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Functional Reconstruction of Structurally Complex Epitopes using CLIPSTM Technology[#]

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Abstract: This review summarizes an illustrative set of data from our research on the reconstruction of structurally complex protein surfaces, i.e. those that fail to be mimicked properly by linear peptides. In the past 5 years, our newly developed CLIPSTM technology has proven an extremely valuable tool for 1) binding site mapping of therapeutically relevant mAbs, 2) generating hyperimmune sera *via* immunization with CLIPS peptides, and 3) the generation of monoclonal antibodies (mAbs) *via* the use of hybridoma technology. We currently have data available for more than 50 therapeutic targets. The examples described in this review illustrate the potential of this powerful new technology.

Key Words: Peptide, protein mimicry, epitope, conformational, discontinuous, monoclonal antibodies, vaccine.

INTRODUCTION

The functional mimicry of a folded protein surface using peptides is a scientifically challenging problem. It has been commonly accepted that “many proteins exert their biological activity through relatively small regions of their folded surfaces, so their actions could in principle be reproduced by much smaller “designer” molecules that retain these localized bioactive surfaces, but have improved pharmacodynamic and kinetic properties” [1]. Peptide-based protein mimics have enormous value for the development of therapeutic antibodies or synthetic vaccines, where an increasing number of protein targets is being identified that is difficult to address by using standard techniques (HIV, GPCR's, etc.). Antibody-based immunotherapies are nowadays regarded as the most promising therapeutic approaches for life-threatening diseases, especially when combined with classical therapies [2, 3].

Occasionally, short linear peptides can mimic the function of a whole protein, but this approach is largely limited to unstructured N- and C-termini of proteins (Fig. 1). In the early 1980s, Bittle and Dimarchi described the first example of (partial) protection of cattle for viral infection with foot-and-mouth-disease-virus (FMDV) *via* immunization with

linear peptides derived from the coat-protein of the virus [4, 5]. Soon after, Langeveld *et al.* obtained complete protection of dogs against canine parvovirus (CPV) infection *via* immunization with linear peptides derived from the VP2-protein [6, 7]. Other studies showed that immunization with a 37-mer peptide derived from the C-terminus of human choriongonadotrophin (hCG; CTP37), the pregnancy hormone, renders women temporarily infertile [8, 9]. Following this observation, anti-hCG vaccination has been studied as means of affordable birth control in underdeveloped countries [9], and recently also as a potential therapy for the treatment of hCG-overexpressing tumors [10, 11].

For mimicry of conformational and/or discontinuous binding sites (Fig. 1B) linear peptides generally fail, because they do not adopt the correct secondary and tertiary structure. For that purpose, more elaborate tools are needed to position correctly the important functionalities in space, which is an absolute prerequisite for activity [12-14].

THERMODYNAMICS OF PROTEIN FOLDING AND THE PURPOSE OF SYNTHETIC SCAFFOLDS

In order to understand better why small peptides (20-30 amino acids) are not stably folded, contrary to the proteins they were derived of, it is important to consider why proteins unfold. Proteins exist in a dynamic equilibrium between a folded (F) and an unfolded state (U) (Fig. 2A). The Gibbs free energy of folding is given by the following equation: $\Delta G_{F-U} = \Delta H_{F-U} - T\Delta S_{F-U}$. The folding process involves two opposing forces, i.e. the drive towards lower enthalpy ($\Delta H \downarrow$) and the drive towards maximum entropy ($\Delta S \uparrow$) [15]. The folded state is lowest in enthalpy. It is stabilized by

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[#]This review is partly based on a paper presented at the Vaccine Development Conference held at the University of Pittsburgh on 28-30 September 2008 (<http://www.biocore.com/lifesciences/events/vaccine2008/home/index.html>).

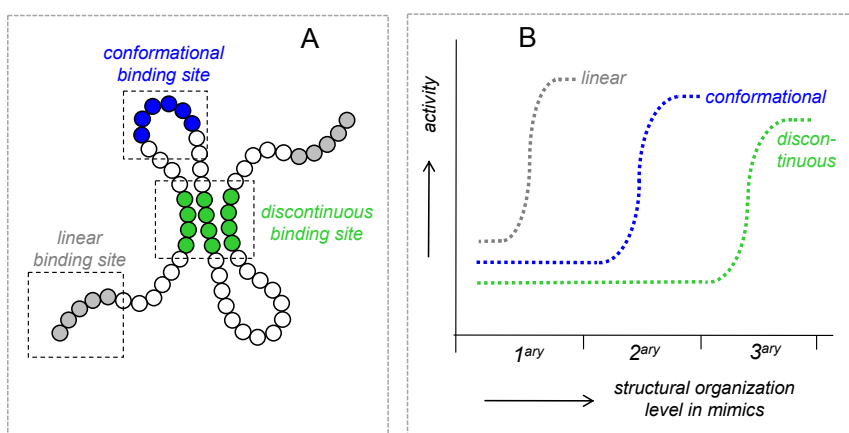


Fig. (1). Schematic representation of **A)** linear, conformational and discontinuous binding sites, and **B)** required level of structural organization in protein mimics for each of these.

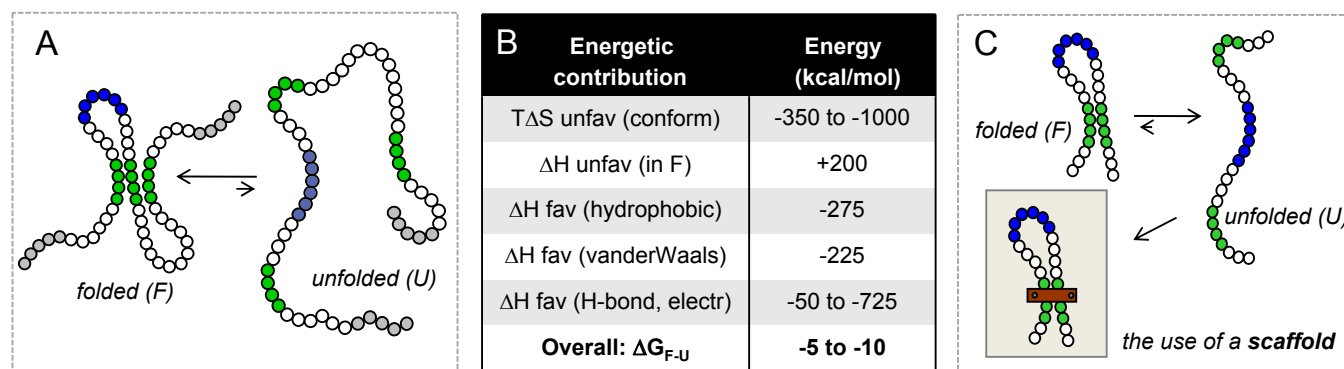


Fig. (2). Schematic illustration of **A)** dynamic equilibrium between folded and unfolded state of proteins, **B)** different enthalpic and entropic contribution to overall Gibbs free energy of folding, and **C)** unfolding of small peptides and stabilization of local protein structure by means of scaffold technology.

