Peptide length-based prediction of peptide-MHC class II binding

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ABSTRACT

Motivation: Algorithms for predicting peptide-MHC class II binding are typically similar, if not identical, to methods for predicting peptide-MHC class I binding despite known differences between the two scenarios. We investigate whether representing one of these differences, the greater range of peptide lengths binding MHC class II, improves the performance of these algorithms.

Results: A nonlinear relationship between peptide length and peptide-MHC class II binding affinity was identified in the data available for several MHC class II alleles. Peptide length was incorporated into existing prediction algorithms using one of several modifications: using regression to pre-process the data, using peptide length as an additional variable within the algorithm, or representing register shifting in longer peptides. For several data sets and at least two algorithms these modifications consistently improved prediction accuracy.

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1 INTRODUCTION

Major histocompatibility complex (MHC) molecules, also known as human leukocyte antigens (HLA), are a vital component to the development of the immune response to pathogens (Kaufmann 2005). These molecules act as receptors for peptides derived from foreign antigens as well as self peptides and enable the long-term display of antigens on the cell surface. T cells recognize antigenic peptides in the context of MHC, and depending on the class of MHC involved, recognition can lead to the death of the presenting cell or its activation. In either case peptide-MHC binding is an important prerequisite event and has far-reaching consequences to the ensuing response.

Prediction of peptide-MHC binding therefore represents an important goal in bioinformatics, particularly as applied to immunology, and a number of computational approaches have been developed (reviewed in Buus 1999; see also Robinson *et al.* 2003 for other MHC-specific bioinformatics tools). The simplest are based on motifs, i.e. requirements for particular amino acids at positions within the peptide as determined from pool sequencing of eluted peptides (Falk *et al.* 1991, Rammensee 1995 and references therein). Such approaches have largely been superseded by algorithms using matrices to score the relative contribution of amino acids at each position within the peptide (Parker *et al.* 1994, Davenport *et al.* 1995, Marshall *et al.* 1995). Machine learning methods including hidden Markov models and artificial neural networks

have also been applied, with peptide sequence serving as input and binding/non-binding as output (Brusic and Harrison 1994, Honeyman *et al.* 1998, Mamitsuka 1998). More recently, attempts have been made to predict the structure of the peptide-MHC complex and free energy changes associated with binding (Altuvia *et al.* 1997, Rognan *et al.* 1999, Schueler-Furman *et al.* 2000, Davies *et al.* 2003, Schafroth and Floudas 2004; for a review of current structural information and nomenclature see Kaas and Lefranc 2005). It is also possible to combine some of these approaches, as Sturniolo *et al.* (1999) did using matrices to represent each pocket lining the peptide-binding groove.

Continued progress in the development of these algorithms faces a number of challenges including how to handle differences between the two classes of MHC. Most prediction algorithms were first developed in the context of peptide-MHC class I binding which involves peptides of a narrow range of lengths, usually 8-10 amino acids. These algorithms were then applied to peptide-MHC class II binding, typically with little or no modification.

Despite the fact that both classes of MHC share superficial similarities and bind a core of nine amino acids within peptides (Jones 1997), important differences exist. In particular the open-ended nature of MHC class II peptide-binding groove allows for a wide range of peptide lengths (Brown *et al.* 1993). Peptides binding MHC class II usually vary between 13 and 17 amino acids in length, though shorter or longer lengths are not uncommon (Chicz *et al.* 1992, Sercarz and Maverakis 2003). As a result peptides are hypothesized to shift within the MHC class II peptide-binding groove, changing which 9mer window (register) sits directly within the groove at any given time. In contrast the capped nature of the MHC class I peptide-binding groove does not allow variation in length or such register shifting.

Variation in peptide length may have important consequences for the binding and function of antigenic peptides (Malcherek *et al.* 1994, Vogt *et al.* 1994). For instance, Srinivasan *et al.* (1993) found that a 23mer peptide derived from cytochrome c was 32 times more immunogenic than a 10mer peptide containing the same putative binding core. A direct relationship between peptide length and binding affinity has been observed for some MHC class II alleles, but whether this holds true for most alleles remains unknown, as does an explanation for why this relationship exists (Bartnes *et al.* 1999, Fleckenstein *et al.* 1999, Arnold *et al.* 2002, Sercarz and Maverakis 2003). In addition to having more binding registers, longer peptides also possess peptide-flanking residues (PFR) which lie outside of the peptide-binding groove and may interact with the MHC class II molecule at more distal locations (Sercarz and Maverakis 2003). Whether information regarding

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peptide length, or any other peptide property lost by considering only 9mers, may aid prediction also remains unknown.

