Combinatorial Peptide Libraries and Biometric Score Matrices Permit the Quantitative Analysis of Specific and Degenerate Interactions Between Clonotypic TCR and MHC Peptide Ligands

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The interaction of TCRs with MHC peptide ligands can be highly flexible, so that many different peptides are recognized by the same TCR in the context of a single restriction element. We provide a quantitative description of such interactions, which allows the identification of T cell epitopes and molecular mimics. The response of T cell clones to positional scanning synthetic combinatorial libraries is analyzed with a mathematical approach that is based on a model of independent contribution of individual amino acids to peptide Ag recognition. This biometric analysis compares the information derived from these libraries composed of trillions of decapeptides with all the millions of decapeptides contained in a protein database to rank and predict the most stimulatory peptides for a given T cell clone. We demonstrate the predictive power of the novel strategy and show that, together with gene expression profiling by cDNA microarrays, it leads to the identification of novel candidate autoantigens in the inflammatory autoimmune disease, multiple sclerosis. The Journal of Immunology, 2001, 167: 2130–2141.

The CD8+ and CD4+ T lymphocytes recognize short peptides of 8–10 and 12–16 aa in the context of self MHC class I and class II molecules, respectively (1, 2). During the last 15 years, this central process of cellular immune responses has received enormous attention and has been dissected using a vast array of different immunological and biochemical techniques. A quantitative analysis of the interaction between TCR and their MHC peptide ligands would be an important basis for the design of vaccines and therapeutic approaches to immune-mediated, infectious, and neoplastic diseases.

Because it has been difficult to describe the trimolecular complex in its entirety, experiments initially focused on the interaction between peptide and MHC molecules. Structural studies of MHC class I and class II molecules complexed with antigenic peptides disclosed that the latter bind in a linear fashion (3). Sequencing of peptide pools and of individual self peptides eluted from MHC molecules (4, 5) together with systematic binding analyses (6, 7) have provided experimental data for the definition of MHC-binding motifs (8–12) and the development of MHC peptide-binding models. A combination of positive and negative influences from amino acid side chains in the antigenic peptide has been shown to determine the interaction between peptide and MHC molecules (13). Indeed, the assumption of independent contribution of each amino acid side chain in the peptide sequence to MHC binding has been used to develop quantitative methods that predict peptide binding to MHC alleles (8, 14–16). More recently, elegant neural network approaches have been used to further refine the prediction of peptide binding to MHC (17–20). Based on the fact that a subset of MHC-binding peptides are also T cell epitopes (21, 22), MHC binding has been used to predict candidate T cell epitopes in bulk T cell populations, such as those contained in the peripheral blood (12, 19). However, to dissect and predict precisely the interaction of all three components of the trimolecular complex has until now been a difficult undertaking. Therefore, the quantitative study of MHC peptide recognition by single TCR has remained a largely unsettled issue.

The specificity of the trimolecular complex interaction has been studied using individual substitution analogues. Although initial studies showed that some amino acids in the antigenic peptide sequence are necessary for recognition by the TCR (primary TCR contacts) and others can tolerate conservative substitutions (secondary contacts) (23, 24), the systematic use of single and multiple amino acid-substituted peptides has shown that all amino acid side chains can contribute to peptide recognition in a largely independent manner (25). In extreme cases, this can lead to recognition of peptides with entirely different amino acid sequences by the same TCR (25).

The development of soluble- and bead-bound combinatorial peptide libraries in various formats representing millions to trillions of peptides has emerged as a powerful approach to both T
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cell epitope determination and the analysis of TCR specificity and flexibility, as recently reviewed (26, 27). Recent studies (28–32) of T cell clones (TCC)\(^9\) demonstrated the efficacy of using positional scanning synthetic combinatorial libraries (PS-SCL) for identifying target Ags and highly active peptide mimics. However, it was technically impossible to fully use this technology without the development of quantitative methods for predicting the stimulatory potential of peptides based on data from these complex libraries.

We report in this study a new strategy that combines data acquisition with PS-SCL and analysis with a quantitative scoring matrix to identify agonist peptides for clonotypic TCR of known and unknown specificity. Peptides can be identified from database searches with unprecedented efficiency and ranked according to a score that is predictive of their stimulatory potency. To our knowledge, this is by far the most efficient available approach to identify stimulatory peptides for individual TCR and predict their actual stimulatory potency with relatively high accuracy. While further improvements of this strategy will be pursued, we have developed a tool for the identification of potential T cell epitopes, the design of vaccines, and the quantitative analysis of TCR degeneracy. Finally, we demonstrate how the search results from the above prediction strategy can be related to tissue-specific expression profiles determined by cDNA microarray assays to identify candidate peptides that are derived from proteins that are overexpressed in a diseased tissue, i.e., the brain in multiple sclerosis (MS), and are thus available for the expansion of autoreactive T cells.