intramolecular interactions (hydrophobic, ionic, H-bond, etc.) between side and main chain elements. Hydrophobic parts stick together and form the inner core of the protein, while other groups remain at the outer, solvent-accessible surface. From an entropic point of view, the folded state is disfavored, since the protein likes to retain a maximal degree of conformational freedom. For structured proteins, the enthalpy term outweighs the entropy term (usually ~ 5 -10 kcal/mol) and so the protein is stable in its folded state. However, this balance can be tipped over by heating, which explains why proteins denature at elevated temperatures (increasing importance of $T\Delta S$, ΔG_{F-U} becomes positive).

The majority of small peptides (20-30 amino acids) derived from intact proteins lack a well-defined structure in solution. In fact, removal of the remainder of the protein backbone has taken away the majority of stabilizing, enthalpic, interactions that normally give structure to the protein. The peptide requires an energetic compensation for this, to prevent it from remaining unstructured. Synthetic scaffolds can usually solve this problem by affixing the loose ends of the peptide. A scaffold is able to strongly reduce the unfavourable entropy term for spontaneous unfolding of the flexible peptide by lowering the number of possible conformations (Fig. 2C). In this way, the scaffolded peptide may be able to adopt the same spatial structure as the corresponding sequence in the intact protein.

CLIPS™ TECHNOLOGY: MIMICRY OF CONFORMATIONAL AND DISCONTINUOUS EPITOPES

A variety of chemical methods is currently available for structural fixation of peptides into stable secondary structures, like α -helices [16-19], β -sheets [20], or β -hairpins [21-23], either *via* backbone or side chain functionalities (SS, Br-SH, $\text{CO}_2\text{H}/\text{NH}_2$, metathesis, CLICK chemistry) [19, 24, 25]. Some of these require special protection/deprotection protocols ($\text{CO}_2\text{H}/\text{NH}_2$) or the addition of a catalyst (SS, metathesis, CLICK), since otherwise the reaction rates are too slow (SS, CLICK). This may severely limit applicability to the purpose of binding site reconstruction.

We recently developed a novel and generic technology for structural fixation of linear peptides (Fig. 3) [26]. This technology, termed **CLIPS™** technology (Chemical **L**inkage of **P**eptides onto **S**caffolds, (<http://www.pepscan.com>), is applicable to peptides in solution as well as on microarrays used for instance for epitope mapping (PEPSCAN method) [26, 27]. Conversion of the peptides is extremely fast (5-10 min) and takes place at room temperature in 100% H_2O at pH 7.8-8.0, or, in case of water-insoluble peptides, in the presence of up to 80% of an organic solvent (ACN, DMF, DMSO). Furthermore, the reactions run smoothly without catalyst and with fully unprotected sidechain functionalities (NH_2 , COOH , OH , imidazole, etc.). Occasionally,

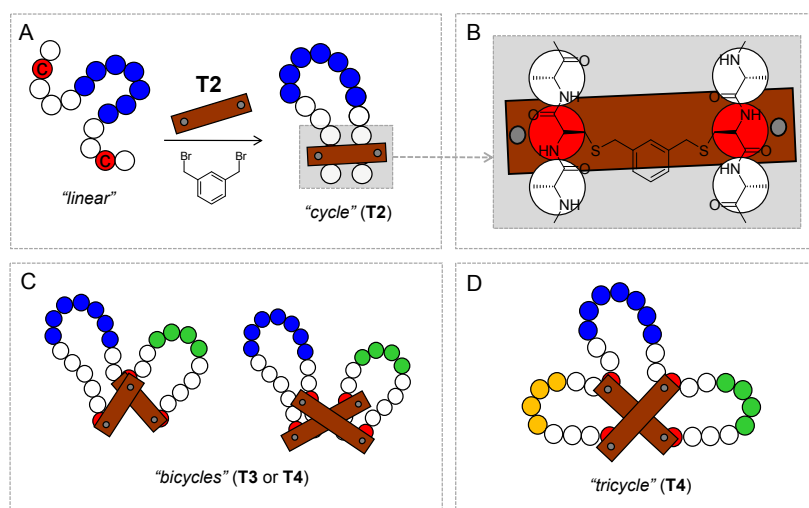


Fig. (3). A) Schematic representation of CLIPS-reaction; B) molecular structure of CLIPS-peptide; C and D) different CLIPS-based topologies (T3 and T4) for mimicry of discontinuous epitopes.

we encounter side reactions at the thioether functionality of methionine, but this is not a general phenomenon and always sequence-dependent.

CLIPS™ technology involves the chemical reaction between peptides containing (homo)cysteine residues and a synthetic scaffold ('T' for template) containing a benzyl bromide group (PhCH₂Br). Reaction of the T-scaffold is entirely selective for the nucleophilic thiol (SH) group of cysteine, and is not observed with amines (NH₂), carboxylic acids (COOH), phenols (PhOH), aliphatic alcohols (RCH₂OH), imidazoles, tryptophans, or thioethers (RCH₂SCH₃). When peptides containing two cysteines are reacted with a scaffold containing two PhCH₂Br groups (*o*-/*m*-/*p*-dibromoxylene, T₂; 2,6-bis(bromomethyl)pyridine, P₂), a cyclic product is formed rapidly (Fig. 3AB). Similarly, reaction of a scaffold containing three (1,3,5-tribromomesitylene, T₃) or four PhCH₂Br groups (1,2,4,5-tetrabromodurene, T₄) with peptides containing either three or four cysteines readily produces the corresponding 1:1 products, in which each cysteine has reacted with one of the PhCH₂Br groups, mainly in an intramolecular fashion (Fig. 3CD). We use CLIPS™ technology for two main purposes, i.e. *i*) to fix the conformation of linear peptides and so improve antibody binding (mainly T₂/P₂), and *ii*) bring together remote parts of discontinuous epitopes in a single, structurally defined entity (mainly T₃/T₄). CLIPS™ technology has found widespread application in 1) epitope mapping studies of therapeutic mAbs [26, 27], 2) eliciting antisera with neutralizing activity for the target protein from which they were derived, and 3) generation and selection of mAbs or scFvs using antibody (hybridoma, phage display library) technologies. Currently, our dataset include more than 50 therapeutically relevant targets, most of which are confidential and proprietary to third parties. Here we present three examples of experimental work that illustrate the kind of results that can be expected when using this technology.