In this study we address several issues related to peptide length and binding to MHC class II. Using aggregate data that are now available from online databases, we first examine whether a relationship exists between length and affinity for several MHC class II alleles. We then attempt to incorporate length into two existing binding algorithms in a number of ways, including using regression to pre-process the data, treating length as an additional variable within the algorithms, and deriving a formula to more accurately represent register shifting (Figure 1). We show that improvements to more than one current algorithm for predicting peptide-MHC class II binding are possible with relatively simple amendments. We also comment on which mechanisms are likely to be affecting binding as peptide length increases.



Fig. 1. Schematic of modifications made to existing algorithms to incorporate peptide length. Modification 1, Alternative Modification 1, and Modification 2 are abbreviated Mod. 1, Alt. 1, and Mod. 2. Also shown are examples of sources of data, algorithms used to remove homologous sequences from data, and algorithms to predict peptide-MHC class II binding.

2 SYSTEMS AND METHODS

Data sources Peptide data sets used in this study are available from the AntiJen database (http://www.jenner.ac.uk, Blythe et al. 2002) and can be downloaded using the perl LWP::Simple module. Other peptide-MHC databases listing affinities are also available, including the Immune Epitope Database (currently in beta version at http://www.immuneepitope.org, Peters et al. 2005), but were not used in this study. Our data sets comprised the sequences and IC₅₀ values of peptides binding the MHC class II alleles HLA-DRB1*0101, -DRB1*0401, and -DRB1*1501 from AntiJen. IC50 refers to the concentration of peptide required to inhibit 50% of reporter peptide-MHC binding. When more than one IC₅₀ measurement was available for a given peptide-MHC complex, the first measurement listed was used, unless otherwise indicated. IC50 values were converted into pIC50 using the formula $pIC_{50} = -\log IC_{50}$ where IC_{50} has units of molar. Homologous sequences and their IC50 measurements were removed using Unique-Prot (Mika and Rost 2003). Other algorithms for removing homologous sequences are also available, including Hobohm 1 and Hobohm 2 (Hobohm et al. 1992), but were not used in this study. The data sets were of the following sizes (before/after filtering by UniqueProt): DRB1*0101 (464/303), Regression of binding affinity versus peptide length Both parametric and nonparametric fits were made to plots of affinity vs. length in the data. Parametric fits were made with one, two, and three fitted parameters (linear, quadratic, and cubic, respectively) using the open-source statistical program R (http://www.R-project.org, R Development Core Team 2005) and the function lm. Nonparametric local regression fits were made using the R function loess with default settings (Cleveland and Devlin 1988). To evaluate the quality of the fits analysis of variance was performed using the R function anova. An F statistic was generated which we used to compare linear with nonlinear parametric fits (Motulsky and Christopoulos 2004).

Nonparametric local regression fits were evaluated using a permutation test. In this test each pIC_{50} value was reassigned to a different peptide sequence at random, and a loess fit was re-derived for the shuffled values. This was repeated 1000 times, and the smallest 25 (2.5%) and largest 25 (2.5%) fitted values at each length were excluded. The local regression fit to the original, non-shuffled data set was then compared to the remaining 95% of permuted values at each length and was determined to be significant if it fell outside of this interval.

