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NetMHCIIpan-3.0, a common pan-specific MHC class II prediction method including all three human MHC class II isotypes, HLA-DR, HLA-DP and HLA-DQ

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Abstract

Major histocompatibility complex class II (MHCII) molecules play an important role in cellmediated immunity. They present specific peptides derived from endosomal proteins for recognition by T helper cells. The identification of peptides that bind to MHCII molecules is therefore of great importance for understanding the nature of immune responses and identifying T cell epitopes for the design of new vaccines and immunotherapies. Given the large number of MHC variants, and the costly experimental procedures needed to evaluate individual peptide-MHC interactions, computational predictions have become particularly attractive as first-line methods in epitope discovery. However, only a few so-called pan-specific prediction methods capable of predicting binding to any MHC molecule with known protein sequence are currently available, and all of them are limited to HLA-DR. Here, we present the first pan-specific method capable of predicting peptide binding to any HLA class II molecule with a defined protein sequence. The method employs a strategy common for HLA-DR, HLA-DP and HLA-DQ molecules to define the peptide-binding MHC environment in terms of a pseudo sequence. This strategy allows the inclusion of new molecules even from other species. The method was evaluated in several benchmarks and demonstrates a significant improvement over moleculespecific methods as well as the ability to predict peptide binding of previously uncharacterised

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MHCII molecules. To the best of our knowledge, the *NetMHCIIpan-3.0* method is the first panspecific predictor covering all HLA class II molecules with known sequences including HLA-DR, HLA-DP, and HLA-DQ. The *NetMHCpan-3.0* method is available at http://www.cbs.dtu.dk/ services/NetMHCIIpan-3.0.

Keywords

MHC class II; Tcell epitope; MHC binding specificity; Peptide–MHC binding; Human leukocyte antigens; Artificial neural networks

Introduction

Major histocompatibility complex (MHC) molecules play a key role in defining the specificity of the cellular immune system by presenting antigens to the immune system cells. In case of MHC class II molecules, these cells are T helper lymphocytes that recognize peptide–MHC complexes on the surface of antigen-presenting cells. Peptides presented by MHC class II molecules are derived from proteins taken up from the extracellular environment. Whereas a large number of peptides can be generated from pathogenic proteins, only a small part of these trigger an immune response. One of the most important events defining which peptides will trigger an immune response is binding to MHCII molecules expressed by the host (Castellino et al. 1997).

The human MHC locus (in humans called HLA for human leukocyte antigens) is extremely polymorphic and encodes thousands of different HLA class II molecules. Characterising the peptide-binding specificities of all the polymorphic MHC class II molecules is a serious experimental challenge. Therefore, during the last decades, large efforts have been put into the development of in silico methods for predicting peptide-binding affinities to MHC class II molecules. Using thousands of peptide-binding data points, several predictors have been developed and benchmarked (for review, see Nielsen et al. 2010b). One very important subset of these predictors consists of the so-called pan-specific methods that are capable of obtaining accurate predictions for molecules with limited or no binding data (Nielsen et al. 2008, 2010a; Zaitlen et al. 2008; Zhang et al. 2005). For MHC class I prediction, it has been demonstrated that a pan-specific approach can benefit from being trained on cross-loci, and even cross-species, data. That is, the predictive performance for HLA-B locus molecules is improved when including HLA-A locus data in the training of the pan-specific MHC class I binding prediction method (and vice versa), and the overall performance of predictions of HLA molecules is improved when including binding data representing non-human MHC molecules (Hoof et al. 2009). Extending this approach to MHC class II is not a trivial task. Differences in sequence polymorphism and corresponding details in the molecular structures across the different MHC class II loci complicate the development of cross-loci and crossspecies training strategies. This, combined with the very limited amount of data available for most MHC class II molecules, has limited the application of pan-specific methods to HLA-DR molecules. The understanding of HLA-DP and HLA-DQ binding specificities is limited to a handful of molecules which have been characterised experimentally, and beyond a few mouse H-2 molecules, to the best of our knowledge, no general MHC class II prediction method is available for non-human primates and other non-human species.

The number of state-of-the-art pan-specific methods for MHC class II molecules available up to date is very limited. The classical MHC class II predictor, *TEPITOPE* (Sturniolo et al. 1999), uses position-specific scoring matrices derived from experimental data. The method is, however, limited to 51 HLA-DR molecules only. In addition to this, a *TEPITOPEpan* predictor has been developed (Zhang et al. 2012) by extrapolating from the binding

specificities of the molecules characterised by TEPITOPE. The method is based on MHC pocket similarities and is capable of providing predictions for any HLA-DR molecule. The same is achieved by the *NetMHCIIpan-2.0* predictor (Nielsen et al. 2010a), which outperforms the *TEPITOPEpan* method in terms of prediction accuracy (Zhang et al. 2012). The method is based on artificial neural networks and uses an MHC binding pocket pseudo sequence combined with the peptide sequence as an input. Like the *TEPITOPEpan* method, *NetMHCIIpan-2.0* predicts binding for all HLA-DR molecules with a known primary sequence.

In this paper, we present a novel pan-specific predictor capable of predicting binding affinities to all HLA class II molecules. The method is based on artificial neural networks and has been trained on more than 50,000 quantitative peptide-binding measurements covering HLA-DR, HLA-DP, HLA-DQ as well as two murine molecules. Using a panel of benchmark setups, we seek to investigate to what extent the pan-specific method outperforms allele-specific approaches and whether it can obtain accurate predictions even for HLA molecules, which have not been experimentally characterised. Arriving at a "true" pan-specific method enabling prediction of the binding specificity for all HLA-II molecules, we end the analysis by conducting the first global analysis covering all prevalent HLA-II molecules, investigating and quantifying the functional diversity of the molecules encoded at the three HLA-II loci.

