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## **Celiac disease : from basic insight to therapy development**

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## T-cell recognition of HLA-DQ2-bound gluten peptides can be influenced by an N-terminal proline at p-1

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# T cell recognition of HLA-DQ2-bound gluten peptides can be influenced by an N-terminal proline at p-1

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## **ABSTRACT**

Recent research has implicated a large number of gluten-derived peptides in the pathogenesis of celiac disease, a preponderously HLA-DQ2-associated disorder. Current evidence indicates that the core of some of those peptides is 10 amino acids long while HLA-class II normally accommodates 9 amino acids in the binding groove. We have now investigated this in detail using gluten specific T cell clones, HLA-DQ2-specific peptide binding assays and molecular modeling.

T cell recognition of both a  $\gamma$ -gliadin peptide and a low molecular weight glutenin peptide was found to be strictly dependent on a 10 amino acids long peptide. Subsequent peptide binding studies indicated that the glutenin peptide bound in a conventional p1-p9 register with an additional proline at p-1. Testing of substitution analogs demonstrated that the nature of the amino acid at p-1 strongly influenced T cell recognition of the peptide. Furthermore, molecular modeling confirmed that the glutenin peptide binds in a p1-p9 register and that the proline at p-1 points upwards towards the T cell receptor. Database searches indicate that a large number of potential T cell stimulatory gluten peptides with an additional proline at relative position p-1 exist suggesting that the recognition of other gluten peptides may depend on this proline as well. This knowledge may be of importance for the identification of additional T cell stimulatory gluten peptides and the design of a peptide-based tolerance-inducing therapy.

## INTRODUCTION

Celiac disease (CD) is a multifactorial inflammatory disorder caused by an uncontrolled T-cell response directed against wheat gluten and analogous grain storage proteins. The HLA-class II molecule HLA-DQ2 ( $\alpha 1^*0501$ ,  $\beta 1^*0201$ ) is the most important susceptibility locus for CD as roughly 95% of all CD patients are DQ2+ [1]. The role of HLA-DQ2 in presenting gluten-derived peptides to CD4+ effector T cells is well established. HLA-DQ2 selectively binds peptides with large hydrophobic residues at positions p1 and especially p9. At positions p4 and p7 negatively charged anchors are preferred, and at p6 a proline residue or a negative charge [2]. While native gluten hardly contains any negatively charged amino acids, these can be introduced by the enzyme tissue transglutaminase (tTG), that selectively deamidates glutamine residues in gluten, resulting in peptides that bind to HLA-DQ2 with high affinity [3,4].

Peptide binding to MHC class II molecules is accomplished by the accommodation of side chains of amino acids in the bound peptide in the respective pockets of the peptide-binding groove of MHC-molecules. In addition, a network of hydrogen bonds stabilizes the binding of the peptide in the groove over the entire contact length [5,6]. All known crystal structures of class II MHC molecules show that the peptide binding core consists of 9 amino acid residues (the so called p1/p9 register).

A large number of HLA-DQ2-restricted, T-cell stimulatory gluten epitopes that are implicated in the pathogenesis of celiac disease have been identified [7–10]. However, relatively little is known about their particular MHC binding characteristics. In a previous study we have reported algorithms that predicted T cell stimulatory gluten peptides [11]. These algorithms were based on requirements for binding to HLA-DQ2 and the specificity of the enzyme tissue transglutaminase. Two types of algorithms were used, one predicting 9-mer peptides, the other predicting 10-mer peptides. Strikingly, T cell responses were only observed against a series of gluten peptides, which were found with the algorithm searching for 10-mer peptides:  $X_1X_2X_3Q_4X_5P_6Q_7X_8P_9(YFWIL)_{10}$ . Moreover, in a recent study the binding of a 10-mer peptide to MHC-class II molecules was demonstrated and suggested to be the consequence of a bulge in the bound peptide [12]. Gluten molecules are very proline-rich and proline is known to introduce a bend in the polypeptide backbone of proteins and peptides, which might facilitate the binding of 10-mer gluten peptides to HLA-DQ2. Since the described predictive algorithm identified some 40 potential T cell stimulatory peptides in the gluten database this can indicate that such 10-mers may be an important part of the repertoire of T cell stimulatory gluten peptides. We have therefore investigated the requirements for binding of these putative 10-mer binding peptides to HLA-DQ2 in detail.