Materials and Methods

T cell clones

TCC were established from peripheral blood or cerebrospinal fluid (CSF) lymphomononuclear cells by a split-well technique, as previously described (33, 34). TCC GPSF11 was established from PBMC of a patient with MS using influenza virus hemagglutinin (HA) peptide (306–318) (PKYYVKQNTLKLAT, single letter amino acid code) as an Ag. The TCC is restricted by DRB1\(^*\)0404. TCC TL3A6 was established with myelin basic protein (MBP) from PBMC of a patient with MS and recognizes the immunodominant epitope MBP\(_{22–34}\) (VHFKRKNYVTTRPG) in the context of DR2a (DR\(_B1\) + DRB\(_5\)\(^*\)0101). The TCC has been extensively characterized for recognition of numerous altered peptides derived from MBP\(_{22–34}\) as well as other molecular mimics (25, 31, 32, 35, 36). The TCR usage is TCRAV18 and TCRBV5S1. TCC CSF-3 was established with a lystate of Borrelia burgdorferi from the CSF of a patient with chronic Lyme disease, as described (34). The TCC recognizes several B. burgdorferi-derived as well as human peptides in the context of DR2b (DR\(_B1\) + DRB\(_1\)\(^*\)0101). The TCR usage is TCRAV13S2 and TCRBV14S1.

Peptides and peptide combinatorial libraries

Peptides were synthesized by the simultaneous multiple peptide synthesis method (37) and characterized using HPLC and mass spectrometry. A synthetic N-acetylated, C-amide l-aminoc acid combinatorial peptide library in a positional scanning format (PS-SCL; 200 mixtures in the O\(_X\) format, in which O represents one of the 20 l-amino acids, and X represents all of the natural l-amino acids, except cysteine) was prepared as described (38).

Proliferative assays

The proliferation of TCC in response to PS-SCL or individual peptides was tested by seeding in duplicate 2 \(\times\) 10\(^4\) T cells, 5 \(\times\) 10\(^4\) irradiated PBMC with or without mixtures from PS-SCL or peptide. Proliferation was measured by \(^{3}H\)thymidine (Amersham, Arlington Heights, IL) incorporation (32).

Statistics analysis and model building

A positional scoring matrix was generated by assigning a value of the stimulatory potential to each of the 20 defined amino acids in each position. The score \(S_i\) for each amino acid \(i\) at each position \(j\) was calculated as follows:

\[
S_i = \frac{L_i - B}{\sqrt{(std(L))^2 + (std(B))^2}}
\]

where \(L_i\) equals the mean of replicate experimental measurements (cpm), \(B\) stands for background noise, \(std(L_i)\) denotes the smoothed estimate of the SD for each measurement using a locally weighted regression smoothing technique (S-plus package) based on the assumption that the SD is dependent on level of response. We call this the Z-index score due to its similarity to statistical Z ratios of means divided by their SE values.

In an alternative score called stimulation index (S-index), we generated the score in each position by using the mean of duplicate cpm values in the presence of mixtures from the PS-SCL fractions divided by the mean of duplicate values in the absence of mixtures from the PS-SCL. The S-index score appeared preferable when the PS-SCL spectrum of the cpm value was more clearly defined.

Under the assumption of independent contribution to stimulation, the predicted stimulatory potential of given peptide is the sum of the scores in each position. A 10-mer peptide sequence can be represented by a 20 \(\times\) 10 matrix of 0s and 1s (each position). A 10-mer peptide sequence can be represented by a 20 element vector (S-index) which can be ranked according to the score in each position by using the mean of duplicate cpm values in the presence of mixtures from the PS-SCL fractions divided by the mean of duplicate values in the absence of mixtures from the PS-SCL. The S-index score appeared preferable when the PS-SCL spectrum of the cpm value was more clearly defined.

Database search

We wrote a Perl script to systematically search the GenPept database. A window with the same length of peptide as used in the PS-SCL was applied to slide over the available translated protein-coding sequences. The sum of the scores within the window was used as a ranking criterion. All peptides with scores higher than a threshold were output into a file. The threshold was chosen based on the statistical significance of the peptide score, compared with that for a random peptide. Those peptides were then sorted. Redundant peptides were removed. The database search can also be restricted to specific organisms (e.g., Homo sapiens or Influenza virus).