Materials and methods

Data sets

Training data used to develop the method consisted of quantitative MHC class II peptidebinding data retrieved from the IEDB database (Vita et al. 2010). In total, the training data set comprises 52,062 data points covering 24 HLA-DR, 5 HLA-DP, 6 HLA-DQ and 2 mouse (H-2) molecules. All molecules were covered by more than 50 peptide binding data points measured as IC_{50}/EC_{50} values which were log-transformed to fall in the range between 0 and 1 using the relation $1-log(IC_{50}nM)/log(50,000)$ (Nielsen et al. 2003). The evaluation set was restricted to HLA-DR molecules and contained 9,860 binding affinity measurements covering 13 molecules, four of which were not included in the training set. A summary of the data used to develop the method is presented in Table S1, and evaluation data set details are given in Table S2.

Mapping of MHC molecules

For constructing the *NetMHCIIpan* method, all MHC class II molecules need to be mapped to a common reference sequence. This is done by aligning alpha and beta chain sequences of all MHC molecules to the reference sequences, DRA101*01 and DRB101*01. For HLA-DR molecules, the mapping on a sequence level is in agreement with the mapping on the structural level. On the other hand, HLA-DP and HLA-DQ molecules demonstrate minor variations from HLA-DR in the peptide-binding domain in both the alpha and beta chains. To evaluate the structural impact of these variations, we employed the analysis described below. The analysis is based on the five available structures solved for HLA-DP and HLA-DQ molecules, which are compared to a representative high-resolution HLA-DR structure selected among the large number of structures available for HLA-DR molecules. The list of available HLA-DP and HLA-DQ structures from the Protein Data Bank (PDB) is given in Table 1.

An HLA-DR (PDB ID: 1A6A Ghosh et al. 1995) structure was chosen as a reference, and the HLA-DQ and HLA-DP structures were aligned to the binding domain of this reference molecule (Fig. 1). The superimpositions were performed in PyMOL (Schrodinger 2010) and

During the analysis, an important variation was observed for HLA-DQ molecules only and was investigated in more detail. We observed that sequences belonging to the HLA-DQA1*04, HLA-DQA1*05 and HLA-DQA1*06 serotype groups (e.g. sequences like HLA-DQA1*04:01) display a single amino acid deletion, which from a pure sequence point of view, corresponds to position 53 in HLA-DRA (Robinson et al. 2001). However, this leads to a shift of the preceding residues in the DQA sequences, which now realign with DRA positions 52 and 53. Although the deletion affects the orientation of the short -helical segment and the loop (residues 45–52) next to the P1 binding pocket (area marked in Fig. 1), these changes have negligible impact on peptide binding, as the reorientation appears to be a localised change, and very few contacts with the peptide are observed within the area. Due to the minor impact of the area discussed above to the binding of the peptide, an automated sequence alignment approach was chosen to identify the deletion in HLA-DQ sequences. Pair-wise sequence alignments were made and visualized using *ClustalW*(Larkin et al. 2007). Each HLA-DQ sequence was aligned one by one to the reference sequence of HLA-DR. The results are presented in Fig. 2. Figure 2a shows the alignments of HLA-DQ alpha chains with the amino acid deletion, while Fig. 2b demonstrates alignments of HLA-DQ alpha chains with no deletions. The alignments demonstrated that for all the HLA-DQ sequences that have a deletion, the deletion is consistently found in the same place (position 53 in the reference sequence).

MHC class II pseudo sequence

For constructing the *NetMHCIIpan* method, MHC class II molecules were represented by a pseudo sequence consisting of amino acid residues important for peptide binding. Amino acid residues comprising the pseudo sequence were defined as having their side chains pointing towards the peptide and being within 4.0 Å of the peptide-binding core in one or more of the MHC class II structures (including HLA-DR, HLA-DP and HLA-DQ molecules) available in the PDB (www.pdb.org; Berman et al. 2000). The MHC molecules were aligned using the PyMOL molecular viewer (Schrodinger 2010) and interacting residue positions extracted according to the distance criterion. Among the interacting residues, only those found to be polymorphic across the sequences of MHC molecules used for the training of the method were considered. The final pseudo sequence is composed of 15 residues from the alpha chain and 19 residues from the beta chain. The interaction map between the peptide and MHC pseudo sequence is given in Fig. 3.

Method

The *NetMHCIIpan-3.0* method was implemented as a conventional feed-forward artificial neural network method as described in detail by Nielsen et al. (2010a). The networks were trained using fivefold cross-validation. The data set was split into five groups of peptides based on a common motif clustering as described by Nielsen et al. (2007b). The difference in network architecture from the study presented by Nielsen et al. (2010a) was that network ensembles were trained with 10, 15, 40 and 60 hidden neurons. The BLOSUM50 matrix was used to encode peptide and MHC sequences for the network trainings. Each training was repeated 10 times with different initial configuration values as described in Nielsen et al. (2010a). In total, 40 (4 different numbers of hidden neurons times 10 different random seeds) networks were used for each training/test set combination leading to 200 (5 folds times 40 networks) networks for each molecule.

Leave-one-out setup

In order to assess the predictive performance of the method in the situation where a molecule is not part of the training data, a leave-one-out (LOO) approach was applied. Using LOO, the binding data for the molecule in question were excluded from the training data. Since our data set has a large number of peptides that have been measured for binding to multiple molecules, we also removed peptides common between the evaluation and training data sets to ensure unbiased LOO trainings. In order not to reduce the training set too much in this type of LOO trainings, the evaluation set was split into three subsets resulting into three different fivefold cross-validation trainings for each molecule. The details about such LOO setup are described in Karosiene et al. (2012).

Nearest neighbour approach (NN-finder)

In order to evaluate the performance of the pan-specific method on the molecules that are not found in the training set, we set up a nearest neighbour prediction approach which in this study we call NN-finder. This approach represents the simplest method where the predictions of a query molecule are obtained by first finding its nearest neighbour and using a subsequent allele-specific method to predict the query binding specificity. First of all, for each molecule in question, we found a corresponding nearest neighbour from the training set. The distance between two MHC molecules was calculated from the amino acid similarity between the two pseudo sequences as described by Nielsen et al. (2008), and the nearest neighbour to the molecule in question was defined as the molecule in the training set having the shortest distance. The binding data of each nearest neighbour were then used as training data for the corresponding query molecule. We retrained an allele-specific method from those training data using the NNAlign method (Andreatta et al. 2011) with settings identical to those used for NetMHCII (Nielsen et al. 2007b). The predictive performance for each query molecule was obtained by using its binding data as an evaluation set. In order for the performance to be directly comparable to the LOO results, the splitting of the evaluation set into three subsets was also used here.