## **MATERIALS AND METHODS**

### **T-cell lines and clones**

The gluten specific T-cell clones were generated from small intestinal biopsies of celiac disease patients and have been described previously [10]. Culture flasks and other disposables were from Greiner (Frickenhausen, Germany).

### **Peptides**

Peptides were prepared using standard Fmoc chemistry as already described [10].

### **Tissue transglutaminase treatment**

tTG treatment was performed by incubating the peptides (500 µg/ml) with guinea pig tTG (100 µg/ml; Sigma) in buffer (50 mM triethylamine-acetate, 2 mM CaCl<sub>2</sub>, pH 6.5) for 4 hours at 37°C.

### **T-cell proliferation assay**

Proliferation assays were performed in triplicate in 150 µl RPMI-1640 (Gibco) supplemented with 10% human serum in 96-well flat-bottom plates (Falcon) using 10<sup>4</sup> gluten specific T cells stimulated with 10<sup>5</sup> irradiated HLA-DQ2-matched allogeneic PBMCs (3000 RAD) in the presence or absence of antigen peptides (10 µg/ml). After 48 hours at 37°C, cultures were pulsed with 0.5 µCi of <sup>3</sup>H-thymidine and harvested 18 hours later.

### **Peptide binding assay**

96-well FluoroNunc™ plates were coated with the HLA-DQ-specific monoclonal antibody SPV-L3, 2 µg/well in 100 µL of carbonate buffer (50 mM Na<sub>2</sub>CO<sub>3</sub>, 50 mM NaHCO<sub>3</sub>, pH 9.6) for 2 h at 37°C, subsequently blocked for 1 h at 37°C with 0.2% solution of gelatin in PBS. HLA-DR3/DQ2 positive EBV-transformed B cells were grown and a sample was checked for proper HLA-DQ2 expression by FACS-analysis using the SPV-L3 monoclonal antibody. Subsequently, the remainder of the cells were lysed in 20 mM Tris-HCl pH 7.5, 5 mM MgCl<sub>2</sub>, 1% NP-40 and protease inhibitor mix (Complete™, Roche), 4°C, 4 x 10<sup>6</sup> cells per 1ml. Cell debris was removed by centrifugation (4°C, 2000g, 15 min). Such prepared lysates were mixed with an equal volume of ice-cold 1% solution of BSA in PBS and pipetted into the SPV-L3 coated plates in 100 µL aliquots. After an overnight incubation at 4°C the plates were washed and 50 µL of binding buffer (0.1% NP-40, 0.1% Tween, 33.6 mM citric acid, 72 mM Na<sub>2</sub>HPO<sub>4</sub>, pH 5.5 and Complete™ protease inhibitor mix) was added to each well. A titration range of peptides to be tested (concentration range 600 - 1.0 µM) was prepared in 10% DMSO containing a fixed amount of the biotin-labeled indicator peptides Glt-156 or MHCIIα (46-63) at a concentration of 2.5 µM

and 1.2  $\mu\text{M}$ , respectively. Subsequently, 50  $\mu\text{L}$  of the samples was applied to the SPV-L3/HLA-DQ2 coated plates. Following a 48 h incubation at 37°C each well was washed extensively. Subsequently, 100  $\mu\text{L}$  of 1000 times diluted streptavidin-europium in assay buffer (both Wallace) was added and incubated for 45 minutes at RT. After extensive washing, 150  $\mu\text{L}$ /well of enhancement solution (Wallace) was applied and the plates were read in a time resolved fluorimeter (1234, Wallace) 15-30 minutes thereafter.  $\text{IC}_{50}$  values were calculated based on the observed competition between the test peptides and biotin-labeled indicator peptides and indicate the concentration of the tested peptide required for half maximal inhibition of the binding of the indicator peptide. Each  $\text{IC}_{50}$  value was determined in 3 independent experiments and the average is presented.