Simulations of register shifting To simulate the effects of register shifting on peptide-MHC class II binding affinity over a range of peptide lengths, we derived a formula for the expected value of the affinity of a single hypothetical peptide with multiple registers:

$$E[K(X)] = \sum K(x_i) p(x_i)$$
[1]

where K(X) is the equilibrium association constant, or affinity, of a peptide X, $K(x_i)$ is the affinity of a complex with a single register x_i , and $p(x_i)$ is the probability of register x_i occurring. We assume that $p(x_i)$ can be approximated by the proportion of complexes having register x_i :

$$\mathbf{p}(x_i) = \mathbf{N}(x_i) / \sum \mathbf{N}(x_i).$$
[2]

where $N(x_i)$ denotes the number of complexes having register x_i and the sum is taken over all possible registers. Belmares and McConnell (2001) found that the kinetics of shifting between two registers could be accurately represented as $x_1 \leftrightarrow P + M \leftrightarrow x_2$ where P and M are peptide and MHC, respectively. Based on this result, at equilibrium $[x_1] = K(x_1)[P][M]$ and $[x_2] = K(x_2)[P][M]$. Because both x_1 and x_2 exist in the same solution, it follows that:

$$N(x_1) / [N(x_1) + N(x_2)] = K(x_1) / [K(x_1) + K(x_2)].$$
 [3]

More generally,

$$N(x_i) / \sum N(x_i) = K(x_i) / \sum K(x_i).$$
[4]

Combining Equations 1, 2, and 4, we obtain the following result for the expected affinity of a given complex when multiple registers are available:

$$E[K(X)] = \sum K(x_i)^2 / \sum K(x_i).$$
 [5]

This result can also be applied to log-transformed measures of affinity such as log K(X). Henceforth we refer to Equation 5 (or its log-transformed counterpart) as the equilibrium-based formula for reconciling multiple registers.

We assume that every overlapping 9mer window within a peptide can result in binding to MHC and therefore set the lower and upper limits of summation at 1 and l - 8, respectively, where l represents peptide length and is varied between 9 and 25, the shortest and longest lengths typically observed in our data sets. $K(x_i)$ was generated from a lognormal distribution with mean $10^{7.5}$ and standard deviation $10^{0.5}$, based on the observation that most values for the equilibrium dissociation constant K_D of peptide-MHC binding fall in the range of 10^{-7} - 10^{-8} M (McFarland and Beeson 2002). Moreover, a lognormal distribution was chosen based on the equation for standard free energy change, $\Delta G^\circ = -RT \ln (1/K_D)$ where R and T are the gas constant and temperature, respectively (Eisenberg and Crothers 1979), and the assumption that free energy change for peptide-MHC binding is normally distributed. For each value of *l* between 9 and 25, a set number of values were generated (in our case, either 10 or 100), resulting in a scatter plot of simulated pIC_{50} values versus length. A curve was then fit to this plot using local regression (the loess function in R) with default settings.

Peptide-MHC binding affinity prediction Two algorithms were selected to generate baseline predictions against which the effects of modifications based on length could be compared. One of these algorithms was the iterative self-consistent (ISC) partial-least-squares (PLS) algorithm of Doytchinova and Flower (2003). We implemented this matrix-based algorithm for predicting peptide-MHC binding affinity in perl and R. Briefly, this algorithm uses partial least squares regression to identify underlying factors (also known as latent variables) relating multiple predictor variables to an outcome variable. In the case of peptide-MHC binding, 180 predictor variables were used to denote the presence or absence of the 20 possible amino acids within each 9mer window, and the outcome variable was binding affinity as pIC_{50} .

The initial steps of the algorithm were performed using perl scripts: splitting each data set into training and test sets; generating all possible 9mers for each training set peptide; selecting only those 9mers having position 1 anchor residues (F, I, L, M, V, W, and Y); and converting 9mers thus selected into bit strings. PLS regression was then performed in R using the bit-encoded 9mers and their corresponding pIC50 values. PLS is available for R as the pls.pcr library (available at http://cran.r-project.org) and was called from within a perl script using the IPC::Open2 module. Default settings were used for PLS; however, some options in the commercial software used by Doytchinova and Flower (2003) were not available in R, namely scaling method and column filtering. Subsequent steps in the algorithm were performed using additional perl scripts: selecting those 9mers in the training set yielding predicted pIC50 values closest to experimental pIC₅₀ values during cross-validation and repeating the algorithm until the selected set of 9mers matched the previously selected set, i.e. when self-consistency was achieved. For computational expediency we limited the number of PLS iterations for any given peptide to 10. At that point the final PLS model was extracted and used to generate predictions on the test set.