EXAMPLE 1: THE CYS-KNOT PROTEIN FAMILY: FOLLICLE STIMULATING HORMONE (FSH)

Follicle stimulating hormone (FSH) is a hormone that plays a key role in fertility regulation in mammals [28].

Together with luteinizing hormone (LH), thyrotrophin hormone (TSH) and choriogonadotrophin (CG), it constitutes the homologous family of glycoprotein hormones, a subfamily of the cys-knot growth factor superfamily. All four hormones are heterodimers, consisting of a common α -subunit and a unique, hormone-specific β -subunit (Fig. 4A) [29,30]. *In vivo* neutralization of human-FSH has long been considered a viable option for contraceptive therapy in men [31]. For example, immunization with ovine-FSH has shown to be able to largely impair spermatogenesis and completely inhibit fertility [32]. However, FSH antisera have the potential to crossreact with LH and TSH, which comprise identical α -subunits as FSH. For this reason, peptide vaccines derived from immunization with the FSH- β subunit only have gained increasing interest. Antigenic mapping studies revealed that two of the nine antigenic sites on FSH are located on the β -chain [33]. Moreover, it was shown that residues 31-52 (β 2-loop) and 66-85 (β 3-loop; Fig. 4B) harbor epitopes involved in hFSH neutralization. Previous studies in our laboratory showed that sera generated with the tandem β 2-loop peptide (β 33-53)₂ possessed neutralizing activity. However, similar studies with β 3-loop derived peptides were so far not successful.

Epitope Mapping Studies for Two Neutralizing Anti-FSH mAbs Using CLIPS™ Technology

From a pool of 20 hFSH mAbs, we selected two mAbs (6602 and 5828) with the strongest neutralizing activity against hFSH in a cell-based proliferation assay [34]. Epitope mapping studies using overlapping libraries of 12- to 18-mer CLIPS peptides (1-12, 2-13, 3-14, etc.) showed that both mAbs bind to a highly conformational binding site (Fig. 5C), located mainly at the top of β 3-loop (core is R₆₂-L₇₃) [35]. It was found that for 12-mer peptides the binding of CLIPS variants was far superior to linear peptides (Fig. 5B), while for longer peptides (18-mers) the difference in binding levelled off. Apparently, the hydrophobic residues flanking the core region (₅₆LVY₅₈ and ₇₆YPV₇₈) can also serve as a scaffold for loop fixation, e.g. *via* mutual hydrophobic interactions.

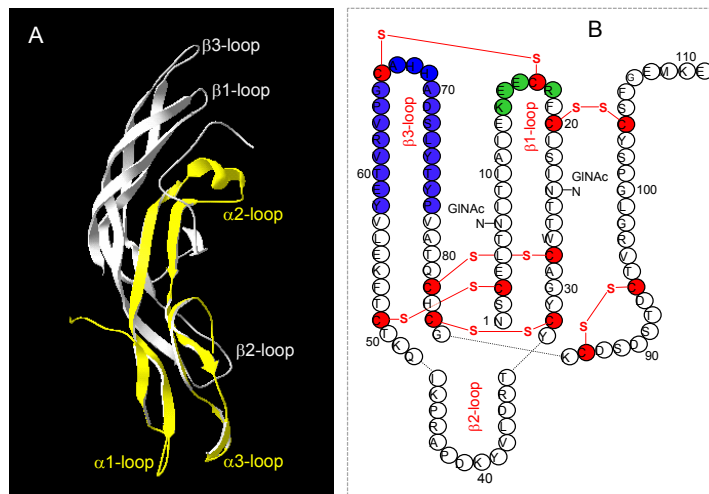


Fig. (4). **A)** Ribbon structure (1fl7.pdb) of the heterodimeric protein follicle stimulating hormone (FSH), consisting of an α - (in yellow) and β -subunit (in grey); **B)** primary amino acid sequence, loop structures and disulfide topologies in FSH- β .

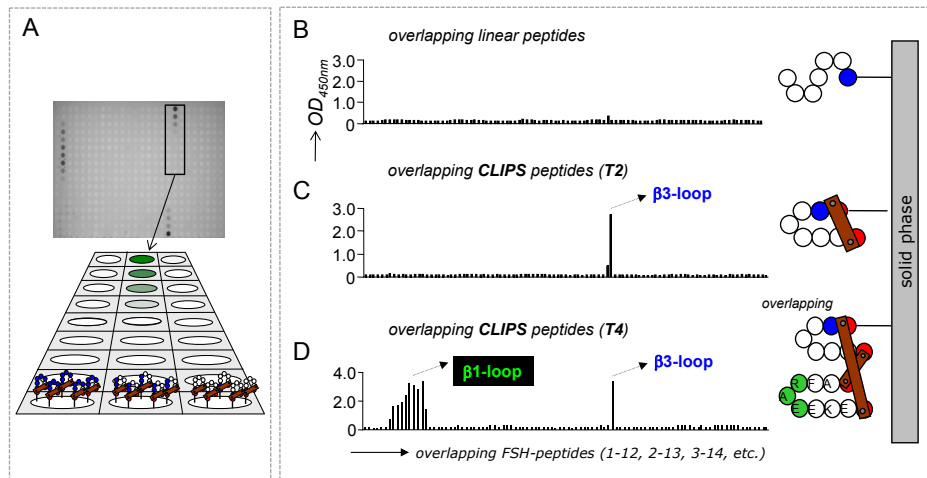


Fig. (5). **A)** “Real-life” picture and schematic figure of a microarray of overlapping CLIPS peptides. **B-D)** Epitope mapping data for anti-FSH mAb in three different formats using **B)** linear peptides, **C)** cyclic CLIPS peptides (**T2**), and **D)** bicyclic CLIPS peptides (**T4**; formed as mixture of *ortho*-, *meta*-, and *para*-substituted isomers). Optical density data ($OD_{450\text{nm}}$) relate to an ELISA-type binding assay.

In order to identify additional residues that are part of the binding site for mAb 5828 and 6602, we also screened these mAbs in the so-called “double-loopscan” format with **T4** as a scaffold (Fig. 5D; unpublished data). Therefore, the CLIPS-peptide construct $C_{T4}EKEEARFAC_{T4}-Br_2$, the sequence which was derived from the FSH $\beta 1$ -loop, was chemically linked to a microarray of overlapping FSH- β derived peptides *via* the two reactive CH_2Br groups. In this way the residues E_{16} and R_{18} were identified as part of the discontinuous binding site of mAb 5828 and 6602, since all peptides containing these residues showed clearly improved binding to both mAbs. This showed also that the binding site of both mAbs was both conformational and discontinuous in nature, covering residues on top of both the $\beta 3$ - and $\beta 1$ -loop region.