### Homology modeling

Homology modeling of the complex between HLA-DQ2 and the Glt-156 peptides was performed essentially as previously described [13]. The recently determined crystal structure of DQ2 was used as the base molecule [14]. The program Quanta (Accelrys, San Diego, CA, USA) was used to obtain a complete structure by providing the missing amino acid residues from the crystal structure (e.g.  $\beta 105-112$  as well as missing atoms from certain residues. The deamidated  $\beta$ -gliadin antigenic peptide in this structure (PFPQPELPY) with its 4 proline residues adopts a conformation in which the p9Tyr residue lies outside the p9 pocket in a niche formed by  $\beta 57\text{Ala}$ ,  $\beta 60\text{Tyr}$ , and  $\beta 61\text{Trp}$ . We reasoned that this mode of anchoring the p9 residue could not be correct for all peptides that bound to DQ2, especially those of high affinity e.g. the MHCII $\alpha$  (46-63) peptide with an affinity of 163 nM [15]. We therefore aligned the crystal structures of HLA-DQ2 and HLA-DQ8 on the polypeptide backbone atom positions of their  $\alpha$ - and  $\beta$ -chains and used the coordinates of the insulin peptide in the HLA-DQ8 structure as the proper ones for the Glt-156 peptide. The p9 pocket of DQ2, the most spacious p9 pocket of any MHC II molecule known to date, is formed by  $\alpha 68\text{His}$ ,  $\alpha 69\text{Asn}$ ,  $\alpha 72\text{Ser}$ ,  $\alpha 73\text{Leu}$ ,  $\alpha 76\text{Arg}$ ,  $\beta 9\text{Tyr}$ ,  $\beta 30\text{Ser}$ ,  $\beta 37\text{Ile}$ , and  $\beta 57\text{Ala}$  [15]. We reasoned that for a peptide with intermediate to high affinity for HLA-DQ2, as the Glt-156 peptide, the p9 pocket had to be occupied by the Phe residue. Energy minimizations proceeded via 1000 steps of the steepest gradient method followed by 1000 steps of the conjugate gradient method of the program Discover (Accelrys, San Diego CA, USA) with no cross-terms. Graphical representations were performed via the WebLabViewer program (version 3.5) of Accelrys. The energy-minimized structure thus obtained contains the peptide in a polyproline II helical conformation as expected for all peptides bound to MHC II molecules [16].

## Database search

To screen for potential gluten T cell epitopes the PIR Non-Redundant Reference Protein Database (PIR-NREF) was used (<http://pir.georgetown.edu/pirwww/search/pirnref.shtml>). The *Triticum aestivum* sequence register was searched with help of the PIR-supplied pattern search tool.

## RESULTS AND DISCUSSION

### Minimal epitopes required for T cell stimulation

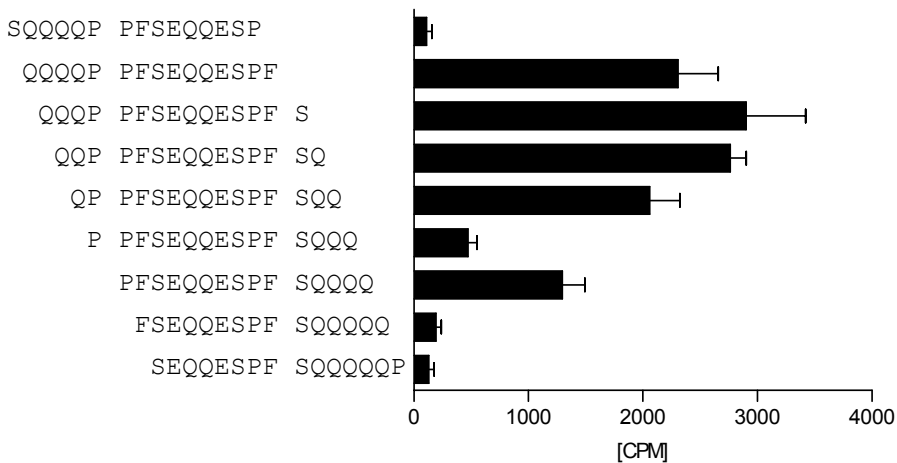
To determine the minimal peptide size required for T cell recognition we synthesized a set of partially overlapping peptides corresponding to 8 T cell stimulatory gluten peptides. Subsequently, these peptides were treated with tissue transglutaminase, which introduces the negative charges required for HLA-DQ2-binding, and tested the T cell stimulatory properties of these peptides with appropriate gluten specific T cell clones.