Performance measures and statistical analysis

The predictive performance was measured in terms of Pearson's correlation coefficient (PCC) and area under the ROC curve (AUC). PCC values vary between 0 and 1, where 1 represents perfect predictions and 0 random predictions. For AUC measures, a performance value of 1 corresponds to a perfect prediction, and a value of 0.5 reflects random predictions. For more details concerning the performance measures, see Nielsen et al. (2010a). Throughout this study, PCC and AUC values were compared for different methods and evaluated using binomial tests with a significance level of 0.05.

Generation of HLA-II distance trees

For generation of the HLA-II distance tree, the most prevalent alpha and beta chains in the European population were selected as defined by the allele frequencies database (http:// www.allelefrequencies.net) (Gonzalez-Galarza et al. 2011). At a frequency threshold of 1 %, we found 21 HLA-DR1, 3 HLA-DPA1, 12 HLA-DPB1, 12 HAL-DQA1 and 13 HLA-DQB1 alleles. We constructed all HLA-DPA1–HLA-DPB1 and HLA-DQA1–HLA-DQB1 combinations arriving at a total of 21 HLA-DR, 36 HLA-DP and 156 HLA-DQ molecules. Sorting (on a per-loci level) the different molecules on descending population frequencies, we constructed a functional redundancy deduced set containing 72 molecules using the Hobohm1 algorithm (Hobohm et al. 1992) with redundancy defined as two molecules sharing a Pearson's correlation coefficient of 0. 99 or above when comparing the predicted binding affinities on a set of 200,000 random natural 15-mer peptides. The set of 72 nonredundant HLA-II molecules is comprised of 21 HLA-DR, 14 HLA-DP and 37 HLA-DQ

molecules. Next, we applied the *MHCcluster* method (Thomsen et al. 2013) to construct a tree describing the functional similarity between the different molecules. In short, the *MHCcluster* method functions as follows. Binding affinities of a set of 200,000 natural random 15-mer peptides are predicted for each of the HLA molecules using *NetMHCIIpan-3.0.* Next, the functional similarity between any two HLA molecules is defined by correlating the union of the predicted top 10 % strongest binding peptides for each molecule. The similarity is 1 if the two HLA molecules are predicted to have a perfectly overlapping peptide repertoire and negative if there is no or very limited overlap. The distance between two molecules is defined as 1–similarity. By using the unweighted pair group method with arithmetic mean clustering, the distance matrix is converted to a distance tree. The trees are next summarized, and a consensus tree is made with branch bootstrap values. Sequence logos were constructed from the predicted binding core of the top 1 % strongest predicted binders using *Seq2Logo* method with default settings (Thomsen and Nielsen 2012).

Results

In the following section, we give the results of applying the new pan-specific method: *NetMHCIIpan-3.0* to predict binding for a large set of MHC class II molecules from three human class II loci as well as a small set of mouse H-2 molecules.

NetMHCIIpan-3.0 method's new approach for getting pseudo sequence

In the most recent pan-specific MHC class II prediction method, NetMHCIIpan-2.0, the pseudo sequence is composed of 21 amino acids from positions within the HLA-DR beta chain that are in potential contact with a peptide using a 4.0 Å distance cut-off and polymorphic across the set of sequenced MHC class II molecules available at the time of the study (Nielsen et al. 2010a). For the NetMHCIIpan-3.0 method described here, the pseudo sequence contains 19 residues from the beta chain of MHC molecules. The main difference between two pseudo sequence obtaining approaches resulting into different number of pseudo sequence positions is that the NetMHCIIpan-3.0 considers polymorphism across the sequences of MHC molecules from the training set only. In order to evaluate this new approach for obtaining the pseudo sequence, we performed a fivefold cross-validation training and compared the results of those reported for NetMHCIIpan-2.0 (Nielsen et al. 2010a). In this comparison, only HLA-DR molecules were considered due to availability of results from both methods. Moreover, for the new method, only the part of the pseudo sequence corresponding to beta chain positions was included as the NetMHCIIpan-2.0 method only includes beta chain residues in the pseudo sequence. The results are shown in Table 2.

The results in Table 2 demonstrate that the new approach for obtaining the pseudo sequence leads to a significantly (*p* values<0.05) improved predictive performance compared to the original approach when the pan-specific training approach is applied to the HLA-DR data set. The average increased from 0.688 to 0.695 and from 0.846 to 0.847 for PCC and AUC values, respectively. *NetMHCIIpan-3.0* achieves the highest performance for most of the molecules (PCC values are higher for 20 out of 24 molecules, and AUC values are higher for 17 out of 23 molecules, excluding ties). The results demonstrate that the new approach of obtaining pseudo sequences for the neural network trainings improves the predictive performance of the method.

Per-locus training versus cross-loci training

To the best of our knowledge, all pan-specific prediction methods for HLA class II molecules available up to date are limited to HLA-DR. In this study, we introduce an approach for combining residues from the alpha and beta chains into one pseudo sequence. The procedure for the pseudo sequence construction is universal to all MHC class II complexes allowing the pan-specific method to be trained in a cross-loci/cross-species manner (see "Materials and methods") arriving at one common method suitable for all MHC class II molecules.

To evaluate how such a cross-loci/cross-species impacts the predictive performance of the method, we compared per-locus (and per molecule, see below) training with the pan-specific training including cross-loci data. The results are shown in Fig. 4. Detailed results are given in Table S3. The figure gives average PCC and AUC values for each locus when the method was trained in a cross-loci manner including all HLA molecules and when trained using binding data restricted to each locus, respectively. As can be seen from the figure, the overall performance of the two training approaches is similar. For HLA-DR, the predictive performance improved when training in a cross-loci manner compared to per-locus training. For HLA-DQ and HLA-DP, the performance on the other hand is slightly reduced. This reduction is, however, only significant for HLA-DQ and only when measuring AUC performance values.