Immunization Studies with CLIPS Peptides Derived from Human FSH

Subsequently, we studied in detail the possibilities to use CLIPS peptides derived from the $\beta 3$ -loop of hFSH for generating antibodies that display *in vitro* hFSH neutralizing activity (Fig. 6) [35]. Initial experiments with CLIPS

peptides derived from both FSH and hCG $\beta 3$ -loop were successful in 50% (1 of 2) of the immunized rats, in comparison to 0% (0 of 2) both for the SS-looped and linear variants of these. Based on this finding, we then performed a detailed comparative study with peptides derived from the sequence $L_{56}-A_{79}$ of FSH- β (Fig. 6A). Unequivocally, these studies revealed the superior ability of the **T2/SS** peptide $C_{(S)}VYETC_{T2}RVPGNAHHADSLC_{T2}TYPVC_{(S)}$ to raise hFSH-crossreactive antibodies. This peptide comprises both a **T2** module (connecting the two cysteines that replace V_{61} and Y_{74}) and an additional SS bond between the cysteines replacing L_{56} and A_{79} . It raised hFSH-crossreactive antibodies in 100% of the rats (6 of 6), in comparison to 50% (1 of 2) and 0% (0 of 2) for the corresponding **T2** and linear variant. Even though the SS-alone peptides also showed activity in 2 out of 2 animals, the ELISA and neutralization titers were significantly lower in this case. Moreover, we recently observed much larger differences in hFSH cross-reactivities between antibodies elicited with **T2/SS**- and **SS**-peptides when studying longer variants of the FSH $\beta 3$ -loop (unpublished data).

Significant crossreactivity with rFSH (but not with hLH or hCG) was observed for all **CLIPS** antisera ($ET_{hFSH} \sim 10 \times ET_{rFSH}$). In this respect, the anti-FSH mAb 6602 was superior to any of the **CLIPS** peptide antisera as it did not show any sign of crossreactivity for rFSH ($ET < 1.0$) [35].

There is strong evidence that the activity of the **T2/SS** peptide is a direct result of appropriate fixation of the peptides conformation, as judged from circular dichroism (CD) spectroscopy studies in combination with ELISA-competition studies (Fig. 7). Despite this marked improvement in mAb-binding of the **T2/SS** peptide compared to linear, it is still orders of magnitude lower than that of intact human FSH. This is likely due to the fact that the β -loop does not cover the complete epitope of mAb 6602, but only the most important part of it (see also Fig. 5D).

Systematic variation of the **T2** position in the **T2/SS** peptide (59-76, 60-75, 61-74, 62-73, 63-72) showed that only two (63-72, 61-74) of the five **T2/SS** peptides were able to generate hFSH-crossreactive antibodies in rats (2 of 2; data not shown) [35]. In these cases, both amino acid side chains that were replaced by the **CLIPS** residue in the solvent-shielded hydrophobic core of the protein, while for the remaining three inactive peptides at least one of the two amino acid side chains was located in the solvent-exposed area. Changing the **CLIPS** from m-**T2** to o-**T2**, p-**T2**, to m-**P2** did not have a major impact on antibody generation. Yet, the best results were still obtained with the meta-oriented **T2**-scaffold.

Finally, we observed that immunization with **T2/SS**-peptides for which the configuration of the **CLIPS** cysteines

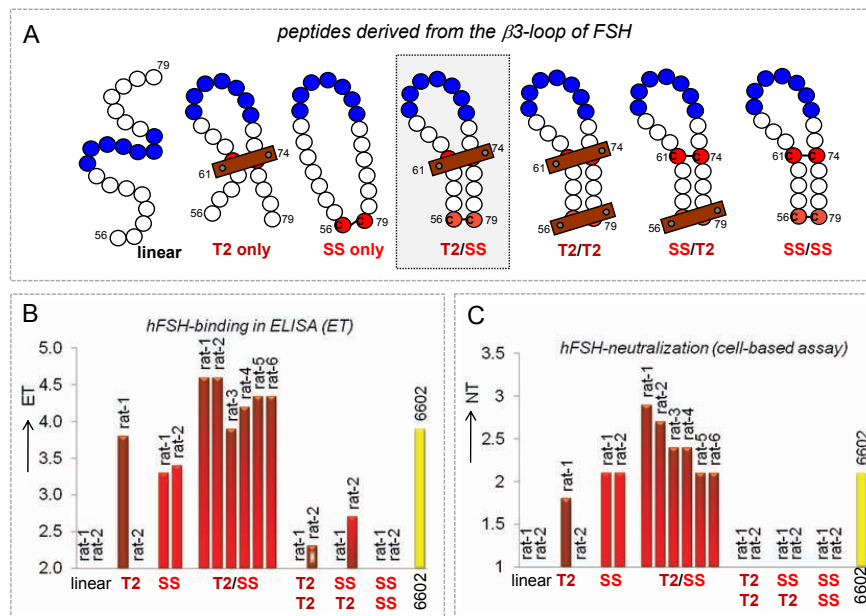


Fig. (6). **A**) Overview of linear, single- and double-constrained peptides (SS, **T2**, **T2/SS**, SS/SS, etc.) covering the β -loop sequence L₅₆-A₇₉ derived from human FSH- β ; **B**) hFSH binding of rat antisera raised with the designated peptides in ELISA (ET = ELISA titer: $^{-10}$ log values of serum dilution that gives 4x background OD_{405nm}-level), and **C**) neutralization titer (NT) of rat antisera compared to that of a 10 μ g/mL solution of mAb 6602. (NT = $^{-10}$ log values of highest serum dilution that shows neutralizing activity; neutralization was measured in Y1 cell-based assay, see [35] for details).