For most of the tested T cell stimulatory gluten epitopes the minimal core sequences required for T cell recognition confirmed previous results [10,11,17] and were found to be 9 amino acids long (Table 1). In contrast, the minimal core of the Glt-156 peptide required for T cell stimulation was found to consist of 10 amino acid residues (Fig. 1). Similarly, a requirement for a 10-mer was also found for the Glia- $\gamma$ 2 epitope ([11] and Table 1).

Alignment of the Glt-156 and Glia- $\gamma$ 2 peptides allows two alternative binding registers (Table 2). In the first binding register, termed p1/p10, the two negative charges that are introduced as the result of the activity of the enzyme tissue transglutaminase are found at position p4 and p7, which favors binding of these peptides to HLA-DQ2. In the second binding register, termed p-1/p9, the presence of phenyl-

**Table1.** Minimal gluten T cell stimulatory epitopes. Underlined - glutamine residues that are deamidated by tissue transglutaminase.

Epitope name	Core sequence	Amino acid length
Glia- $\gamma$ 30 (222-236)	IIQP <u>Q</u> QPAQ	9-mer
Glu-5 (unknown)	QXP <u>Q</u> QPQQF	9-mer
Glia- $\gamma$ 1 (138-153)	PQSF <u>P</u> Q <u>Q</u> Q	9-mer
Glia- $\alpha$ 2 (62-75)	PQP <u>Q</u> LPYPQ	9-mer
Glia- $\alpha$ 9 (57-68)	PYP <u>Q</u> Q <u>Q</u> LPY	9-mer
Glt-156 (40-59)	PFS <u>Q</u> Q <u>Q</u> QSPF	10-mer?
Glia- $\gamma$ 2 (89-102)	PFP <u>Q</u> QP <u>Q</u> QPF	10-mer?



**Figure 1.** Minimal core sequence of Glt-156 epitope capable of stimulating T cell proliferative response.

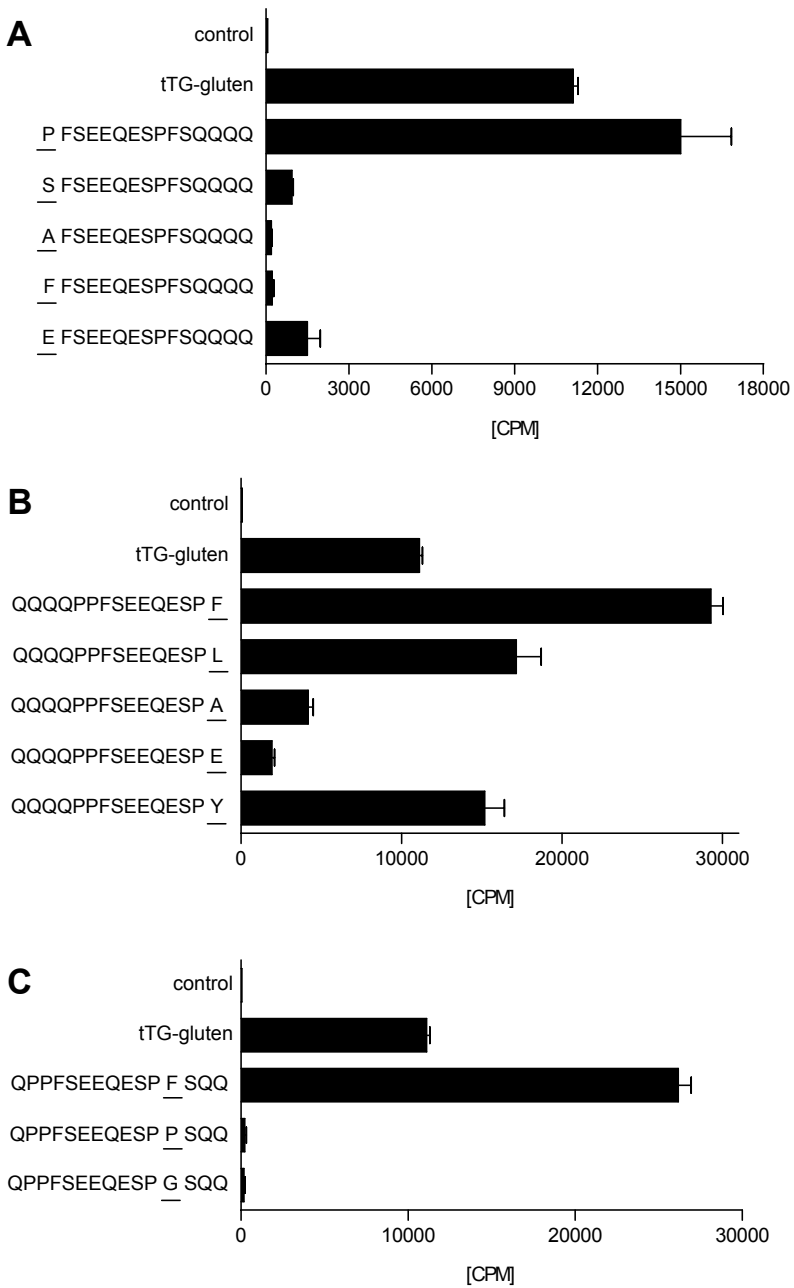
alanine at position p1 and p9 and a glutamate at p6 could facilitate binding to HLA-DQ2. In order to confirm the importance of the N-terminal proline in the Glt-156 peptide for the T cell recognition we tested the impact of amino acid substitutions at this position. Substitution of the proline with serine, alanine, phenylalanine and glutamic acid strongly reduced T cell recognition (Fig. 2A).