Pan-specific versus allele-specific method

As a pan-specific approach, the method presented in this study benefits from the information even from molecules covered by limited binding data or molecules from different loci/ species. To demonstrate this, we present a comparison of the performance values obtained for the *NetMHCIIpan-3.0* method and allele-specific *NN-align* method using fivefold cross-validation (see Table 3). The *NN-align* prediction method was trained as described by Nielsen and Lund (2009), using the same data partitioning based on the common motif clustering approach as used for *NetMHCIIpan-3.0*.

The results presented in Table 3 demonstrate that the pan-specific *NetMHCIIpan-3.0* predictor significantly outperforms the allele-specific *NN-align* method (*p* value<0.0001 for both PCC and AUC values). These results show that the pan-specific method benefits from the binding data measured to different molecules. It also demonstrates that adding data from other molecules significantly boosts the performance for molecules represented by limited peptide-binding measurements. Out of 10 molecules described by less than 400 data points and less than 100 binders, 10 and 9 are shown to obtain higher performance using panspecific predictor in terms of PCC and AUC values, respectively. The allele-specific *NN-align* method gives higher PCC and AUC values for three molecules all defined by more than 1,700 peptide-binding data.

Leave-one-out performance

In order to demonstrate how the method performs when predicting binding to novel and uncharacterised molecules, we performed a LOO experiment. In the LOO experiment, a molecule in question was excluded from the training data set, and its binding data acted as an evaluation set. Likewise, all peptides, included in the binding data set of the given molecules, were excluded from the training data in order to avoid biassed overlapping between the evaluation and the training sets. This was done in three rounds by removing one third of the peptides from the evaluation set at a time and performing fivefold crossvalidation training in each round. The performance for the query molecule was obtained by combining the predictions of all three evaluation subsets. For the benchmark, we compared the results with the predictive performance of the simple allele-specific approach based on

finding the nearest neighbour, *NN-finder*. For this method, the molecule in question and its binding data were also acting as an evaluation set, while the training data were composed of the peptide binding data of the molecule from the training set having the shortest distance to the molecule in question. The results are depicted in Fig. 5 and presented in detail in Table S4. It is apparent from the results that *NetMHCIIpan-3.0* outperforms the *NN-finder* approach for all loci in terms of average PCC and AUC. Even though we find general improvement when comparing the pan-specific method to the nearest neighbour approach, a significant difference (due to the small number of molecules for each subset) is observed only for HLA-DR molecules (*p* value<0.0001). The significance for the mouse allelic locus (H-2) was not assessed due to only two molecules being available.

The method shows decreased predictive performance with the distance to the nearest neighbour from the training set (Fig. 6). The figure illustrates how the predictive performance of the pan-specific method depends on the distance to the nearest neighbour calculated in terms of pseudo sequence similarities as explained in "Materials and methods". Regression analysis showed that the performance is decreased significantly with the increasing distance (*p* value=0.031, exact permutation test).

Independent evaluation of the final NetMHCIIpan-3.0 predictor

For the final evaluation of the pan-specific method common for HLA-DR, HLA-DP, HLA-DQ and mouse molecules, the method was trained using all the available data (52,062 data points) and evaluated on an independent HLA-DR evaluation set containing 9,860 data points. The method was compared with the most recent version of the class II pan-specific predictor *NetMHCIIpan-2.0* (Nielsen et al. 2010a). From the results given in Table 4, it is apparent that *NetMHCIIpan-3.0* outperforms *NetMHCIIpan-2.0* (average PCC is 0.603 compared with 0.586 and average AUC 0.807 compared with 0.802). Although the difference in performances was observed not to be significant, the new pan-specific method showshigher performance for most of the molecules from the evaluation set (*NetMHCIIpan-3.0* wins 9 out of 13 and 6 out of 12 times in terms of PCC and AUC measures, respectively).

The *NetMHCIIpan-3.0* method presented and benchmarked in this paper was implemented as a web server and is available online at http://www.cbs.dtu.dk/services/NetMHCIIpan-3.0.

Functional clustering of HLA class II molecules

Given the potential of the NetMHCpan-3.0 method to predict binding for any MHC class II molecules with known alpha and beta chain protein sequences, we next applied the method to give an overall estimate of the functional diversity of molecules from the HLA-DR, HLA-DP and HLA-DQ loci molecules. The analysis of the most prevalent alpha and beta chains in the European population was done as described in the "Materials and methods", and the result is shown in Fig. 7. From the figure, it is apparent that the molecules encoded at the three loci display very limited functional overlap. Also, one can notice that the HLA-DP locus molecules display a very limited functional diversity compared to the HLA-DR and HLA-DQ loci molecules. This is also reflected when measuring the functional diversity of an HLA locus in terms of the mean and standard deviation of the intra-locus distances. Here, we find that the mean intra-distance is significantly shorter (p < 0.001, Student's t test) for HLA-DP compared to HLA-DQ and HLA-DR. We can further relate these differences in functional diversity to the degree of polymorphism at a population level of the HLA pseudo sequences of each locus. Estimating polymorphism in terms of the Kullback-Leibler information content (or divergence sum) (Kullback and Leibler 1951) for the 34 positions in the pseudo sequence for the three loci, we find that this value is significantly higher (p<0.001, t test) for DP compared to DQ and DR, hence demonstrating that DP molecules

share a significantly lower degree of polymorphism compared to the molecules at the two other loci.