For test set peptides having more than one 9mer with an anchor residue in position 1, multiple predictions were generated and a rule was needed to make a final prediction. One option is to assume only one register predominates and to take the highest score from among the predictions. More complicated rules are also possible such as the combination rule of Doytchinova and Flower (2003) whereby the mean of the pIC_{50} predictions is chosen if they fall within a one log range; otherwise, the highest is chosen.

To measure the performance of the algorithm we used five-fold crossvalidation (5x-CV), setting aside one-fifth of each data set to use as a test set and using the other four-fifths as the training set. This process was repeated on the same data set four additional times until a prediction was made for each peptide in the data set and complete coverage was achieved. (This instance of cross-validation was independent of the leave-one-outcross validation used in the ISC-PLS algorithm.) The accuracy of each set of predictions was scored by calculating the area under receiver operating characteristic curve (A_{ROC}). This calculation can be done in R using the prediction and performance functions of the ROCR library. By repeating each 5x-CV multiple times, we were able to calculate the standard error of the A_{ROC} scores which could then be used to determine whether two mean A_{ROC} scores significantly differed by Student's *t* test.

A second algorithm that was selected was the TEPITOPE algorithm of Sturniolo *et al.* (1999). In this algorithm amino acid-binding profiles are generated for each pocket within the peptide-binding groove, and these profiles are combined according to MHC sequence. We did not regenerate these matrices but rather used the matrices available on the ProPred website (http://www.imtech.res.in/raghava/propred, Singh and Raghava 2001). Using the appropriate matrix a sum was calculated for each peptide in a selected AntiJen data set. To this value we added an approximation of the binding affinity of an all-alanine 9mer ($pIC_{50} = 6.169$, Doytchinova and Flower 2003) generating a final prediction. Performance was scored by calculating the A_{ROC}.

Incorporating length into existing prediction algorithm Peptide length was incorporated into the ISC-PLS algorithm using one of three modifications. In Modification 1 (Mod. 1) a local regression fit was first made to the peptide lengths and pIC50 measurements in each training set. (In the event that the pIC_{50} value for either the shortest or the longest length peptide was excluded from the training set but included in the test set, a local regression fit at that length could not be generated; instead, we assigned the average fitted values at the remaining lengths.) The value of the fit was then subtracted from the original pIC_{50} value for each peptide, and the resulting difference, i.e. the residual, was then used in place of the original pIC_{50} value. The ISC-PLS algorithm was performed as described earlier providing initial predictions on the test set. To these predictions the value of the regression fit was added yielding final predictions. Alternatively, in Alternative Modification 1 (Alt. 1), peptide length was appended as the 181st predictor variable to the bit-encoded training set and test set 9mers. The remainder of the algorithm was then performed as described earlier. Finally, in Modification 2 (Mod. 2) the formula derived to represent register shifting (Equation 5) was used to reconcile predictions made on multiple candidate 9mers, i.e. registers, within a test set peptide. This modification occurred at the last stage of the ISC-PLS algorithm and was used in place of the combination rule described above.

Only Mod. 2 was used to incorporate length into the TEPITOPE/ProPred algorithm. When TEPITOPE/ProPred is applied to peptides with multiple registers, the highest score among the different registers is typically taken to be the score of the entire peptide (Brusic *et al.* 1998, Nielsen *et al.* 2004, Murugan and Dai 2005). We reconciled individual register scores using the equilibrium-based formula (Equation 5) but did not regenerate the pocket profiles and therefore did not apply Mod. 1 or Alt. 1 in this case.

3 IMPLEMENTATION

3.1 Peptide length significantly affects binding affinity to MHC class II

To determine the nature of the relationship between peptide length and peptide-MHC class II binding affinity, we derived a number of regression fits to binding data for several MHC class II alleles from the AntiJen database. In all cases homologous sequences were first removed from the data sets using a pre-filtering algorithm, UniqueProt (Mika and Rost 2003). Parametric fits were then made based on polynomials with one, two, or three fitted parameters (linear, quadratic, and cubic, respectively). Analysis of variance from these fits showed that for these MHC class II alleles the nature of the relationship was most likely nonlinear (Table 1). A quadratic or cubic fit resulted in a significant reduction in sum of squares in all three cases at the 0.05 level.