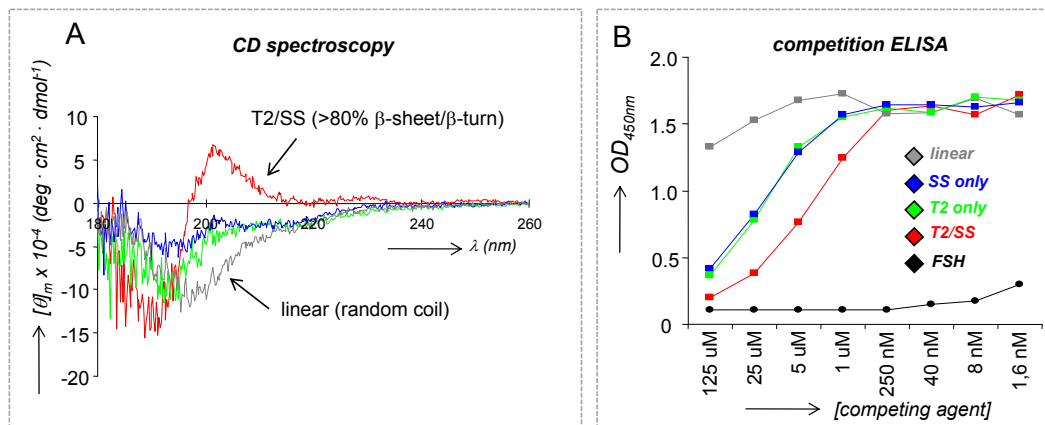


Fig. (7). **A**) CD spectroscopy measurements of aqueous solutions of linear, SS, **T2** and **T2/SS** peptide derivatives of hFSH β -loop sequence L₅₆-A₇₉. **B**) Ability of the peptides to inhibit the binding of mAb 5828 to surface-immobilized hFSH. These data provide evidence for the fact that increased structural organization in **T2/SS** is likely to be the reason for improved antigenicity and superior ability to raise hFSH-crossreactive antisera in rats.

was inverted (C_{61}/C_{74}) from L- to D-, either change at a single or both positions, totally prevented the formation of neutralizing antibodies (data not shown) [35]. Crossreactivity with or neutralization of hFSH was no longer observed ($ET/NT < 1.0$) for these antisera. This result is quite interesting in the sense that the **CLIPS** cysteines are not directly involved in antibody binding, but apparently induce a proper orientation of the crucial functionalities (R_{62} , G_{65} , A_{71} , D_{72} , L_{74}). This issue clearly underscores the importance of correct 3-dimensional peptide folding for successful antibody generation and demonstrates that **CLIPS** technology is apparently able to enforce this.

Binding Characteristics of Antisera Elicited with Linear, T2 and T2/SS Peptides

In order to obtain detailed information about the binding mode of the antisera elicited using **CLIPS** peptides, we performed competition experiments with a panel of substitution variants in ELISA [35]. Therefore, we synthesized 28 Ala-replacement (Gly for native Ala) variants of the 32-mer **CLIPS** peptide TFKELVYETC_{T2}RVPGAAHHADSLC_{T2}TYPVATQAH (52-83) and investigated the ability of these to effectively block the binding of the various antisera. The competition studies were performed in two different experimental setups, i.e.:

1. using surface-immobilized hFSH (visualizes only “desired” antibodies that crossreact with hFSH).
2. using surface-immobilized **CLIPS** peptide 52-83 (visualizes both “desired” and “undesired” antibodies).

Using these two different formats allowed us to highlight the characteristic differences in binding, i.e. defining the

crucial amino acids, between “desired” and “undesired” antibodies.

As a reference, we first studied binding competition for the two neutralizing mAbs (Fig. 8A). The data showed perfect correlation with replacement studies in peptide microarrays (Fig. 8E), and identified the amino acids R_{62} , G_{65} , A_{71} (mAb 5828 only), D_{72} , and L_{74} as being crucial for binding. Results obtained with **CLIPS** peptide 52-83 (data not shown) were very similar to the data obtained with surface-immobilized FSH. Only, the observed effects with 52-83 were much less pronounced than with hFSH.

We then studied a selection of peptide antisera, all elicited with β 3-loop derived peptides that differ in the extent of structural organization (linear, **T2**, **T2/SS**, Fig. 8B-D). These data demonstrate how structural organization drastically changes the type of antibodies that is formed. For example, the sera obtained with linear peptides did not show any cross-reactivity with hFSH (Fig. 8B). Obviously, this is due to the fact that binding occurs mainly at the N- (T_{52} , F_{53}) or C-terminal (Q_{81} , H_{83}) amino acids, which are not solvent-exposed at the hFSH protein surface. With **CLIPS** peptide 52-83 only a small subset of antibodies does crossreact with hFSH (Fig. 8C, format 1). Interestingly, the crucial amino acids are located mainly in the loop region, with some of these (R_{62} , D_{72}) being identical to those for the neutralizing mAbs. Finally, the antibodies obtained with **CLIPS/SS** peptides show a remarkably close parallel to the neutralizing mAbs 5828 and 6602 with respect to the crucial amino acids. There are some distinct differences (importance of E_{59} and T_{60}), which might be due to steric shielding of these amino acids as a result of glycosylation of FSH. Moreover, there is little to no difference between the data from formats 1 and 2,

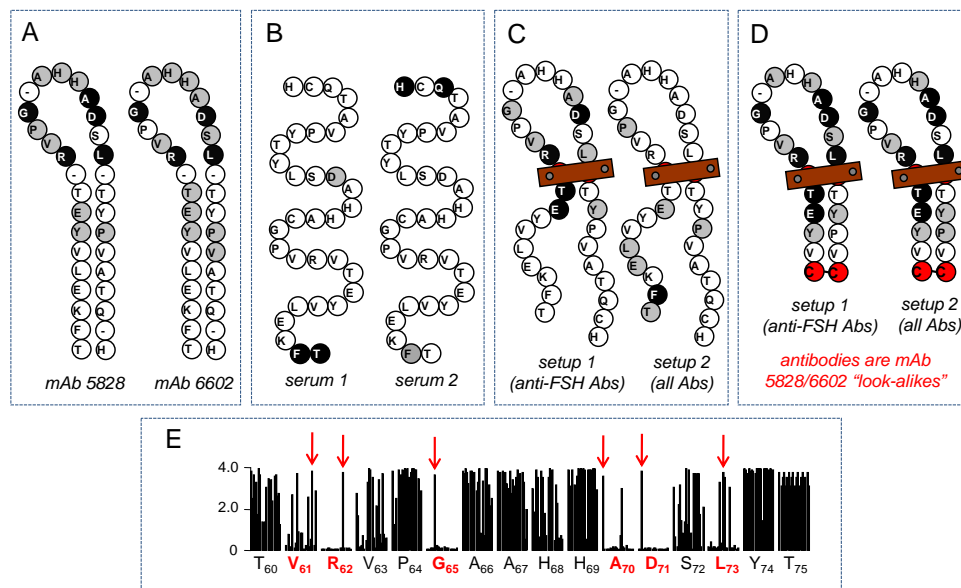


Fig. (8). Overview of the results from ELISA competition experiments with **A**) neutralizing mAbs, **B**) sera elicited with linear peptides (no observed crossreactivity with hFSH), **C**) with **CLIPS** peptides, and **D**) with **CLIPS/SS** peptides. Amino acids in black represent crucial amino acids (>50% inhibition of antibody binding), in grey important (>10% inhibition), and in white not important (<10% inhibition). Ala-replacement variants for cysteines were not included in this study. **E**) Microarray data for binding of mAb 6602 to full replacement analysis of **CLIPS** peptide C_{T2}TVRVPGAAHHADSLYTC_{T2}, showing perfect correlation with the data from the ELISA-competition experiments. Black bars are grouped in 16 sets of 19 each. Every bar represents one of 19 substitution variants of all 16 amino acids of **CLIPS** peptide C_{T2}TVRVPGAAHHADSLYTC_{T2} that were replaced by all other naturally occurring amino acids (cysteines were not substituted). Red arrows indicate positions of native amino acids.

which proves that the vast majority of generated antibodies is able to bind to hFSH.

