>β, Ile<sup>65</sup>β and Glu/Lys<sup>69</sup>β.

Table 9 Pair energies in peptide position 7

| DP chain position | aa peptide position | aa | E<sub>total</sub> | E<sub>str</sub> | E<sub>rep</sub> | E<sub>sol</sub> | E<sub>hbd</sub> | E<sub>pair</sub> |
|------------------|---------------------|----|-----------------|--------------|-------------|-------------|-------------|--------------|
| A 65 Ile 7       | Leu                 | −0.24 | −0.34 | 0 | 0.10 | 0 | 0 | 0 |
| A 69 Asn 7       | Leu                 | −1.11 | −0.66 | 0 | 0.76 | −1.21 | 0 | 0 |
| B 26 Glu 7       | Leu                 | 0.15 | −0.47 | 0 | 0.62 | 0 | 0 | 0 |
| B 28 Tyr 7       | Leu                 | −1.73 | −1.36 | 0 | 0.87 | −1.24 | 0 | 0 |
| B 45 Phe 7       | Leu                 | −0.34 | −0.29 | 0 | −0.05 | 0 | 0 | 0 |
| B 59 Trp 7       | Leu                 | −1.48 | −1.87 | 0.13 | 0.26 | 0 | 0 | 0 |
| B 65 Ile 7       | Leu                 | −0.33 | −0.24 | 0 | −0.10 | 0 | 0 | 0 |
| B 69 Glu 7       | Leu                 | 0.08 | −0.25 | 0 | 0.33 | 0 | 0 | 0 |

**sum** | | −5.00 | −5.48 | 0.13 | 2.79 | −2.45 | 0 | 0 |

Polymorphic residues are given in bold.
Aliphatic residues are well accepted here. Additionally, Asp is preferred by Lys69β-containing DP proteins. Position 69β is dimorphic: DP1, DP41, DP42 and DP5 have Lys69β, while DP2 has Glu69β. Protonated His is accepted better here than the unprotonated form. Position 8 (p8) is solvent-exposed, yet shows preference for a variety of peptide residues: Trp, Tyr, Pro, Arg, Asn, Gly, Ala, His. Pro8 makes favourable contacts with Ile65α and Trp59β and disfavoured contacts with Asn69α and Asp55β (Table 10 and Figure 12). Position 55β is polymorphic: DP1 and DP41 contain Ala; DP2 and DP42 have Asp; and DP5 has Glu. However, this position is situated far from the side chain of p7 and does not influence the preferences there. Protonated His is preferred here.

Binding pocket 9 (p9) is formed from Asn68α, Asn69α, Leu70α, Thr72α, Leu73α, Phe/Tyr9β, Ala/Asp/Glu55β and Trp59β (Table 11 and Figure 13). It accepts large aromatic, polar, and even charged residues [10]. The side chain of p9 is oriented towards the α-chain. Ser9 is too short to fill the pocket. It makes H-bonds with Asn69α and Thr72α. Phe, Tyr, Trp, His fit well into this pocket. The Asp/Glu55β-containing DPs accept Arg and protonated His.

The influence of pH on the affinity of peptides binding to HLA-DP has two main aspects: influence on peptide protonation/deprotonation and influence on protein binding site protonation/deprotonation. At pH 5.0, His is positively charged and it is preferred at peptide positions 3 to 9. Among the five His residues in the HLA-DP binding site, only the protonation state of His79β affects peptide binding. At pH 5.0 an additional hydrogen bond is formed between the backbone carbonyl oxygen of the peptide position before p1 (p-1) and the imidazole E-nitrogen of His79β (Figure 4). This H-bond increases the peptide binding affinity by more than 3 orders of magnitude, perhaps explaining the higher experimentally-observed equilibrium binding level seen at pH 5.0 [30]. The peptide-protein association rate constants greatly increases at pH 5.0 (~ 40-fold), while the dissociation rates are almost unchanged in the pH range 5.0 – 7.0 [13]. Thus, one may speculate that the peptide-protein complex formed in the acidic environment of endosomes will also be stable in the neutral environment of the cell surface.

**Conclusion**

For peptide binding to the four most frequent HLA-DP proteins (DP1, DP41, DP42 and DP5), statistically the DS-QMs derived through molecular docking simulations...
at pH 5.0 gave better predictions than those derived at pH 7.0 and performed better than current state-of-the-art servers for MHC binding prediction. Clear differences are observed in our X-ray-based protein-peptide models: an additional hydrogen bond is formed between the backbone carbonyl oxygen belonging to the peptide position before p1 and the protonated ε-nitrogen of His\textsuperscript{β}\textsuperscript{β79}. This additional hydrogen bond may provide additional stabilization for all peptide regardless of their sequences, provided that they have a sufficiently long N-terminal extension. Protonated His residues make favourable interactions at most of the peptide binding core positions.