 Table 1. Evidence of non-linear relationships in length-affinity data for several MHC class II alleles

	DRB1*0101	DRB1*0401	DRB1*1501
Quadratic, F	11.745 (<0.001)	8.575 (0.004)	3.670 (0.057)
Cubic, F	5.849 (0.016)	0.708 (0.401)	4.871 (0.028)

F statistics are shown for analysis of variance results with p values in parentheses.

To better characterize the apparent nonlinearities in the lengthaffinity data we then made nonparametric fits to the data and analyzed the fits. Local regression was used to make nonparametric fits, and analysis was done using a permutation test. In this test binding affinities were reshuffled among peptide lengths to create 1000 new data sets, and a local regression fit was re-derived for each data set. If the fit to the original data fell outside of the middle 95% of permutation fits at any particular length, the nonlinearity at that length was determined to be significant. In each data set we found that the nonlinearity between length and affinity was significant at one or more lengths (Figure 2). Lengths associated with strongest affinity could be identified, as could lengths associated with weakest affinity. For example, for DRB1*0401 affinity appeared strongest for peptides of 12 amino acids and weakest for peptides of 20 amino acids. When the data sets were combined and the local regression fits were regenerated, the same trends were seen (Figure 2D): shorter peptide lengths, of approximately 12 amino acids, were associated with higher affinity, while longer peptide lengths, of approximately 20 amino acids, were associated with lower affinity.



Fig. 2. Local regression fits of peptide-MHC class II binding affinity versus peptide length for three HLA data sets: *A*, DRB1*0101; *B*, DRB1*0401; *C*, DRB1*1501; and *D*, the three data sets combined. 95% boundaries of permutation distributions are shown (dotted) with fits to the original, non-shuffled data (solid).

Nonlinearities may have been present in the length-affinity data for several reasons, including the ability of peptides to shift registers within the MHC class II peptide-binding groove. To simulate the effect of register shifting on the mean affinity observed for peptides of different lengths, we used a simple statistical model based on two assumptions: first, that longer peptides are likely to contain more registers than shorter peptides, and secondly, that the measured affinity of a given peptide-MHC complex approximates the weighted average of the affinities of all the registers in a peptide (Equation 5). For a simulated peptide of a given length *l*, the affinities of l - 8 registers were generated and averaged. This process was repeated until the average affinities of either 10 or 100 peptides at each length (i.e. each value of *l*) were obtained, resulting in data sets of two sizes (one of the same magnitude as those typically obtained from databases, the other an order of magnitude larger). At this point a regression curve was derived (Figure 3). For the larger sized data set the fitted curve was nonlinear and monotonically increasing (Figure 3A). The same trend was seen in the smaller data set; in this case, however, deviations were also possible, resulting in maxima at mid-length peptides (Figure 3B). Together these results suggest that register shifting may be one mechanism behind the nonlinearities in the length-affinity relationship from experimental data sets.



Fig. 3. Statistical simulations of the effects of register shifting on MHC class II binding affinity over a range of peptide lengths: *A*, for a 1700-peptide data set; *B*, for a 170-peptide data set. Curve fits by local regression are shown overlaid.

We also estimated the lengths of the amino- and carboxylterminal portions of each peptide extending outside of the MHC class II peptide-binding groove to determine if particular lengths at either end of the peptide were favorable or unfavorable for binding. 9mer cores were identified by position 1 anchor residues (F, I, L, M, V, W, Y), and the lengths remaining at each end were calculated. Local regression fitting and permutation testing were done as with overall peptide length. In most cases fits to amino- and carboxyl-terminal peptide extensions were determined to be significant at one or more lengths (Figure 4 and additional data not shown). In comparing fits we found that extensions of 2-4 amino acids at the amino terminus and extensions of 1-2 at the carboxyl terminus generally appeared favorable for binding (Figure 4 and additional data not shown). Likewise, longer extensions (8 and 10 amino acids at the amino and carboxyl termini, respectively) generally appeared unfavorable for binding (Figure 4 and additional data not shown). We also found that in at least some cases fits to overall peptide length could be decomposed into amino- and carboxyl-terminal contributions. For example binding to DRB1*0401 was strongest when amino and carboxyl termini were 2 and 1 amino acids, respectively (Figure 4). Together with the 9mer core, these lengths sum to match the overall length associated with strongest binding, 12 amino acids (Figure 2B).