EXAMPLE 2: MEMBRANE PROTEINS PART I: THE CD20 ANTIGEN

CD20 is a non-glycosylated 33-37 kD integral plasma membrane protein belonging to the MS4A family. It crosses the plasma membrane four times with both the N- and C-terminus located inside the cell. CD20 is specifically expressed on B cells from pre-B cell until plasmacytoid immunoblast stage (not on lymphoid stem cells and plasma cells) and is found on the surface of >99% of all B-cell lymphomas. Recent evidence confirmed that CD20 plays a role in regulating cell cycling and differentiation as well as being part of a membrane ion-channel [36-38].

Therapeutic successes using monoclonal antibodies (mAbs), either alone or in combination with chemotherapy, continue to raise interest for this relatively new class of drugs. One of the first targets for mAb-therapy was the CD20 antigen, which arguably represents the best validated target for immunotherapy [39]. During the past 10 years, therapeutic mAbs, and in particular the chimeric CD20 antibody rituximab, have changed the treatment paradigm for patients with B-cell neoplasms dramatically [40]. Building on this success, novel CD20 mAbs are being developed which are improved in terms of activity in Fc-dependent effector function and in terms of being fully human [27, 41].

Ofatumumab is a fully human mAb which has demonstrated exceptional activity in preclinical studies [27, 41-43]. It was found to induce stronger complement-dependent cytotoxicity (CDC) than rituximab by promoting more efficient binding of C1q to the surface of CD20-positive cells. Improved activity of this mAb is likely due to the fact that it docks to a different binding site on CD20 and demonstrates a reduced off-rate [39]. Ofatumumab has been shown to have substantial clinical activity as a single agent in the treatment of chronic lymphocytic leukemia and other B cell lymphomas [44-46].

Epitope Mapping Studies for A Series of Mouse and Human CD20 mAbs Using CLIPS™ Technology

In collaboration with GenMab B.V. (Utrecht, The Netherlands) we mapped the binding site of various CD20 mAbs [27]. To this end, we synthesized a set of overlapping **CLIPS** peptides (4-13 amino acids long) covering the extracellular sequences, encompassing amino acid residues 70-84 (small loop) and 140-184 (large loop) of CD20 (Fig. 9A) and screened these for binding with both human and murine mAbs. Most CD20 mAbs were able to bind well to similar, overlapping, epitopes as measured with both linear and **CLIPS** peptides. These epitopes encompassed residues A170 and A172, which were previously identified by Polyak and Deans as CD20 mAb epitope by means of mutagenesis studies [47]. However, for ofatumumab, binding was only observed when **CLIPS** peptides were used which, furthermore, revealed completely different binding sites for ofatumumab and the panel of known CD20 mAbs, including rituximab (Fig. 9B). Most intriguing was the observation that this novel epitope encompassed the small loop (AGIYAPI) of CD20. Based on these data it was concluded that ofatumumab is a unique monoclonal antibody that targets a distinct small loop epitope on the CD20 molecule. It is thought that the close proximity of the epitope to the cell membrane, situates ofatumumab very favorably for inducing complement-dependent cytotoxicity. Upon complement activation, short-lived activated complement protein intermediates are generated which can only travel a short distance before they are hydrolyzed. Presumably, the proximal positioning of CD20-ofatumumab-complement protein complexes is then optimal for effective deposition and complement-mediated attack of the tumor cell surface. **CLIPS** mapping of ofatumumab thus identified a novel critical epitope on CD20 and provided important insight into the mechanism of action of this therapeutic antibody.

EXAMPLE 3: MEMBRANE PROTEINS PART II: THE HUMAN CHEMOKINE RECEPTOR CCR5

CC Chemokine receptor 5 (CCR5) is a seven-transmembrane (7TM) chemokine receptor that is part of the

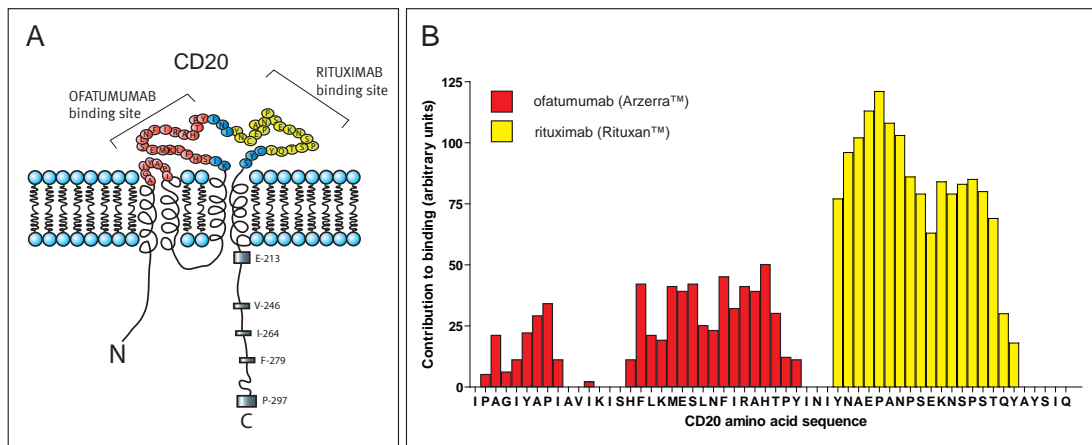


Fig. (9). **A**) Schematic representation of the two extracellular loops of the membrane-bound CD20 antigen; **B**) Epitope mapping data for ofatumumab (in red) and rituximab (in yellow), showing that both CD20 mAbs bind to a different binding site on CD20. The contribution of CD20 amino acids to binding in B is shown in arbitrary units [27]. It should be noted that the magnitude in AU represents a relative frequency of binding peptides containing the indicated amino acid and not a measure for relative binding strength. The lower bars for ofatumumab thus express a greater selectivity, i.e. it only binds **CLIPS** peptides with proper conformational constraints, whereas rituximab binds a larger proportion of peptides including **CLIPS** as well as linear peptides.