In terms of the predicted functionality, we recover for HLA-DRB1 the overall clustering proposed earlier (Nielsen et al. 2008) with 9 well-defined subgroups (supertypes). For HLA-DQ, the overall functionality seems reduced compared to HLA-DR, with only 5/6 well-defined subgroups, and as stated above, HLA-DP seems to encode for the least functionally diverse set of molecules with only one specificity group being present. Sequence logos for selected subgroups and subgroup representatives are included in the figure to illustrate the functional difference between the different molecules. In general, the predicted binding motifs are in agreement with the motifs proposed earlier for the limited set of HLA class II molecules experimentally characterised by peptide binding data (Andreatta and Nielsen 2012; Andreatta et al. 2011).

Discussion and conclusion

Identification of peptides binding to MHC is a critical step in understanding T cell immune responses. The human MHC genomic region (HLA) is extremely polymorphic comprising several thousands alleles, many encoding a distinct molecule. The potentially unique specificities remain experimentally uncharacterised for the vast majority of HLA molecules.

The sequences of human MHC class II molecules stored in the IMGT database (Robinson et al. 2001) cover over 600 different HLA-DR variants and more than 6,000 different combinations of HLA-DP and HLA-DQ alpha and beta chains. Of these many molecules, less than 30 HLA-DR and only 5 HLA-DP and 6 HLA-DQ molecules have been experimentally characterised with binding data allowing for an accurate estimate of their binding specificity. In order to span this gap, several methods have been developed and benchmarked during the last decade for the prediction of peptide binding to MHC class II molecules (for review, see Nielsen et al. 2010b). Here, pan-specific methods play an important role, as they are capable of giving predictions to those molecules, which have not yet been characterised experimentally. However, until now, MHC class II pan-specific binding prediction approaches have been limited to HLA-DR molecules, leaving a gap in the general understanding of binding specificities for HLA-DP and HLA-DQ molecules (Nielsen et al. 2010b).

In this paper, we present a pan-specific method, *NetMHCIIpan-3.0*, capable of predicting peptide binding to all HLA molecules. To the best of our knowledge, this is the first predictor common for HLA-DR, HLA-DP and HLA-DQ molecules. The method is based on artificial neural networks and is trained on 52,062 quantitative peptide binding data covering all HLA as well as two mouse molecules.

NetMHCIIpan-3.0 uses a new approach for defining the peptide-binding environment of MHC in terms of pseudo sequence as compared with the most recent *NetMHCIIpan-2.0* method (Nielsen et al. 2010a). The main difference between the two approaches for obtaining pseudo sequence is that for *NetMHCIIpan-3.0*, only polymorphism within the training set is considered whereas the *NetMHCIIpan-2.0* method includes polymorphism across all known MHC class II sequences. Our results demonstrated that the new approach for defining the pseudo sequence leads to a significantly improved predictive performance.

Several large-scale benchmarks were carried out that demonstrated that the *NetMHCIIpan-3.0* method fulfils the requirements for the pan-specific methods. Its performance was found to be significantly better than that of the allele-specific *NN-align* predictor (Nielsen and Lund 2009). In particular, the method outperformed *NN-align* for molecules characterised with only a limited number of binding data. These results hence

agree with the results obtained when benchmarking the original *NetMHCIIpan* method (Nielsen et al. 2010a) and underline the unique power of the pan-specific approach in providing accurate predictions also for molecules characterised with limited peptide-binding data as it has also been demonstrated previously for MHC class I predictions (Karosiene et al. 2012; Nielsen et al. 2007a; Zhang et al. 2009b).

To mimic the situation where the *NetMHCIIpan-3.0* method is applied to predict binding for uncharacterised MHC molecules, we conducted a panel of LOO experiments. In these experiments, binding data for one MHC molecule at a time were removed from the training, and the predictive performance next evaluated on the left-out data. The LOO results demonstrated that the proposed method is capable of predicting binding affinity for the molecules for which no binding data are available in the training process. In addition to this, the method showed decreased predictive performance with the distance to the nearest neighbour from the training set, which is in agreement with previous studies on MHC class I (Hoof et al. 2009; Karosiene et al. 2012; Zhang et al. 2009a).

From the results included here, one can notice that HLA-DP molecules demonstrate higher performance values compared with DR and DQ performances. As this high performance is maintained also for the allele-specific (and pseudo sequence independent) NN-align approach, the high performance is not due to the fact that DP molecules are found to be very close to each other in terms of pseudo sequence similarity and function. The reason for this different performance is rather related to the differences within distributions of the binding data available for each locus. The HLA-DP data are very well separated with the majority of the data being either strong or very weak binders. This is in strong contrast to the data for DQ and DR molecules where the majority of the data have intermediate binding affinity (data not shown). This difference in binding affinity distribution strongly influences the predictive performance, as well-separated data sets (as is the case for DP) in general achieve a higher predictive performance. To prove this further, we have performed the analysis where we, in the evaluation of the prediction methods, mimicked the distribution of the HLA-DP data for the DR loci. The analysis demonstrated that the performance for DR molecules is significantly increased when the data match the distribution observed for the DP molecules compared to the original DR data distributions (data not shown).

The *NetMHCIIpan-3.0* predictor showed higher performance when compared with the *NetMHCIIpan-2.0* method on the external evaluation set. The increased performance of the *NetMHCIIpan-3.0* method demonstrates its promising ability to improve when more data become available for molecules from other loci/species.

We further presented a powerful application of the developed pan-specific predictor. We applied the *NetMHCIIpan-3.0* method to functionally cluster the most prevalent HLA alleles of the European population. For HLA class I, clustering of molecules into supertypes was proposed by the analysis carried out using experimental data (Lund et al. 2004; Sette and Sidney 1999) and extended by applying pan-specific class I predictor (Nielsen et al. 2007a). However, for MHC class II, the amount of experimental data remains too limited to perform such a cluster analysis, which therefore so far has been limited to HLA-DR molecules (Nielsen et al. 2008). The analysis performed here hence is the first study suggesting reduction of polymorphism of HLA class II molecules by definition of clusters based on similarities in predicted functional binding specificities. Such clustering builds a base for facilitating identification of T helper cell epitopes within different ethnic groups having a high value in the design of epitope-based vaccines.