Statistical significance

We developed a statistical significance test of the hypothesis that the score for a peptide is no greater than would be expected if the peptide were obtained from 10 random draws of amino acids. Under the null hypothesis, it is not assumed that all amino acids are equally likely, but rather the relative frequencies \(f_1, f_2, . . . , f_{20}\) are derived from the database being searched. Under the null hypothesis, the distribution of S will be approximately normally distributed. The mean and the variance of this null distribution can be expressed as:

\[
m = \frac{\sum f_i \sum S_i}{\sum f_i}
\]

\[
var = \frac{\sum f_i S_i^2}{\sum f_i} - m^2
\]

where \(f_i\) is the frequency of each amino acid, and \(S_i\) is the score of each amino acid.

The variance can be shown to equal:

\[
var = \sum_{i=1}^{20} f_i S_i^2 + \sum_{j=1}^{9} \sum_{j+1}^{10} m_i m_j - m^2
\]

The statistical significance of any score \(S\) can be approximated as

\[
p = \Phi \left( \frac{m - S}{\sqrt{var}} \right)
\]

in which \(\Phi\) denotes the standard normal distribution function. However, this significance level does not account for the number of 10-mer sequences contained in the database.

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\(^9\) Abbreviations used in this paper: TCC, T cell clone; PS-SCL, positional scanning synthetic combinatorial libraries; CSF, cerebrospinal fluid; HA, hemagglutinin; MBP, myelin basic protein; MS, multiple sclerosis; S-index, stimulation index.
Analysis of gene expression using cDNA microarrays

Brain tissue was obtained at autopsy from two MS patients. Patient W was a 46-year-old male with primary progressive MS (39); patient R was a 46-year-old female with relapsing-remitting MS. Normal white matter was dissected, postmortem, from three nondiseased brains. RNA extracted from these three normal white matter samples was pooled, in equal amounts, for use in hybridization experiments. Lesions were identified by H&E and Luxol fast blue-periodic acid Schiff staining of paraffin-embedded sections. Further characterization of lesions was performed using immunohistochemistry for cell-specific Ags. All staging of lesions was performed as previously described (40). From the first patient, patient W, one acute (W1) and one chronically active lesion (W2) were studied. From the second patient, R, 16 chronic lesions were studied. These lesions had inflammatory cells present, but the inflammatory cells were not participating in any form of ongoing demyelination.

The detailed methodology of cDNA microarray analysis has been described in detail elsewhere (41) Arrays for this study contained 2889 human cDNAs that were primarily derived from I.M.A.G.E. consortium cDNA libraries (42). A list of genes present on the arrays can be found at http://intra.ninds.nih.gov/Biddison/cDNA_microarray.asp. [33P]dCTP-labeled cDNAs were produced by reverse transcriptase from RNAs obtained from individual MS lesions, pooled normal white matter, experimental allergic encephalomyelitis, and normal mouse brains, and hybridized to the cDNA microarrays. Hybridizations of RNA obtained from MS lesions and experimental allergic encephalomyelitis brains were performed in two independent experiments, except for lesions R10, R11, and R16, in which enough RNA was obtained for only one hybridization. Quantitation of radioactivity bound to the arrays was performed on a Molecular Dynamics STORM PhosphorImager (Molecular Dynamics, Sunnyvale, CA) at 50 μm resolution. All data were analyzed from the PhosphorImager images using Pscan (Ref. 43, see also http://abs.cit.nih.gov/pscan). Pscan calculates spot intensities and compares spot intensities between samples, giving a ratio of gene expression between comparative samples. Using Pscan, spot intensities between arrays were automatically normalized to the background level.

FIGURE 1. Proliferative response of TCC GP5F11 (A) and TL3A6 (B) to the 200 mixtures of a decapeptide PS-SCL in which each position has one defined amino acid (20 for each of the 10 positions; the single letter amino acid code is used). Proliferation is shown as cpm induced by each mixture of the PS-SCL (mean and SD values of duplicate wells). *Proliferation in the absence of peptide mixtures. TCC GP5F11 is specific for an influenza virus HA-derived peptide, Flu-HA<sub>Δ308-317</sub>; TCC TL3A6 is specific for a MBP-derived peptide. The corresponding sequences of HA<sub>Δ308-317</sub> = YVKQNTLKLA, and MBP<sub>Δ90-99</sub> = FFKNIIVTPRT, are indicated by diamonds at the top of each panel. Proliferation in the absence of Ag was 124 + 42 cpm (A) and 1453 + 493 cpm (B).
median of all spot intensities on each individual array. Ratios of gene expression that were greater than 2-fold were considered significant based on a 99% confidence interval (44).

Results
Data obtained with combinatorial peptide libraries suggest different levels of TCR degeneracy for different CD4+ TCC

In this study, we sought to develop an approach that would combine the information generated from the screening of a decapeptide PS-SCL with all protein sequences in public databases. This strategy should allow the identification of the entire spectrum of stimulatory peptide ligands for a given TCC and the ranking of naturally occurring peptides with regard to predicted stimulation. The ultimate goal is to develop a methodology for identifying biologically relevant peptides for TCC of unknown specificity that have been isolated, e.g., from a tissue.