Subsequently, we determined the need for the phenylalanine at the C-terminus of the Glt-156 peptide. For this purpose homologs were synthesized, in which the phenylalanine was substituted with proline, glutamine, alanine, glutamic acid, leucine or tyrosine and the T cell stimulatory properties of these peptides were tested in two independent experiments (Fig. 2B, C). While the conservative tyrosine and semi-conservative leucine substitutions only moderately reduced the T cell stimulatory properties, these properties were strongly diminished or completely abolished by non-conservative proline, glycine, alanine and glutamic acid substitutions (Fig. 2B,C).

**Table 2.** Alignment of peptides sequences with HLA-DQ2 binding motif.

Position: Preferred:	P-1	P1 FWYILV	P2	P3	P4 DEVLI	P5	P6 PAE	P7 DE	P8	P9 FYLWI	P10
Glt 156		P	F	S	E	E	Q	E	S	P	F
Glia $\gamma$ -2		P	F	P	E	Q	P	E	Q	P	F
Glt 156	P	F	S	E	E	Q	E	S	P	F	
Glia $\gamma$ -2	P	F	P	E	Q	P	E	Q	P	F	





**Figure 2.** Influence of substitutions of N-terminal proline and C-terminal phenylalanine on proliferation of gluten specific T cell clone. tTG gluten - peptic/tryptic gliadin digest treated with tissue transglutaminase.

**Table 3.** Influence of p9 substitution on HLA-DQ2 binding.

Amino Acid Sequence	IC <sub>50</sub> [ $\mu$ M]
<u>Q P P F S E E Q E S P F S Q</u>	3.5
Q P P F S E E Q E S P <b>P</b> S Q	48.0
Q P P F S E E Q E S P <b>G</b> S Q	28.0

To exclude the possibility that the substitutions of the C-terminal phenylalanine affected the deamidation of glutamine at the putative p7 residue, and thereby abrogated the T-cell recognition, we checked the effect of the amino acid replacements on the deamidation pattern. As expected, the deamidation was compromised by the introduction of a proline since a proline located 3 amino acids C-terminal of a glutamine is known to inhibit the deamidation of this glutamine [11]. The replacement with a glycine, however, had no influence on the deamidation (not shown). As the T cell stimulation was compromised by both substitutions, this confirms the significance of the bulkiness of the amino acid on the C-terminus for the T cell recognition.

Together, these results indicate that a 10-mer peptide is required for T cell recognition of the Glt-156 peptide. Similar results were obtained for the Glia- $\gamma$ 2 peptide (not shown). We therefore investigated the possibility that the peptide binds in a p1/p10 register in more detail.