As we have discussed earlier, developing a cross-loci method for MHC class II is complicated due to the differences of sequences and structures of different loci (Nielsen et

al. 2010a). However, as also suggested in this earlier publication, with increasing amounts of binding data covering HLA-DP and HLA-DQ molecules, pan-specific methods may benefit from cross-loci training. In this study, we demonstrated that this is indeed the case. The performance of the proposed *NetMHCIIpan-3.0* method when trained on cross-loci was shown to be comparable with that of a method trained on per-loci data. So far, no significant improvement was found between the per-loci and cross-loci trained method. However, this is most likely due to the very low number of HLA-DQ and HLA-DP molecules included in the study and is expected to change with the inclusion of more data covering HLA-DP and HLA-DQ molecules. The situation is hence parallel to that observed for MHC class I. Here, at first, only limited data characterising HLA-A and HLA-B molecules were available for development of the original version of *NetMHCpan*, and an optimal performance was obtained when the method was trained in a loci-specific manner (Nielsen et al. 2007a). Only when binding data became available covering more HLA molecules as well as MHC molecules from non-human species (including non-human primates) was a cross-loci/cross-species training strategy found to be optimal (Hoof et al. 2009).

The training strategy outlined here for the MHC class II pan-specific prediction method is highly flexible and readily allows inclusion of novel data both in terms of peptides and MHC molecules. This flexibility makes the method a powerful and unique platform for the development of a pan-specific MHC class II predictor covering not only the human class II molecules but also MHC molecules from other species of interest. Lessons learned from MHC class I suggest that such a "true" pan-specific approach is feasible and that prediction accuracies for both human and non-human MHC molecules can be greatly boosted given the ability of the pan-specific method to leverage information across species and loci (Nene et al. 2012).

In conclusion, we believe the proposed *NetMHCIIpan-3.0* method is an important step forward in boosting MHC class II binding predictions covering a large number of molecules from different species and therefore reduces experimental costs for the immunologists working within the field of epitope-based vaccine design.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Fig. 1.

Superimposition of HLA-DR, HLA-DP and HLA-DQ alpha chains. HLA-DR alpha chain (PDB ID: 1A6A Ghosh et al. 1995) is shown in *yellow* and was used as a reference chain. HLA-DP chain (PDB ID: 3LQZ Dai et al. 2010) is shown in *green*, HLA-DQ chain without a gap (PDB ID: 1JK8 Lee et al. 2001) is shown in *orange* and HLA-DQ chain with a gap (PDB ID: 1S9V Kim et al. 2004) is shown in *blue*. The area affected by the deletion in DQA sequence is circled

а	35 40 45 50 55 60 65 70 75 80 85 90
DRA1_01_01	DMAKKETVWRLEEFGRFASFEAQGALANIAVDKANLEIMTKRSNYTPITNVPPEVTVLTN
DQA1_05_04	DLGRKETVWCLPVLRQFR-FDPQFALTNIAVLKHNLNSLIKRSNSTAATN
DQA1_05_07	DLGRKETVWCLPVLRQFR-FDPQFALTNIAVLKHNLNSLIKRSNSTAATNEVPEVTVFSK
DQA1_05_01	DLGRKETVWCLPVLRQFR-FDPQFALTNIAVLKHNLNSLIKRSNSTAATN
DQA1 05 06	DLGRKETVWCLPVLRQFR-FDPQFALTNIAVLKHNLNSLIKRSNSTAATNEVPEVTVFSK
DQA1_05_03	DLGRKETVWCLPVLRQFR-FDPQFALTNIAVLKHNLNSLIKRSNSTAATNEVPEVTVFSK
DQA1 05 02	DLGRKETVWCLPVLRQFR-FDRQFALTNIAVLKHNLNSLIKRSNSTAAT
DQA1 05 08	DLGRKETVWCLPVLRQFR-FDPQFALTNIAVLKHNLNSLIKRSNSTAATNEVPEVTVFSK
DQA1 05 05	DLGRKETVWCLPVLRQFR-FDPQFALTNIAVLKHNLNSLIKRSNSTAATNEVPEVTVFSK
DQA1 05 11	DLGRKETVWCLPVLRQFR-FDPQFALTNIAVLKHNLNSLIKRSNSTAATNEVPEVTVFSK
DQA1 05 10	DLGRKETVWCLPVLRQFR-FDPQFALTNIAVLKHNLNSLVKRSNSTAATNEVPEVTVFSK
DQA1 05 09	DLGRKETVWCLPVLRQFR-FDPQFALTNIAVLKHNLNSLIKRSNSTAATNEVPEVTVFSK
DQA1_04_02	DLGRKETVWCLPVLRQFR-FDPQFALTNIAVTKHNLNILIKRSNSTAATNEVPEVTVFSK
DQA1 04 04	DLGRKETVWCLPVLRQFR-FDPQFALTNIAVTKHNLNILIKRSNSTAATNEVPEVTVFSK
DQA1 06 01	DLGRKETVWCLPVLRQFR-FDPQFALTNIAVTKHNLNILIKRSNSTAATN
DQA1 06 02	DLGRKETVWCLPVLRQFR-FDPQFALTNIAVTKHNLNILIKRSNSTAATNEVPEVTVFSK
DQA1 02 01	DLERKETVWKLPLFHRLR-FDPQFALTNIAVLKHNLNILIKRSNSTAATNEVPEVTVFSK
h	*: :**** * : :: *: * **:**** * **: : **** *. *
ט 1 01 01 גפת	
DOA1 01 04	
$DQA1_01_04$	
$DQA1_01_01$	DIERKEIAWRWFEFSKIGGIDFQGALKNMAVAKHNLNIMIKKINSIAAINEVFEVIVISK
DQA1_01_01	DIERKETAWRWPEFSKFGGFDPQGALKNMAVAKINLNIMIKKINSTAAINEVPEVIVFSK
DQA1_01_02	DLERKETAWRWPEFSKFGGFDPQGALRNMAVARHNLNIMIKRINSTAATNEVPEVIVFSK
DQAI_UI_U3	DLEKKETAWRWPEFSKFGGFDPQGALKNMAVAKHNLNIMIKRYNSTAATNEVPEVTVFSK
DQA1_03_02	DLERKETVWQLPLFRRFR R FDPQFALTNIAVLKHNLNIVIKRSNSTAATNEVPEVTVFSK
DQA1_03_03	DLERKETVWQLPLFRRFR K FDPQFALTNIAVLKHNLNIVIKRSNSTAATNEVPEVTVFSK
DQA1_03_01	DLERKETVWQLPLFRRFRRFDPQFALTNIAVLKHNLNIVIKRSNSTAATNEVPEVTVFSK
	** ****** * ***************************

Fig. 2.