Three CD4+ TCC were tested in proliferative assays with the 200 mixtures of the decapeptide PS-SCL. Two TCC had known specificity, one specific for influenza HA (Flu-HA) (306–318) (TCC GP5F11), and one for MBP83–99 (TCC TL3A6). We also studied one clone of unknown specificity that recognizes B. burgdorferi, the causative organism of Lyme disease (TCC CSF-3).

Data obtained with combinatorial peptide libraries suggest different levels of TCR degeneracy for different CD4+ TCC. The stimulation profiles for TCC GP5F11 and TL3A6 are shown in Fig. 1, A and B, respectively. The profile for CSF-3 is shown previously (34). The profile of TL3A6 shows that more than one mixture in several positions of the PS-SCL generated a clear proliferative response. The amino acids of MBP89–98 are marked by diamonds (FFKNIVTPRT). Although the target amino acids correspond to the defined amino acid in the most stimulatory mixtures in most positions, this is not observed in certain positions, such as N in position 4 and P in position 8. In contrast, the profiles for GP5F11 and CSF-3 show a very different pattern with fewer, but more differential activity between stimulatory and not stimulatory mixtures.

Limitations of motif searches
Motif searches are widely used to search protein databases in a nonquantitative manner. However, this approach was not successful for identifying the known target peptides of the TL3A6 and

FIGURE 1. (continued)
GP5F11 clones. Motifs searches are generated from screening results of PS-SCL, and contained in each position are amino acids corresponding to mixtures with S-index greater than a specified threshold (see Materials and Methods for definition of S-index). Thresholds of 2 and 3 were used to generate the search motifs. The resulting motifs were then used to search the SwissProt and GenPept databases.

Tables I and II show the number of peptides that satisfied the motif searches, and indicates whether the target peptide was identified. The target peptide was not found with either of the motifs for TL3A6 in either database. The target peptide for GP5F11 was identified only when the search criterion was so permissive/lax that over 500 other peptides were also selected. Furthermore, the inability of motif searches to rank peptides renders it almost impossible to identify the most likely epitopes in a rational way and without synthesizing and testing very large numbers of individual peptides.

**Developing a score matrix-based approach for predicting T cell-stimulatory candidate peptides**

It is clear that a more systematic approach that employs all the data generated from the screening of PS-SCL needs to be developed for the search of databases. Our strategy is outlined in the flow diagram (Fig. 2).

We recently demonstrated that each amino acid within a peptide contributes to recognition almost independently and in an additive fashion, so that amino acid substitutions that abrogate recognition can be compensated for by highly stimulatory substitutions in a mixture derived from the PS-SCL screening is used, and the peptides, and then they are ranked accordingly. This strategy offers important advantages compared with motif searches: 1) all the information derived from the PS-SCL screening is used, and the selection based on a cutoff of activity is not required; 2) peptides are now ranked according to their predicted stimulatory score.

An example of a score matrix for one of the CD4+ TCC (GP5F11) is shown in Fig. 3A. The amino acids of Flu HA308–317 peptide are boxed. Note that the amino acids of the target peptide sequence L in position P7 and A in P10 are below an S-index value of 3, thus show that peptides that shared no amino acid in corresponding positions of their sequences could still be recognized by the same TCR (25). Also, the findings that the specificity information derived from PS-SCL libraries is similar to that obtained with individual peptide analogues and the fact that highly active peptides can be identified allow the development of a new search algorithm.

Our algorithm provides a predicted stimulatory score for the peptide of the same length as used in PS-SCL libraries. Based on the above assumptions, the peptide score is the sum of position-specific scores of the component amino acids. The scoring is accomplished by calculation of a matrix in which the columns represent positions, and the rows the 20 aa used in PS-SCL libraries. The scoring matrix entry for a particular amino acid in a specific position is based on the stimulation assay results for the mixture of PS-SCL corresponding to that amino acid defined in that position (Fig. 3A). The scoring matrix entry can either use the S-index or use the Z-index, which takes into account the experimental errors (see Materials and Methods).

The matrix is then used to search for predicted stimulatory peptides in the public protein databases. By moving a decamer scoring window across the known protein sequences in 1-aa increments (Fig. 3B), a stimulatory score is calculated for all published 10-mer peptides, and then they are ranked accordingly. This strategy offers important advantages compared with motif searches: 1) all the information derived from the PS-SCL screening is used, and the selection based on a cutoff of activity is not required; 2) peptides are now ranked according to their predicted stimulatory score.