Fig. 4. Local regression fits of peptide-MHC class II binding affinity versus lengths of portions of the peptide extending outside of the peptide-binding groove for the HLA-DRB1*0401 data set: *A*, at the N terminus and *B*, at

the C terminus. 95% boundaries of permutation distributions are shown (dotted) with fits to the original, non-shuffled data (solid).

3.2 Modifying algorithms to account for peptide length consistently improves performance

We incorporated peptide length into two peptide-MHC class II binding prediction algorithms in one of three ways. First, as a preprocessing event (Mod. 1 in Figure 1) a local regression fit was made for affinity vs. length in the training/fitting data and the value of the fit was subtracted from each affinity measurement. The resulting residuals were used in place of the original pIC_{50} values in the training set. After the algorithm was used to make initial predictions for the target set peptides, the value of the regression fit for each target set peptide length was added to yield final predictions. Alternatively (Alt. 1 in Figure 1) length was also incorporated directly into the existing algorithm as an additional variable (in the case of ISC-PLS, as the 181st variable). Training/fitting was then performed as published, and predictions were made on test set peptide sequences and peptide lengths. Lastly we used a formula derived from the equilibrium-based statistical model to reconcile predictions made by existing algorithms on multiple registers within the peptide (Mod. 2 in Figure 1). We point out that Mod. 1 and Alt. 1 are similar modifications that both consider peptide length directly (by fitting length as a discrete variable); in contrast Mod. 2 considers binding registers (i.e. 9mers with a valid position 1 anchor) and the relationship among them. Therefore, Mod. 1 and Alt. 1 are not used together, although either can be used with Mod. 2.

Incorporating peptide length by one or more modifications into the ISC-PLS algorithm improved the performance of the algorithm for all alleles examined (Table 2). Performance was measured by area under receiver operating characteristic curves (A_{ROC}) when a threshold of 500 nM was used to differentiate binding from nonbinding affinities (Sette et al. 1994). The performance of ISC-PLS in conjunction with a combination rule (mean if less than one order range; highest otherwise) to reconcile register predictions was used as a baseline (Doytchinova and Flower 2003). Taking the highest scoring register to be representative of the entire peptide was also done as a reference. In general using any of three modifications resulted in increases in algorithm performance. However the modification resulting in the greater increase differed by MHC class II allele. In the case of DRB1*0101, deriving a regression fit (Mod. 1) resulted in significantly greater improvements than either using length as an additional variable or using the equilibrium-based formula to reconcile register predictions. In the case of DRB1*0401, all three modifications resulted in the same magnitude of increase in performance. Finally in the case of DRB1*1501 only an application of both the regression fit (Mod. 1) and the equilibrium-based formula (Mod. 2) resulted in the greatest increase in performance. Differences in which modifications resulted in the greatest increase in performance may be suggestive of alleleor data set-specific mechanisms behind the length-affinity relationships.

Table 2. Binding prediction accuracy of ISC-PLS algorithm for different

 MHC class II alleles when peptide length was incorporated

DRB1*0101	ISC-PLS	Mod. 1: Re- gression fit	Alt. 1: Length as variable
Combination rule	0.615 ± 0.009^{1}	$\textbf{0.754} \pm \textbf{0.009}$	0.690 ± 0.013
Highest scoring register	0.652 ± 0.008	$\textbf{0.758} \pm \textbf{0.006}$	0.705 ± 0.013
Mod. 2 : Equilib- rium formula	0.709 ± 0.005	$\textbf{0.770} \pm \textbf{0.009}$	$\textbf{0.752} \pm \textbf{0.003}$
DRB1*0401	ISC-PLS	Mod. 1: Re- gression fit	Alt. 1: Length as variable
Combination rule	0.730 ± 0.007^{1}	$\textbf{0.741} \pm \textbf{0.010}$	$\textbf{0.749} \pm \textbf{0.009}$
Highest scoring register	0.732 ± 0.015	$\textbf{0.750} \pm \textbf{0.006}$	$\textbf{0.751} \pm \textbf{0.005}$
Mod. 2: Equilib- rium formula	$\textbf{0.757} \pm \textbf{0.008}$	$\textbf{0.757} \pm \textbf{0.004}$	$\textbf{0.754} \pm \textbf{0.008}$
DRB1*1501	ISC-PLS	Mod. 1: Re- gression fit	Alt. 1: Length as variable
Combination rule	0.574 ± 0.009^{1}	0.596 ± 0.015	0.584 ± 0.020
Highest scoring register	0.575 ± 0.021	0.626 ± 0.014	0.603 ± 0.011
Mod. 2 : Equilib- rium formula	0.609 ± 0.019	$\textbf{0.677} \pm \textbf{0.014}$	0.609 ± 0.018