rhodopsin subfamily of G-Protein Coupled Receptors (GPCRs) [48]. It is the major coreceptor for HIV-1 entry [49,50]. CCR5 has been intensely studied since its central role in HIV-1 infection became apparent with the discovery that the absence of CCR5 expression (*via* a homozygous $\Delta 32$ deletion) provides resistance to HIV-1 infection [51-53]. CCR5 has been validated as a potential target for prophylaxis against HIV-1 infection and therapy of HIV-1/AIDS [54]. Small-molecule approaches to block the CCR5-HIV interaction have shown promise in clinical trials [55,56], but suffer from the development of resistance [57]. Recent experiments even suggest the future possibility of genetic disruption of CCR5 using zinc-finger nucleases in an attempt to reconstitute immune function in HIV-infected people [58]. Furthermore, anti-CCR5 antibodies are being investigated to block the CCR5-HIV interaction as a potential therapeutic strategy for HIV-1/AIDS [59,60]. A similar approach based on active immunization with GPCR-derived immunogens could have potential as a prophylactic agent for preventing HIV infection [61-63].

Epitope Mapping Studies Using CLIPS™ Technology

In light of this, we tried to reconstruct the binding site of the CCR5-neutralizing mAb 2D7, a CCR5 antibody with strong neutralizing activity [64], by means of CLIPS technology. We synthesized overlapping libraries consisting of linear, monocyclic (T2) and bicyclic (T3) peptides, the sequences of which were all derived from the CCR5-extracellular domain (ECD; Fig. 10A). In total more than 1500 CLIPS-peptides were screened for binding with 2D7. The results showed that only peptides comprising the ECL2a-derived sequence 170-QKEGLHY-176 displayed affinity for the antibody. Binding was not observed for any of the sequences not incorporating these 7 residues. Moreover, binding of CLIPS peptides incorporating 170-QKEGLHY-176 was intense ($OD > 2$; Fig. 10B) and detectable even at very low antibody concentration ($\sim 0.1 \mu\text{g/mL}$). In contrast to this, binding of linear peptides was most often not detectable, occasionally only at the highest possible concentration ($10 \mu\text{g/mL}$).

Detailed mapping studies of the ECL2a-epitope provided valuable information on the mode of antibody binding of mAb 2D7 (Fig. 10C), nicely complementing the data from Ala-mutation studies with intact CCR5 (Fig. 10D) [64]. For example, replacement studies on the CLIPS peptide 168-C_{P2}RSQKEGLHYTC_{P2}-177 revealed that any replacement of the amino acids K₁₇₁, E₁₇₂ and G₁₇₃ led to a complete loss of mAb-binding (Fig. 10C). The amino acid Q₁₇₀ also showed significant decreases in 2D7 binding for most substitutions, albeit with less stringent demands (between 1-2 substitutions tolerated without major loss of binding). Despite this, the peptides with the native amino acids at these positions were still the strongest binders of all. Moreover, CLIPS peptide 170-C_{P2}QKEGLHYC_{P2}-176 was found to be the shortest possible sequence showing binding to 2D7 (data not shown), which is indicative for the importance of both Q₁₇₀ and Y₁₇₆ for binding. The relatively low sensitivity to substitution of Y₁₇₆ can be best explained by assuming a conformational role for this amino acid, which somehow may interact with one of the crucial amino acids by positioning the side chains of these in the correct orientation for binding. This seems fully in accordance with the fact that Y₁₇₆ was not identified as a crucial amino acid in Ala-mutation studies on intact CCR5 (Fig. 10C) [64].

All other amino acids (T₁₆₇, R₁₆₈, L₁₇₄, H₁₇₅, T₁₇₇) showed a very low level of resistance to substitution (>10 substitutions tolerated), most likely indicating that they do not specifically interact with the antibody. For some of these, significant binding improvements were observed upon substitution with others (L₁₇₄/Q, H₁₇₅/N, T₁₇₆/FIKL), which might be due to the fact that the native amino acids stabilize conformations of the peptides that are suboptimal for 2D7 binding.

Competition ELISA Studies with CLIPS Peptides Derived from ECL2a

In order to confirm the mapping data we performed a series of ELISA competition experiments. Moreover, we wanted to address in more detail the effect of conformational

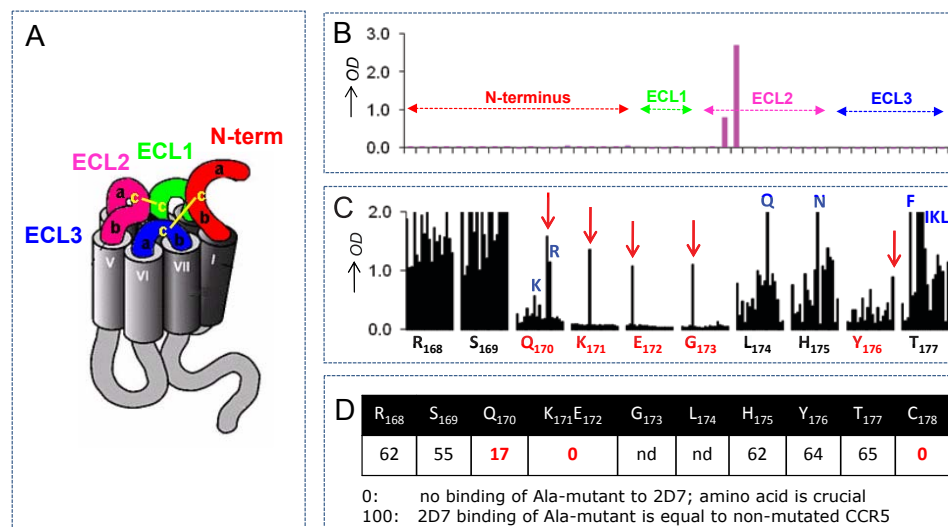


Fig. (10). A) Schematic illustration of a GPCR, B) Epitope mapping data for mAb 2D7 using overlapping libraries of CLIPS peptides derived from the ECL domain of CCR5, C) binding data for mAb 2D7 to a full replacement analysis of CLIPS peptide *C_{P2}RSQKEGLHYTC_{P2} (red arrows indicate native amino acids), and D) data from Ala-mutation studies on intact CCR5 [64].

fixation of the peptides on mAb 2D7 binding. Therefore, we studied the ability of various ECL2a-derived peptides to inhibit the binding of mAb 2D7 to a surface-immobilized CCR5 antigen. We used the **CLIPS** peptide C_(S)FTRC_{T2}QKEGLHYTC_{T2}SSHC_(S) (purified by HPLC) for surface-immobilization as a valuable substitute for CCR5, since purified CCR5 is not available as it is an integral membrane protein. We compared two sets of peptides. In the first set, we studied the effect of conformational fixation (linear, **T2** and **T2/SS**) on binding (Fig. 11A and B). Secondly, we compared the effect of substituting one (mimic-1) or several (mimics 2-3) of the (non-essential) native amino acids on 2D7 binding (Fig. 11C and D).