Part of sequence alignments of HLA-DQ alpha chains to HLA-DR reference sequence of HLA-DRA1*0101 molecule. **a** Sequence alignments of HLA-DQ sequences with gaps, **b** demonstrates the alignment of other HLA-DQ molecules to the same reference sequence. Reference sequence and the position corresponding to the insertion are marked in bold. The alignments were visualized using *ClustalW*(Larkin et al. 2007)

Karosiene et al.



Fig. 3.

Interaction map between the peptide and MHC class II pseudo sequence. The *columns* give the MHC position numbering separately for alpha and beta chains and refer to HLA-DR. The *rows* show peptide binding core positions. *Red squares* marking interaction between a particular position of the peptide and MHC define contacts between corresponding two residues



Fig. 4.

Comparison of the method performance when trained on perlocus data and cross-loci data. Average PCC and average AUC values for each locus are demonstrated on the *left* and *right* panel, respectively. Significant *p* values are given above the *bars* for corresponding loci. The difference in predictive performance between the per-locus and cross-loci training is significant only for HLA-DQ when measuring AUC performance values



Fig. 5.

Leave-one-out results for the *NetMHCIIpan-3.0* method in comparison with the *NN-finder* approach. Average performance measures in terms of PCC and AUC are given in the *left* and *right* panel, respectively. Significant *p* values are given above the *bars* for corresponding loci (not available for H-2 locus)

Karosiene et al.



Fig. 6.

Predictive performance of the *NetMHCIIpan-3.0* method for the molecules from our data set as a function of distance to the nearest neighbour. The performance was obtained using LOO setup as explained in the "Materials and methods" section. The distance to the nearest neighbour was calculated as described by Nielsen at al. (2008). The *solid line* represents the least square fit for the data

Karosiene et al.



Fig. 7.

Functional clustering of the 72 HLA molecules from the European population. HLA-DR molecules are displayed in *red*, HLA-DP molecules are displayed in *green* and HLA-DQ molecules are shown in *blue*. Sequence logos showing the binding motif are presented for selected molecules representing the different specificity groups

Structures of all HLA-DP and HLA-DQ molecules available in the PDB database

PDB ID	Alpha chain	Beta chain
3LQZ	HLA-DPA1*0103	HLA-DPB1*0201
1UVQ	HLA-DQA1*0102	HLA-DQB1*0602
1JK8	HLA-DQA1*0302	HLA-DQB1*0302
1S9V	HLA-DQA1*0501	HLA-DQB1*0201
2NNA	HLA-DQA1*0301	HLA-DQB1*0302

Fivefold cross-validation performance for HLA-DR molecules of a pan-specific *NetMHCIIpan-2.0* compared with *NetMHCIIpan-3.0*

Molecule name	#pep	#bind	NetMHCIIpan-2.0		NetMHCIIpan-3.0	
			PCC	AUC	PCC	AUC
DRB1*0101	7,685	4,382	0.711	0.846	0.716	0.848
DRB1*0301	2,505	649	0.709	0.864	0.723	0.868
DRB1*0302	148	44	0.525	0.757	0.569	0.786
DRB1*0401	3,116	1,039	0.670	0.848	0.671	0.846
DRB1*0404	577	336	0.630	0.818	0.656	0.829
DRB1*0405	1,582	627	0.698	0.858	0.712	0.862
DRB1*0701	1,745	849	0.740	0.864	0.732	0.862
DRB1*0802	1,520	431	0.526	0.780	0.542	0.784
DRB1*0806	118	91	0.796	0.924	0.792	0.933
DRB1*0813	1,370	455	0.746	0.885	0.751	0.888
DRB1*0819	116	54	0.608	0.808	0.610	0.803
DRB1*0901	1,520	622	0.634	0.818	0.647	0.828
DRB1*1101	1,794	778	0.777	0.883	0.780	0.883
DRB1*1201	117	81	0.764	0.892	0.768	0.896
DRB1*1202	117	79	0.769	0.900	0.778	0.910
DRB1*1302	1,580	493	0.634	0.825	0.636	0.822
DRB1*1402	118	78	0.694	0.860	0.724	0.879
DRB1*1404	30	16	0.613	0.737	0.511	0.629
DRB1*1412	116	63	0.757	0.894	0.754	0.890
DRB1*1501	1,769	709	0.653	0.819	0.682	0.830
DRB3*0101	1,501	281	0.690	0.850	0.700	0.858
DRB3*0301	160	70	0.736	0.853	0.752	0.869
DRB4*0101	1,521	485	0.675	0.837	0.699	0.847
DRB5*0101	3,106	1,280	0.765	0.882	0.769	0.885
Total	33,931	13,992				
Average			0.688	0.846	0.695	0.847
<i>p</i> value			0.002	0.035		

NetMHCIIpan-2.0 is the method described by Nielsen et al. (2010a), which uses pseudo sequences composed of polymorphic amino acids that have one or more potential contacts with a peptide (length=21). The performance values for this method are taken from the publication. *NetMHCIIpan-3.0* employs pseudo sequences obtained by finding contacts that side chains of MHC molecules have with the peptide and taking polymorphic positions within the training set (length= 19 for beta chain only). The values in bold show the higher score for each molecule for corresponding performance measures (PCC or AUC). *p* values were obtained using a binomial test excluding ties

#pep-number of peptide binding data available for each molecule, #bind-number of peptides that have a binding affinity stronger than 500 nM