Five-fold cross-validation (5x-CV) was used and repeated five times. Mean A_{ROC} scores between predicted and experimentally determined pIC_{50} values are shown with standard errors of the mean. Highest scores are shown in bold with multiple scores in bold if pair-wise differences were not statistically significant. A threshold of 500 nM (Sette 1994) was used to distinguish binding from non-binding peptides. ¹The ISC-PLS algorithm with combination rule (Doytchinova 2003) was used as a baseline prediction.

We also incorporated peptide length into the TEPITOPE/ProPred algorithm (Sturniolo *et al.* 1999) and without re-deriving the pocket-specific matrices that define that algorithm found that increases in performance could be obtained by use of the equilibrium-based formula alone (Table 3). Typically in applications of TEPITOPE/ProPred to MHC class II, predictions on multiple registers are reconciled by taking the highest scoring register to be representative of the whole peptide (Brusic *et al.* 1998, Nielsen *et al.* 2004, Murugan and Dai 2005). We therefore used this rule to generate baseline predictions against which we could compare the performance of the equilibrium-based formula. Applying the formula for register shifting increased algorithm performance for all three data sets examined.

	ProPred: DRB1*0101	ProPred: DRB1*0401	ProPred: DRB1*1501
Combination rule	0.685	0.741	0.669
Highest scoring register	0.667 ¹	0.754 ¹	0.635 ¹
Mod. 2: Equilib- rium formula	0.702	0.764	0.680

Table 3. Binding prediction accuracy of ProPred algorithm for different

 MHC class II alleles when peptide length was incorporated

Matrices were obtained from the ProPred website and used to calculate a score for each register within a peptide. To each score the approximate affinity of an all-alanine 9mer to MHC was added ($pIC_{50} = 6.169$, Doytchinova and Flower 2003). A_{ROC} scores between predicted and experimentally determined pIC_{50} are shown, using a threshold of 500 nM (Sette *et al.* 1994) to distinguish binding from non-binding peptides. ¹Highest ProPred-predicted scores from all eligible registers were used as baseline predictions following recent precedents (Brusic *et al.* 1998, Nielsen *et al.* 2004, Murugan and Dai 2005).

We also investigated whether our modifications might be applied to alleles for which fewer data exist. In analyzing the data for two other alleles, DRB1*0404 and DRB1*0405, we found no significant nonlinearities in regression fits of length versus affinity (Supplementary Data). Consistent with the results of these fittings, we observed no increase in performance after applying either Mod. 1 or Alt. 1 to the ISC-PLS algorithm when training sets were derived from these data sets (Supplementary Data). An increase in performance was observed, however, for the larger of the two data sets using Mod. 2 (Supplementary Data). These results suggest that our proposed modifications, like matrix-based prediction algorithms, are subject to limitations based on the size of the training set.

4 DISCUSSION AND CONCLUSION

Information is typically lost during the prediction of peptide-MHC class II binding because most algorithms focus exclusively on 9mers within the peptide. An underlying assumption is that properties of the parent peptides that cannot be captured in their 9mers are irrelevant. This assumption may be true for MHC class I binding which involves peptides of nine amino acids almost exclusively but may not be true for MHC class II binding. Peptides that bind MHC class II are variable in length and may contain segments that extend past the ends of the peptide-binding groove, also known as peptide-flanking residues or PFR (Brown et al. 1993). PFR-MHC interactions may in turn affect peptide-MHC binding in a manner that is consistent and useful to prediction. Longer peptides also allow for register shifting, i.e. the ability of peptides to bind MHC using different core 9mers. PFR-MHC interactions and register shifting represent two possible mechanisms by which variability in peptide length affects affinity to MHC class II.