### Minimal epitopes required for HLA-DQ2 peptide binding

To distinguish between the two possible peptide-binding registers we have carried out peptide binding studies to HLA-DQ2. Since the Glia- $\gamma$ 2 is known to be a relatively poor HLA-DQ2 binder [18], these studies were carried out with the Glt-156 epitope.

First, we checked the HLA-DQ2 binding capacities of Glt-156 with C-terminal

**Table 4.** Influence of p1 replacement on HLA-DQ2 binding.

Amino Acid Sequence	IC <sub>50</sub> [ $\mu$ M]
<u>Q P P F S E E Q E S P F S Q Q</u>	3.0
Q P P <b>L</b> S E E Q E S P F S Q Q	3.8
Q P P <b>A</b> S E E Q E S P F S Q Q	10.0
Q P P <b>K</b> S E E Q E S P F S Q Q	50.0
Q P P <b>D</b> S E E Q E S P G S Q Q	100.0

**Table 5.** Minimal core sequences facilitating HLA-DQ2 binding.

Amino Acid Sequence	IC 50 [ $\mu$ M]
S Q Q Q Q P P F S E E Q E S P	49.0
Q Q Q Q P P F S E E Q E S P F	0.9
Q Q Q P P F S E E Q E S P F S	2.1
Q Q P P F S E E Q E S P F S Q	3.2
Q P P F S E E Q E S P F S Q Q	2.1
P P F S E E Q E S P F S Q Q Q	3.1
P F S E E Q E S P F S Q Q Q Q	6.4
F S E E Q E S P F S Q Q Q Q Q	14.7
S E E Q E S P F S Q Q Q Q Q P	>500

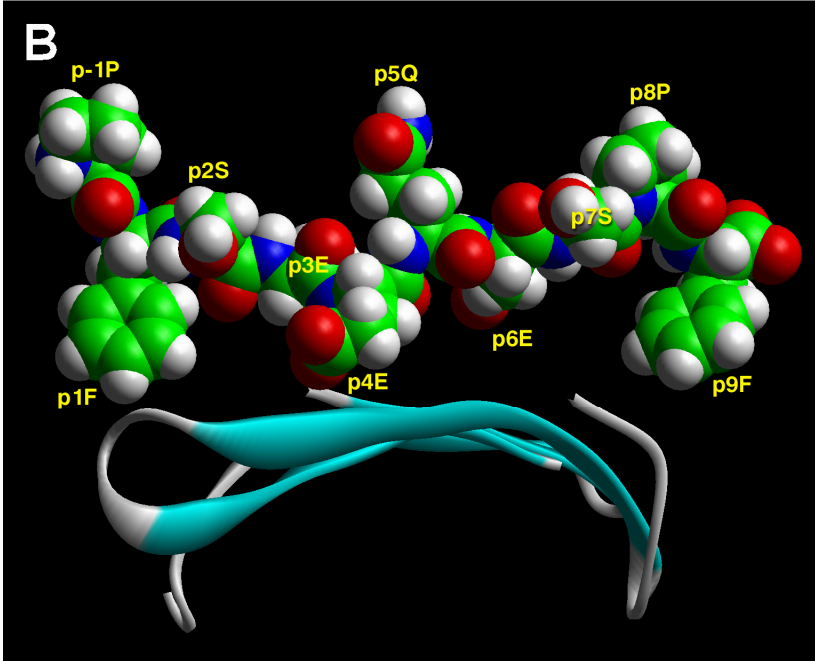
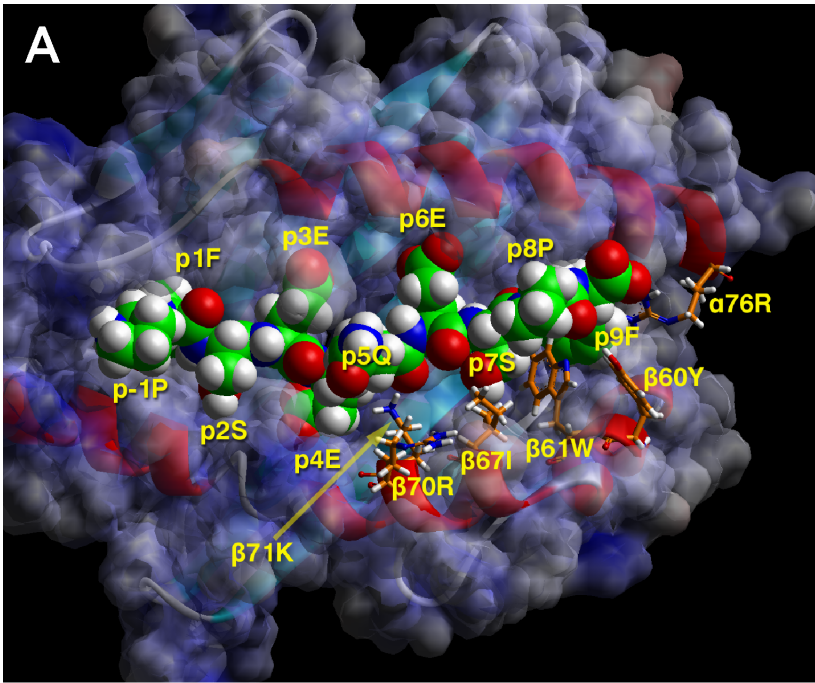
substitutions (F→P or F→G) in a cell free HLA-DQ2 peptide-binding assay. Both substitutions were found to result in significantly higher IC<sub>50</sub> values compared to the wild type peptide (Table 3), indicating a lower binding capacity as the result of the substitutions. We therefore conclude that the C-terminal phenylalanine functions as an anchor residue in the p9 pocket of HLA-DQ2.