**Table I. Database search performed on SwissProt and GenPept to identify agonist peptides for TCC GP5F11**

| S-Index | Search Supermotif<sup>a</sup> | SwissProt | GenPept |
|---------|-------------------------------|-----------|---------|
|         |                               | Target sequence | No. hits | Target sequence | No. hits, viral DB | No. hits, H. sapiens DB |
| >2      | [WYFRH]=[MLVADFYH]-K-[QVLYHVKFT]-[NHQM]-[TSNIGQVAVHM] | Yes | 513 | Yes | 560 | 177 |
|         |                               |             |         |               |               |               |
| >3      | [WYFR]=[MLVADF]-K-[QVLYHVK] | No | 82 | No | 23 | 34 |

<sup>a</sup> Amino acids corresponding to Flu HA(308–317) (YVKQNTLKLA) are shown in bold underlined characters. SwissProt contains 83,857 protein sequences (3-3-00); GenPept viral database: 90,174 proteins (20,198,794 decamer peptides); Homo sapiens database: 43,795 proteins (13,879,822 decamer peptides).

**Table II. Database search performed on SwissProt and GenPept to identify agonist peptides for TCC TL3A6**

| S-Index | Search Supermotif<sup>a</sup> | SwissProt | GenPept |
|---------|-------------------------------|-----------|---------|
|         |                               | Target sequence | No. hits | Target sequence | No. hits, H. sapiens DB | No. hits, viral DB | No. hits, bacterial DB |
| >2      | [WHYFARTLCQVKV]-[KIFSRYLMWTAVN]-[KDLGCGVYIQNH]-[LKKMVSAATDG]-[VMLIWTYR]-[VYMLPITSKWGEQNA]-[TSFVRQLKRGAPNY]-[KICTSPLQFMRWAHV]-[FKRVPYLI]-[TISVHWKMAMLR] | No | 260,085 | No | 104,229 | 183,876 | 289,887 |
| >3      | [WHYFARTLC]-[KIFSRY]-[KDLG]-[LKKMV]-[VMLI]-[VYMLPITSK]-[TSFVR]-[KICTSPL]-[FKRPY]-[TISVHW] | No | 797 | No | 285 | 502 | 776 |

<sup>a</sup> Amino acids corresponding to MBP(89–98) (FFKNIVTRPT) are shown in bold underlined characters. SwissProt contains 83,857 protein sequences (3-3-00); GenPept viral database: 90,174 proteins (20,198,794 decamer peptides); Homo sapiens database: 43,795 proteins (13,879,822 decamer peptides); bacterial database: 111,807 proteins (32,604,667 decamer peptides).
Obtain T cells from blood or tissue

Clone by limiting dilution

Test proliferative response of TCC to a decapetide PS-SCL (Fig. 1)

Design scoring matrix based on stimulatory potency of each mixture of the decapetide PS-SCL (Fig. 3A)

Use matrix to score all decapetides of proteins contained in public databases (Fig. 3B)

Synthesize selected peptides and verify response of TCC (Fig. 4)

FIGURE 2. Flow diagram of the strategy used to quantitatively analyze TCR recognition of Ags by clonotypic T cells. Experimental data collected by measuring functional T cell responses to PS-SCL are then analyzed by a scoring matrix approach. This allows the identification and ranking of the spectrum of antigenic ligands for TCC of known and unknown specificity.

For GP5F11, the rank among viral peptides is given; for TL3A6, we show the rank among human peptides. Consistent with previous observations with another autoreactive clone (45), MBF_{99-98} was far from optimal, i.e., it ranked only 202nd in the set of human peptides using the S-index matrix. In contrast, the target peptide Flu-HA_{309-318} ranked as the sixth highest scoring peptide for GP5F11 among viral proteins, and 24th when not only viral, but also human proteins were scored. This also suggests that molecular mimics that are potentially more stimulatory than the native foreign peptide can be identified.

We assessed the predictive power of the algorithm by synthesizing peptides tested for stimulation of the three clones (76 peptides for GP5F11, 144 peptides for TL3A6, and 88 peptides for CSF-3). For the two TCC of known specificity, TL3A6 and GP5F11, the peptide was considered stimulatory if its EC_{50} (concentration that yields half-maximal stimulatory activity) was equal to or <10 times that of the target peptide (MBP_{99-98} and Influenza virus HA_{308-317}, respectively). For CSF-3, the TCC of unknown specificity, the peptide was considered stimulatory if it activated the TCC with a Z-index >47.5 at any concentration between 0.001 and 100 μg/ml.