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**References**

1. Janeway CA, Travers P, Walport M, Capra JD: Immunobiology: The immune system in health and disease. New York: Elsevier Science Ltd/Garland Publishing; 1999.
2. Jardetzky TS, Brown JH, Gorga JC, Stern LJ, Urban RG, Strominger JL, Wiley DC: Crystallographic analysis of endogenous peptides associated with HLA-DR1 suggests a common, polyproline II-like conformation for bound peptides. Proc Natl Acad Sci USA 1996, 93(1):734–738.
3. Robinson J, Mistry K, McWilliam H, Lopez R, Parham P, Marsh SG: The IMGT/HLA Database. Nucleic Acids Res 2011, 39(Suppl 1):D1171–D1176.
4. Petersdorf EW, Smith AG, Mickelson EM, Longton GM, Anasetti C, Choo SY, Martin PJ, Hansen JA: The role of HLA-DPB1 disparity in the development of acute graft-versus-host disease following unrelated donor marrow transplantation. Blood 1993, 81(7):1923–1932.
5. Lympnay PA, Petrek M, Southcott AM, Newman Taylor AJ, Welsh KI, du Bois RM: HLA-DPB polymorphism: Glu 69 association with sarcoidosis. Eur J Immunogenet 1996, 23(5):353–359.
6. Begovich AB, Bugawan TL, Nepom BS, Kitzl W, Nepom GT, Erlich HA: A specific HLA-DPB\textsuperscript{Iβ} allele is associated with pauciarticular juvenile rheumatoid arthritis but not adult rheumatoid arthritis. Proc Natl Acad Sci USA 1989, 86(23):9489–9493.
7. Dong BP, Kimura A, Okubo R, Shingawara H, Tamai H, Nishimura Y, Sasazuki T: HLA-A and DPB1 loci confer susceptibility to Graves’ disease. Hum Immunol 1992, 35(3):165–172.
8. Potočnik I, Mosconi G, Forni A, Nemery B, Seghizzi P, Sorrentino R: Susceptibility to hard metal lung disease is strongly associated with the presence of glutamate 69 in HLA-DP\beta chain. Eur J Immunol 1997, 27(10):2741–2743.
9. Richeldi L, Sorrentino R, Saltini C: HLA-DPB1 glutamate 69: a genetic marker of beryllium disease. Science 1993, 262(5131):242–244.
10. Dai S, Murphy GA, Crawford F, Mack DG, Falta MT, Marrack P, Kappler JW, Fontenot JP: Crystal structure of HLA-DP2 and implications for
chronic beryllium disease. *Proc Natl Acad Sci USA* 2010, 107(16):7425–7430.

11. Aricozian M, Beretta F, Dweik R, Saltini C. Role of high-affinity HLA-DP specific CLIP-derived peptides in berylliosis binding to the HLA-DPGLu69 berylliosis-associated molecules and presentation to beryllium-sensitized T cells. *J Immunol* 2008, 128(1):P22-e662–e670.

12. Jensen PE. Regulation of antigen presentation by acidic pH. *J Exp Med* 1990, 171(3):1779–1784.

13. Reay PA, Wettstein DA, Davis MW. pH dependence and exchange of high and low responder peptides binding to a class II MHC molecule. *EMBO J* 1992, 11(8):2829–2839.

14. Rötzschke O, Lau JM, Hofstätter M, Falk K, Strominger JL. A pH-sensitive histidine residue as control element for ligand release from HLA-DP molecules. *Proc Natl Acad Sci USA* 2002, 99(36):16946–16950.

15. Patronov A, Dimitrov I, Flower DR, Doytchinova I. Peptide binding prediction for the human class II MHC allele HLA-DP2: a molecular docking approach. *BMC Struct Biol* 2011, 11:S2.

16. Khan JM, Ranganathan S. pDOCK: a new technique for rapid and accurate docking of peptide ligands to Major Histocompatibility Complexes. *Immunome Research* 2010, 6(Suppl 1):S2.

17. Atanasova M, Dimitrov I, Flower DR, Doytchinova I. MHC class II binding prediction by molecular docking. *Mol Inf* 2011, 30(4):368–375.

18. Sidney J, Steen A, Moore C, Ngo S, Chung J, Peters B, Sette A. Five HLA-DP molecules frequently expressed in the worldwide human population share a common HLA superpredis binding specificity. *J Immunol* 2010, 184(5):2492–2503.

19. Lyskov S, Gray JJ. The RosettaDock server for local protein-protein docking. *Nucleic Acids Res* 2008, 36(Web Server Issue):W233–W238.

20. Wang J, Ciaplik P, Kollman PA. How well does a restrained electrostatic potential (RESP) model perform in calculating conformational energies of organic and biological molecules? *J Comput Chem* 2008, 29(10):1049–1074.

21. PyMOL software. www.pymol.org.

22. Nelson DL, Cox MM: Lehninger principles of Biochemistry. In Forth. Forth edition. New York: Freeman WH; 2004:75–115.

23. Morris GM, Huey R, Lindstrom W, Sanner MF, Belew RK, Goodsell DS, Olson AJ: AutoDock4 and AutoDockTools4: automated docking with development. *J Mol Graphics Mod* 2009, 27(6):103–112.

24. Vita R, Zarebski L, Greenbaum JA, Emami H, Hoof I, Salimi N, Damle R, Lazaridis T, Karplus M. Effective energy function for proteins in solution. *Proc Natl Acad Sci USA* 2004, 101(22):7683–7687.

25. Sheu SY, Yang DY, Settle HL, Schlag EW. Energetics of hydrogen bonds in peptides. *Proc Natl Acad Sci USA* 2005, 102(22):7683–7687.

26. Stern LJ, Brown JH, Jargetzky TS, Gorga JC, Urban RG, Strominger JL, Wiley DC. Crystal structure of the human class II MHC protein HLA-DR1 complexed with an influenza virus peptide. *Nature* 1994, 368(6468):215–221.

27. Dessen A, Lawrence CM, Cupo S, Zaller DM, Wiley DC. X-ray crystal structure of HLA-DR4 (DRA*0101, DRB*0401) complexed with a peptide from human collagen II. *Immunity* 1997, 7(4):473–481.

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