To address the question which amino acids may serve as anchors at the N-terminus of the peptide in the p1 binding pocket we have substituted both the N-terminal proline and phenylalanine with other amino acids and determined the HLA-DQ2 binding capacities of these homologue peptides.

Substitutions of N-terminal proline had no effect on HLA-DQ2 binding capacity (data not shown), indicating that this proline is not strongly involved in interactions between the peptide and HLA-DQ2 molecule. Substituting the phenylalanine with other hydrophobic amino acids only slightly decreased binding. Replacements with charged residues, however, had a very pronounced negative effect (Table 4), indicating that the phenylalanine side chain is buried in the hydrophobic p1 pocket, docking the N-terminus of the peptide in the HLA molecule.

To further confirm that the N-terminal proline is not required for binding to HLA-DQ2 we have carried out a binding test using an overlapping set of Glt-156-based peptides and determined the minimal HLA-DQ2 binding core sequence (Table 5). Indeed, contrary to the minimal T cell stimulatory epitope, the minimal binding core was shown to consist of only 9 amino acids and did not include the N-terminal proline.

Finally, we modeled the Glt-156 and the Gli $\alpha$ - $\gamma$ 2 peptide using the recently solved HLA-DQ2 crystal structure [14]. In agreement with this crystal structure, optimal docking of the Glt-156 and Gli $\alpha$ - $\gamma$ 2 peptides into the HLA-DQ2 peptide-binding



**Figure 3.** Computer modeling of the Glt-156 peptide in the HLA-DQ2 molecule. The peptide in the groove is shown in space filling mode with the following color convention: carbon, green; nitrogen, blue; oxygen, red; hydrogen, white. Positive electrostatic surface potential is depicted as a blue surface, negative potential as a red surface, and intermediate values as grey surface. For the HLA-DQ2 protein,  $\alpha$ -helix is shown in red,  $\beta$ -pleated sheet in turquoise, random coil in grey. **A:** TCR view of the peptide in emerged in the HLA-DQ2 peptide-binding groove. p1 and p9 phenylalanine residues are buried in respective pockets, docking the sides of the peptide. p-1 proline is easily accessible to the TCR. **B:** Side view depicting the Glt-156 peptide (top) and non-immunostimulating peptide FSEEQESPFS at the level of the  $\beta$ -sheet floor in the exact orientation found in the groove. Note that residues p1F, p4E, p6E and p9F are buried in the respective pockets while the p-1Pro is pointing upwards. The major effect arising from the p-1 deletion is the absence of a residue in that space, and the positive charge at the amino-side of p1Phe, instead of p-1Pro. There are also slight movements of p1Phe and TCR-exposed residues, e.g. p3, and p4 that might influence TCR recognition, but such an effect would be secondary to the impact of the lack of the p-1Pro.