Table IV shows the relationship between stimulatory potential predicted by the scoring matrices and actual measurement of TCC stimulation. Thresholds for matrix score prediction were based on relative operating characteristic analysis (46) to balance sensitivity and specificity. For clone CSF-3, for example, of the 62 peptides predicted to be stimulatory (have scores above the threshold of 47.5), 58 did stimulate the TCC (a positive predictive value of 58/62, or 93.5%). Of the 26 peptides predicted to be nonstimulatory, only 5 stimulated the TCC (negative predictive value: 21/26, 80.8%). The sensitivity for predictions with this clone was 92%; that is, of the 63 peptides that actually stimulated the TCC, 58 were correctly predicted. The specificity was 84%; that is, of the 25 peptides that did not stimulate the TCC, 21 were correctly predicted. Although the sets of synthesized peptides are small compared with the number of peptides that would be predicted to be stimulatory, Table IV documents the excellent sensitivity, specificity, and negative predictive values for the three TCC.

Table V shows the information on the 10 highest scoring peptides derived from B. burgdorferi database analysis for TCC CSF-3 with the half-maximal stimulatory value that was determined by dose titration, proliferative experiments. Examples of the stimulatory activity of peptides predicted to activate TCC GP5F11 are shown in Fig. 4. Note that a predicted stimulatory peptide with optimal amino acids in each position (WMKQNIGRFL) and a higher score than the target peptide is in fact two orders of magnitude more potent than the target sequence. One of the known peptides with a score of 132.40 ranks much lower than the putative optimal amino acids in each position (WMKQNIGRFL) and a higher score than the target peptide is in fact two orders of magnitude more potent than the target sequence. One of the shown peptides with a score of 132.40 ranks much lower than the putative stimulatory threshold for TCC GP5F11, and consequently it did not stimulate the clone. However, even a few high scoring peptides (data not shown) are not stimulatory from reasons that are currently under further investigation.

Combining scoring matrix predictions of TCC stimulation with cDNA microarrays to identify biologically relevant candidate peptide mimics

The novel strategy described in this work allows us to find peptides from every known source that have stimulatory activity for the clone that was tested with PS-SCL. This leads to the problem of how one identifies from this wealth of data which peptides may be biologically relevant. In cases in which the target Ag for the clone is not known or molecular mimics with potential relevance for an organ-specific disease are of interest, several strategies may be used.
One approach to identify proteins involved in autoimmune diseases is to examine the expression of genes that are overexpressed in the target organ using cDNA microarray technology (41). We examined gene expression in 18 lesions from two MS patients and compared them with levels of gene expression in pooled normal white matter from three individuals with cDNA microarrays containing 2889 human genes. One of the genes that was overexpressed (>2-fold) in 17 of the 18 MS lesions examined was titin (Fig. 5A), a giant muscle protein (47). When we asked which genes that are overexpressed in MS plaques are also identified as candidate epitopes/molecular mimics for CD4+ TCC that were tested with the PS-SCL (Fig. 5B), we identified peptides derived from the same interesting candidate, titin, among the highest scoring peptides for both a CD4+ TCC recognizing the immunodominant MBP peptide (83–99) in the context of the MS-associated DR allele DRB5*0101, but also for the *B. burgdorferi*-specific TCC CSF-3 (Fig. 5C). Titin, a giant muscle protein (47), is surprisingly overexpressed in MS brain tissue, and the identification of titin-derived peptides as candidate molecular mimics for two TCC that are potentially...
pathogenic in two different CNS inflammatory/autoimmune disorders, i.e., MS and chronic CNS Lyme disease, offers unique opportunities to study the involvement of such candidate Ags in the pathogenesis of these diseases.