Fivefold cross-validation performance for the pan-specific *NetMHCIIpan-3.0* method compared with the allele-specific *NN-align* method using our benchmark data set

Molecule name	#pep	#bind	NetMHCIIpan-3.0		NN-align	
			РСС	AUC	PCC	AUC
HLA-DPA1*0103-DPB1*0201	1,404	538	0.922	0.957	0.912	0.952
HLA-DPA1*0103-DPB1*0401	1,337	471	0.929	0.962	0.914	0.958
HLA-DPA1*0201-DPB1*0101	1,399	597	0.905	0.948	0.902	0.941
HLA-DPA1*0201-DPB1*0501	1,410	443	0.868	0.954	0.865	0.950
HLA-DPA1*0301-DPB1*0402	1,407	523	0.912	0.957	0.905	0.956
HLA-DQA1*0101-DQB1*0501	1,739	522	0.791	0.901	0.802	0.907
HLA-DQA1*0102-DQB1*0602	1,629	813	0.698	0.872	0.659	0.855
HLA-DQA1*0301-DQB1*0302	1,719	386	0.723	0.813	0.729	0.833
HLA-DQA1*0401-DQB1*0402	1,701	559	0.807	0.914	0.794	0.903
HLA-DQA1*0501-DQB1*0201	1,658	549	0.802	0.902	0.809	0.902
HLA-DQA1*0501-DQB1*0301	1,689	863	0.816	0.919	0.810	0.918
H-2-IAb	660	126	0.713	0.884	0.664	0.856
H-2-IAd	379	70	0.577	0.816	0.420	0.856
DRB1*0101	7,685	4,382	0.717	0.849	0.682	0.831
DRB1*0301	2,505	649	0.708	0.859	0.671	0.836
DRB1*0302	148	44	0.601	0.800	0.266	0.627
DRB1*0401	3,116	1,039	0.659	0.841	0.609	0.817
DRB1*0404	577	336	0.663	0.838	0.595	0.784
DRB1*0405	1,582	627	0.711	0.862	0.683	0.843
DRB1*0701	1,745	849	0.729	0.861	0.732	0.860
DRB1*0802	1,520	431	0.515	0.771	0.478	0.750
DRB1*0806	118	91	0.778	0.927	0.707	0.886
DRB1*0813	1,370	455	0.740	0.881	0.719	0.867
DRB1*0819	116	54	0.608	0.809	0.334	0.661
DRB1*0901	1,520	621	0.652	0.828	0.572	0.788
DRB1*1101	1,794	778	0.770	0.879	0.749	0.868
DRB1*1201	117	81	0.787	0.909	0.694	0.848
DRB1*1202	117	79	0.783	0.916	0.682	0.849
DRB1*1302	1,580	493	0.612	0.814	0.607	0.804
DRB1*1402	118	78	0.753	0.890	0.546	0.800
DRB1*1404	30	16	0.611	0.728	0.259	0.603
DRB1*1412	116	63	0.764	0.896	0.574	0.789
DRB1*1501	1,769	709	0.677	0.831	0.629	0.803
DRB3*0101	1,501	281	0.683	0.851	0.613	0.816
DRB3*0301	160	70	0.754	0.864	0.543	0.773
DRB4*0101	1,521	485	0.693	0.846	0.687	0.840
DRB5*0101	3,106	1,280	0.760	0.882	0.740	0.865

Molecule name	#pep	#bind	NetMHCIIpan-3.0		NN-align	
			PCC	AUC	PCC	AUC
Average			0.735	0.871	0.664	0.838
<i>p</i> value			<0.0001	<0.0001		
<i>p</i> value ^{<i>a</i>}			0.002	0.021		

NN-align is the method described by Nielsen and Lund (2009); *NetMHCIIpan-3.0* is the method described here. The values in bold show the higher score for each molecule for corresponding performance measures (PCC or AUC). The *p* values for PCC and AUC are given below the first columns of PCC and AUC values respectively

#pep-number of peptide binding data available for each molecule, #bind-number of peptides that have a binding affinity stronger than 500 nM

 $^{a}_{p}$ -values of the 10 molecules characterised by <400 data points and <100 binders

Independent evaluation of the *NetMHCIIpan-3.0* method compared with the performance of the *NetMHCIIpan-2.0* predictor

Molecule name	#nen	#bind	NetMHCIInan-3.0 NetMHCIInan-2			CIInan_2 ()
Molecule name	прер	#Dillu	<u>neimin</u>			<u></u>
			PCC	AUC	РСС	AUC
DRB1_0101	717	550	0.820	0.908	0.817	0.910
DRB1_0301	703	408	0.699	0.850	0.703	0.862
DRB1_0701	682	375	0.754	0.873	0.771	0.882
DRB1_0801	838	363	0.738	0.875	0.713	0.861
DRB1_1101	813	426	0.790	0.901	0.787	0.902
DRB1_1301	803	462	0.573	0.792	0.488	0.753
DRB1_1302	765	404	0.392	0.713	0.289	0.668
DRB1_1501	758	218	0.499	0.764	0.496	0.767
DRB3_0202	726	287	0.490	0.755	0.495	0.750
DRB3_0301	782	449	0.555	0.776	0.602	0.800
DRB4_0101	778	235	0.292	0.654	0.254	0.635
DRB4_0103	764	474	0.538	0.798	0.505	0.795
DRB5_0101	731	461	0.699	0.841	0.697	0.841
Average			0.603	0.808	0.586	0.802
<i>p</i> value			0.267	1.000		

NetMHCIIpan-2.0 method is an updated version of the method proposed by Nielsen et al. (2010a). *NetMHCIIpan-3.0* is the method presented in this study. Molecule names in bold show molecules that were not part of the training set. The values in bold show the higher score for each molecule for corresponding performance measures (PCC or AUC) between the two methods. *p* values were obtained using binomial test for PCC and AUC values #*pep*-number of peptide binding data available for each molecule, #*bind*-number of peptides that have a binding affinity stronger than 500 nM