In this study we found that nonlinear relationships exist between peptide length and peptide-MHC class II binding affinity in a number of aggregate data sets available online. When these nonlinearities were examined in more detail, they were found to be significant at several lengths, suggesting some lengths were more favorable for binding than others. This is consistent with the data from a number of experimental studies (Malcherek *et al.* 1994, Vogt *et al.* 1994, Bartnes *et al.* 1999, Fleckenstein *et al.* 1999). In these studies affinity was generally found to increase with length up to the longest lengths examined, typically between 15 and 17 amino acids. In our simulations register shifting was found to be one mechanism that could account for the direct relationship between length and binding affinity. However, our analysis of aggregate data sets suggests that additional mechanisms also contribute to the effect of length on affinity. For example, register shifting alone cannot explain why certain lengths at the amino and carboxyl termini are advantageous or disadvantageous for binding DRB1*0401. In this case other mechanisms such as hypothesized PFR-MHC interactions that are either attractive or repulsive may also be playing a role (Sercarz and Maverakis 2003).

Incorporating peptide length into existing binding prediction algorithms by one or more of our modifications consistently improved performance across MHC class II alleles. Three modifications were used—one at the level of the training set data (Mod. 1), another within the algorithm itself (Alt. 1), and the last after 9mer predictions were generated (Mod. 2)-and all resulted in performance gains over reference algorithms ISC-PLS and TEPITOPE/ProPred. Baseline A_{ROC} scores for two different algorithms varied between 0.57 and 0.73. By comparison A_{ROC} scores for modified algorithms varied between 0.68 and 0.77, consistent range of with the scores listed in MHCBench (http://www.imtech.res.in/raghava/mhcbench/). The modification resulting in the largest performance increase differed by allele, and this may in part reflect differences in the mechanisms by which length affects affinity. For DRB1*0401, for example, using the formula for register shifting resulted in performance gains that were statistically indistinguishable from those obtained using other modifications. For DRB1*0101, however, modifications based on regression modeling resulted in significantly greater performance increases. These data therefore support roles for both register shifting and other mechanisms.

Previous studies have provided indirect evidence that accounting for variability in peptide length could improve prediction. Godkin *et al.* (1998), for example, found that matrices based on 15mers generally outperformed matrices based on shorter lengths, showing the usefulness of considering information outside of the core 9mer. Likewise, Bui *et al.* (2005) have proposed deriving a separate matrix for each length of peptide (Bui *et al.* 2005). Despite the suggestion that explicit consideration of peptide length could improve binding prediction (McFarland and Beeson 2002), to our knowledge no previous study has implemented this idea. Our results affirm the use of peptide length in binding prediction. In addition our modifications are sufficiently general that they could be incorporated into other current algorithms based on scoring 9mers.

Thus far experimental evidence of either register shifting or PFR-MHC interactions has involved only a small sampling of MHC class II alleles and been of indeterminate generality. For example, register shifting has been demonstrated to occur with alleles I-A^d and I-A^u in mice and DR2 in humans (McFarland *et al.* 1999, Li *et al.* 2000, Seamons *et al.* 2003, Bankovich *et al.* 2004). Solved structures exist for a somewhat wider array of alleles, including I-A^d and I-A^k in mice and DR1, DR3, and DR4 in humans (see McFarland and Beeson 2002 for a review). Although these structures show the presence of PFRs in peptide-MHC class II

complexes, they fail to capture the dynamics of either register shifting or PFR-MHC interactions.

Our analysis of regression fits to different aggregate binding data sets suggests that longer PFRs (i.e. in peptides longer than approximately 16 amino acids) may generally be deleterious to binding. At the same time, however, PFRs of a certain minimum length increase the probability of a peptide having multiple binding registers which, our simulations show, increases overall binding affinity. An optimal peptide length for binding each MHC class II variant may therefore exist. Further computational analysis of aggregate data sets may provide a complement to more direct, observation-based studies in continuing to elucidate the role of peptide length in MHC class II binding.

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