Discussion

The experiments presented in this work have been conducted to better understand, measure, and predict both specific and degenerate interactions between clonotypic TCRs and MHC peptide ligands. For this purpose, an approach was devised that would allow us to 1) describe in a quantitative way the complex interactions of the trimolecular Ag recognition complex, and 2) identify the spectrum of stimulatory ligands for individual TCC with high predictive accuracy. We used combinatorial peptide libraries and biometric strategies in conjunction with large scale database searches to achieve this goal and could show for the first time that T cell recognition can be predicted in quantitative terms. This study builds on and expands previous investigations on the flexibility and degeneracy of TCR recognition of Ag. A role for degenerate T cell recognition has been postulated for such diverse immunological phenomena as thymic selection (48), peripheral T cell survival (49), protection from infectious diseases, and induction of autoimmunity (49, 50). It was previously shown that peptide combinatorial libraries in the positional scanning format can be used to define the spectrum of agonist ligands for clonotypic TCR (26, 49). In recent studies, we showed that functional responses elicited in CD4+ TCC by PS-SCL could be used to build motifs for database searches and thus identify a spectrum of ligands of different potency for clonotypic TCR (45, 46). In the present study, we confirmed that functional T cell responses can be elicited by PS-SCL from certain CD4+ TCC specific for both foreign (Fig. 1A) (34) and self (Fig. 1B) Ags. We then used a matrix-based methodology for the analysis of the experimental data generated with the PS-SCL (Fig. 2). This methodology is based on a model of independent and additive contribution of each amino acid in the peptide sequence to the interactions with both the TCR and the MHC molecule (25). Although numeric matrices (8) and other mathematical approaches based on independent amino acid contribution to antigenicity have been previously used to describe the interaction of antigenic peptides with specific MHC molecules (17, 18), the present study fills the important gap of applying a quantitative, matrix-based model to the interaction of an MHC peptide ligand (keeping the MHC molecule constant) with a specific, clonotypic TCR using the data generated from PS-SCL. The biometric analysis described in this work systematically compares the information derived from a PS-SCL composed of trillions of decapeptides with all the decapeptides (13, 879, 822 for a H. sapiens database, and 20, 198, 794 for a viral database) contained in a public protein database to rank and predict the most stimulatory peptides for a given TCC. The predictions based on this methodology are so accurate (Tables III and IV, Fig. 4) (34) that they actually lend strong support to an additive, combinatorial model of peptide antigenicity. Available TCR crystal structures indeed suggest that peptides may modulate the preexisting affinity between MHC and TCR that is based on a large contact surface between these two components of the trimolecular complex (51, 52). It should be noted that this model does not contradict, but indeed extends and develops the concept of primary and secondary TCR contacts (23, 53). In fact, although complex substitutions of amino acids along the entire sequence of the peptide can lead to molecular mimicry in the absence of any sequence homology (25), the relative weight of different amino acids in each position of the peptides sequence is apparent from the experimental data (Fig. 1A and B).

An important application of the above described model is that one can identify peptide ligands for a specific TCR by searching public database not only with MHC and TCR anchor motifs (54) or motifs obtained from PS-SCL data (34, 45, 49), but also using the scoring matrix derived from the screening of a PS-SCL composed of trillions of peptides (Fig. 3A and B). We also illustrate the limitations of using motifs derived from PS-SCL screening to identify TCR agonist peptides. Such a strategy does not fully use the information generated by screening specific TCR with PS-SCL. Therefore, the native ligand may not be found if the motif is

Table III. Database search performed on GenPept with a sum of S-index score matrix

| TCC   | Target Sequence | Rank in Database |
|-------|-----------------|------------------|
| GP5F11 | Yes             | 6a               |
| TL3A6  | Yes             | 202b             |

a A total of 90,174 proteins scored in viral database (20,198,794 decamer peptides).
b A total of 43,795 proteins scored in H. sapiens database (13,879,822 decamer peptides).

Table IV. Indices of the predictive power of the scoring matrix approach for the definition of the stimulatory potency of antigenic peptides

| TCC CSF-3 | TCC GP5F11 | TCC TL3A6 |
|-----------|------------|-----------|
| Matrix score >47.5 | Matrix score >220 | Matrix score >45.2 |
| Matrix score <47.5 | Matrix score <220 | Matrix score <45.2 |
| Total       | Total       | Total       |

| Experimental measurement | Stimulatory | Nonstimulatory | Total |
|--------------------------|-------------|---------------|-------|
| Sensitivity              | 58/63 (92)  | 4/21 (25)     | 62/88 (88) |
| Specificity              | 21/25 (84)  | 32/34 (94.1)  | 53/59 (87.1) |
| Positive predictive value| 58/62 (93.5)| 38/40 (90.5)  | 96/102 (93.7) |
| Negative predictive value| 21/26 (80.8)| 32/36 (88.9)  | 53/62 (85.3) |
| Overall accuracy         | 79/88 (89.8)| 70/76 (92.1)  | 118/144 (81.9) |

a Fraction of all stimulatory peptides that is correctly identified.
b Fraction of all nonstimulatory peptides that is correctly identified.
c Probability that a peptide predicted to stimulate actually does so.
d Probability that a peptide predicted to be nonstimulatory actually does not activate the TCC.
e Fraction of all predictions that is correct.
This study adds a new and important contribution to the definition and prediction of T cell epitopes using synthetic combinatorial libraries (26, 27). It should be noted that many of the previous approaches to the identification of T cell epitopes were based on the prediction of which peptides would be good binders for specific MHC/HLA molecules (8, 16). Because only a fraction of the potential MHC-binding peptides is a T cell epitope for an individual TCR, these approaches provide information that is specific for particular MHC molecules, but cannot predict which fraction of the peptides that bind a restriction element is actually stimulatory for a TCR with its unique structural features. Conversely, TCR ligands are not always high affinity MHC binders (58). The approach presented in this study takes into account the whole trimolecular complex of T cell activation by reading out a functional T cell response. This requires a certain degree of MHC peptide binding as well as the interaction of the MHC peptide ligand with a specific TCR. When both are considered, the overall accuracy of T cell epitope predictions is far superior to previously adopted methods (Table IV), although further improvements are currently being pursued. This is particularly helpful when the protein(s) recognized by a TCC is/are not known (34). Indeed, less than a third of the peptides that were identified and found to be stimulatory by the PS-SCL and scoring matrix approach would have been predicted to be good MHC binders based on a recently published MHC-binding prediction algorithm (12) (data not shown).