groove was found with the conventional p1/p9 register (Fig. 3 and not shown). As expected, the total calculated energy of the complex in the conventional p-1/p9 register is lower than in the p1/p10 register, or a register in which the N-terminal proline forms the p1 anchor and the C-terminal phenylalanine forms the p9 anchor with a presumed bulge between p7Glu and p8Ser (data not shown). The modeled structure of the complex shows that several features of the DQ2 molecule can exhibit profound influence on the binding of the Glt-156 peptide. Thus, we find the two phenylalanine residues inside pockets 1 and 9, the p4Glu deeply buried in the p4 pocket and p6Glu pointing parallel to the  $\beta$ -sheet floor. Moreover, the hydrogen bonding interactions of the constant MHC II residues and the Glt-156 peptide backbone are all in place, justifying the high binding affinity measured for this peptide. Interestingly, the proline at p-1 is pointing upwards which may explain its impact on T cell recognition (Fig. 3B). In addition, removal of the proline has a small impact of the position of the p1 anchor residue (Fig. 3B). Moreover, the presence of p-1 proline adds a potential hydrogen bonding interaction of the positively charged  $\alpha$ -amine group of the peptide with the carbonyl oxygen of HLA-DQ2  $\alpha$ -chain residue 53. This interaction may have a small impact on the IC<sub>50</sub> value (Table 5).

### Database search

Proline is a highly abundant amino acid in gluten constituting about 12% of wheat glutenins and 17% of wheat gliadins (in barley this is 15% and 23% respectively) [19]. We have already demonstrated in our previous paper that among the epitopes identified by 10-mer predicting algorithms only those that contained proline at the N-terminus (position p-1) were capable of evoking T-cell responses [11]. To check how many potential gluten epitopes contain proline at the p-1 position we designed

**Table 6.** Results of database search with algorithms predicting potential epitopes binding to HLA-DQ2 with Pro in p-1 position.

Algorithm	Glutenins	Gliadins	Number of hits	Distinct sequences
P F X X Q X Q X P [FYLWI]	37	26	168	14
X* F X X Q X Q X P [FYLWI]	14	5	22	9
P Y X X Q X Q X P [FYLWI]	13	16	29	3
X* Y X X Q X Q X P [FYLWI]	0	0	0	0

X – any amino acid

X\* - any amino acid except Pro

4 algorithms with glutamine at the anchor positions p4 and p6, phenylalanine or tyrosine at the anchor position p1, a bulky hydrophobic amino acid (F, Y, L, W or I) at the anchor position p9. The proline at the position p8 was meant to facilitate the deamidation of glutamine at the p6. Using these algorithms we screened a protein database and found 17 potential epitopes with proline at position p-1. These epitopes were repeated almost 200 times in more than 60 different gluten molecules (Table 6). The repertoire of identified sequences with other amino acids at p-1 was more limited, consisting of 9 potential epitopes repeated only 22 times in 19 gluten molecules. Thus, gluten contains many potential T-cell epitopes with N-terminal proline at p-1 position.

## CONCLUDING REMARKS

The results show that certain gluten epitopes, although binding to HLA-DQ2 in the canonic p1/p9 nonamer register, require a p-1 proline for optimal T cell recognition. The prominence of the peptide-flanking residues for the T cell receptor (TCR) interaction and consequently epitope immunogenicity has been already well established [20]. Arnold et al. demonstrated that recognition of peptide flanking residues is a common event [21]. In contrast, in the few available MHC II-peptide-TCR complex crystal structures the CDR3 regions of the TCR alpha and beta chains align over the p5 residue of the bound peptide [22–24]. This has led to the suggestion that the specificity for peptide recognition is dependent on TCR contact with the central p5 residue. In other cases, however, TCR recognition of bound peptide was found to be specific for p2/3 and p7/8 residues [25]. Thus, TCR recognition of HLA-class II bound peptides is not uniform. Here we show that TCR recognition of gluten peptides can be influenced by an N-terminal proline at p-1.

Since there is a high number of potential gluten epitopes with a N-terminal flanking proline, we suggest that this phenomenon should be taken into account while searching for new gluten epitopes or designing novel peptide-based tolerance-inducing therapies for celiac disease.

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