Finally, we show that combining the above-described methodology with the use of cDNA microarrays to assess differential gene expression in pathological and normal tissue of two patients with MS led to an interesting candidate molecule (titin, to date only known as an

![FIGURE 4.](image)

**FIGURE 4.** Proliferative response of the TCC GPSF11 to representative agonist peptides identified by the peptide library strategy. The potency is highest for a theoretical peptide that is predicted to be a potent one because it has a high score. The native peptide (influenza virus HA308–317) and a double-substituted naturally occurring variant have intermediate potency. A low-scoring peptide derived from *H. sapiens* phosphatidylinositol-4-phosphate 5-kinase type III (PIPSKIII 246–255) and a theoretical peptide predicted to be nonstimulatory because it has a very low score are indeed nonstimulatory.

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**Table V. Information on the 10 highest scoring peptides derived from *B. burgdorferi* database analysis for TCC CSF-3**

| Score | Sequence | Protein ID No. | Protein Description | EC<sub>50</sub> µg/ml<sup>a</sup> |
|-------|----------|----------------|---------------------|--------------------------|
| 54.82 | NNIYKALIS | AE001155 | Hypothetical protein (section 41 of 70) of the complete genome | 1 |
| 54.14 | SNIKSLSF  | AE001174 | Hypothetical protein (section 60 of 70) of the complete genome | 0.1–1 |
| 53.73 | SNIKKTEDE | AE001169 | Similar to SP.P07017 (section 55 of 70) of the complete genome | 1 |
| 53.70 | FNIIYKRVVDN | AE001145 | Hypothetical protein (section 31 of 70) of the complete genome | 1 |
| 53.68 | NNIIDKVEYN | AE001135 | (section 21 of 70) of the complete genome; similar to GB:Z32522 | 1–10 |
| 53.09 | FFIRKRLIII | AE000785 | Hypothetical protein of plasmid Ip25 | 1 |
| 52.82 | RNIFKKTVEN | AE001130 | Similar to GB:L10328 (section 16 of 70) of the complete genome | >100 |
| 52.69 | SNIKSLILV  | AE001146 | Similar to PID:1652132 (section 32 of 70) of the complete genome | 1 |
| 52.63 | YNIIVSSLLL | AE001161 | Hypothetical protein (section 47 of 70) of the complete genome | 1–10 |
| 52.57 | DNIFKEKETLI | AE001165 | Similar to GB:L42023 (section 51 of 70) of the complete genome | 1 |

<sup>a</sup> Peptide concentration inducing half-maximal proliferation.
important component of skeletal muscle (47) that is overexpressed in MS plaques and is recognized by a B. burgdorferi-specific TCC (Fig. 5). Preliminary pathological studies by immunohistochemistry indicate the expression of an isoform of this molecule in the pathologic, as opposed to normal white matter tissue, but further work to define its role is clearly needed. Thus, the combination of two powerful methodologies can guide the discovery of candidate autoantigens that would otherwise not easily be identified by either approach.

In summary, we describe a methodology, PS-SCL-based biometrical analysis for ligand identification, which is consistent with a combinatorial model of TCR activation by antigenic peptides and allows the identification of T cell epitopes for both autoreactive
and foreign Ag-specific TCC with unprecedented efficacy. The same approach has also been successfully used for the prediction and identification of Ags by CD8+ TCC (Ref. 59 and R. Martin, B. Gran, M. Nagai, E. Borras, S. Jacobson, W. E. Bidddison, R. Houghten, H. F. McFarland, and C. Pinilla, unpublished results). For the first time, recognition of Ags by clones of unknown specificity can be decypted. This is an important advance in the study of autoimmune disease, in which one tries to suppress specific immune responses, as well as for infectious and neoplastic diseases, in which a stimulation of specific responses by vaccines is pursued. Furthermore, it is important to note that this approach can be used to identify ligands within proteins in public database for any molecular interaction that has been or can be studied with PS-SCLs composed of t-ami acids.

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