The competition data confirm the results from the epitope mapping and replacement analysis studies. The linear peptide TRSQKEGLHYTSGSC did not show any competition at all, not even at the highest concentration used (500 μ M; Fig. 11B). In contrast to this, the **T2**-constrained peptide TRC_{T2}QKEGLHYTC_{T2}GSC clearly showed activity (>50% competition) at 500 μ M. Finally, the **T2/SS** C_(S)RC_{T2}QKEGLHYTC_{T2}SC_(S) proved most active of all three, showing an ~80% decrease in binding of 2D7 at 165 μ M. Replacement of some of the amino acids in **T2**-peptide TRC_{T2}QKEGLHYTC_{T2}GSC, previously identified as non-essential, also improved the ability to reduce 2D7 binding (Fig. 11C and D). Mimic-1 (Q₁₇₀/R only) and -2 (4 replacements: Q₁₇₀/R, L₁₇₄/Q, H₁₇₅/N, and T₁₇₇/I) showed improved activity compared to native **T2**-peptide, in particular at the two highest concentrations (500 and 160 μ M). For mimic-3 (only 3 replacements: L₁₇₄/Q, H₁₇₅/N, and T₁₇₇/I) the improved activity in competition was still visible at 18 μ M, which confirmed our earlier finding that Q₁₇₀ is important for 2D7 binding. The improved activity is likely due to replacement of either L₁₇₄, H₁₇₅, or T₁₇₇, or a combination of these.

Antibody Generation Using CLIPS™ Technology

In a pilot study, we investigated the potential of the strongest binding ECL2a-derived **CLIPS** peptides to generate antibodies capable of neutralizing the bioactivity of CCR5. We included 2 different ECL2a-mimics (1/2) in two different formats (1/2 or 1*/2*) and two mimics derived from the N-terminus (mimic 4: MDYQVSSPIY-DINYYTSEPC_{T2}QKINVK_CT₂) and the N-terminus + ECL1 (mimic 5: MDYQVSSPIYDINYYTSEPC_{T3}QKINVK_CT₃AAQWDFGNTMC_{T3}). ECL2a-mimics 1-2 all have incorporated 1 (mimic 1: TRC_{T2}RKEGLHYTC_{T2}) or 4 (mimic 2: TRC_{T2}RKEGQNYIC_{T2}) amino acid replacements in comparison to the native ECL2a-sequence (167-TRSQKEGLHYTC-178). Mimics 1/2 and 1*/2* differ only in the mode of conjugation to the carrier protein (KLH), i.e. either *via* a C-terminal extension (GSC-sequence; mimics 1/2) or *via* the use of an SH-functionalized **T2** scaffold (mimics 1*/2*). BALB/c mice were immunized (4 mice per mimic) and the sera were tested for CCR5-positive staining in flow cytometry (Fig. 12A).

The sera from ECL2a mimic-2 gave by far the best results (3 of 4 sera CCR5-positive). Splenocytes were isolated from all four CCR5-positive mice and used to generate hybridomas. 180 viable mAbs were selected, from which 32 produced antibodies that showed positive staining for CCR5 in flow cytometric analysis (Multispan, Hayward, CA, USA; Fig. 12B). The charts for mAb 2D7 and mAb 9 at two different concentrations (~1 and 0.1 μ g/mL) are shown (Fig. 12C and D).

In summary, these data show the potential of **CLIPS** peptide mimics to generate mAbs for GPCRs, in particular for CCR5. Even though the data were not optimized for the application, this example provides a good perspective of what might be achievable for related GPCRs or even for

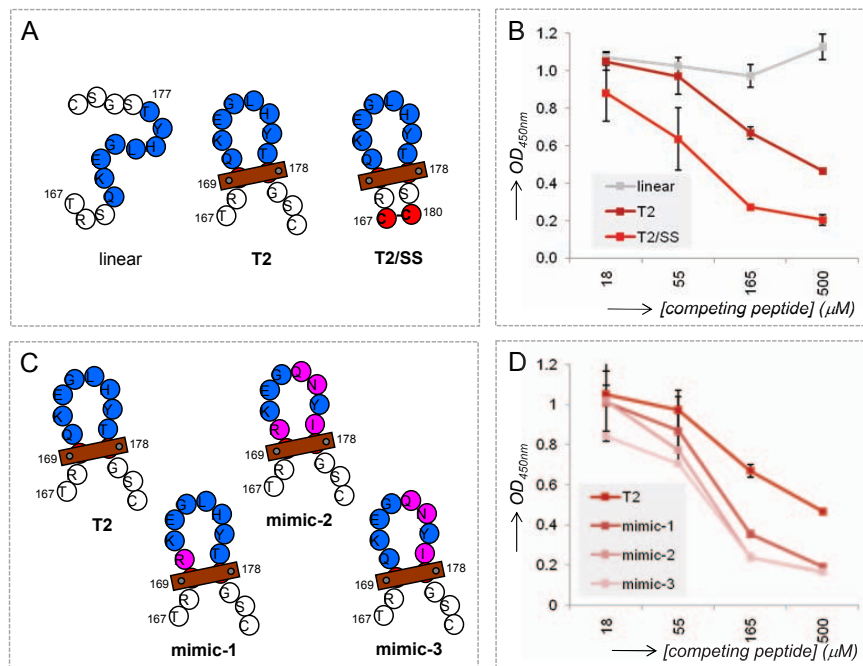


Fig. (11). Competition studies in ELISA with two different sets of peptides derived from the ECL2a sequence 167-TRSQKEGLHYTC-178; **A-B**) results for linear, **T2** and **T2/SS** constrained peptides; **C-D**) results for **T2** peptide TRC_{T2}QKEGLHYTC_{T2}GSC and corresponding mimics 1 (Q₁₇₀/R), 2 (Q₁₇₀/R; L₁₇₄/Q; H₁₇₅/N; T₁₇₇/I) and 3 (L₁₇₄/Q; H₁₇₅/N; T₁₇₇/I). ELISA-measurements were performed in duplicate.

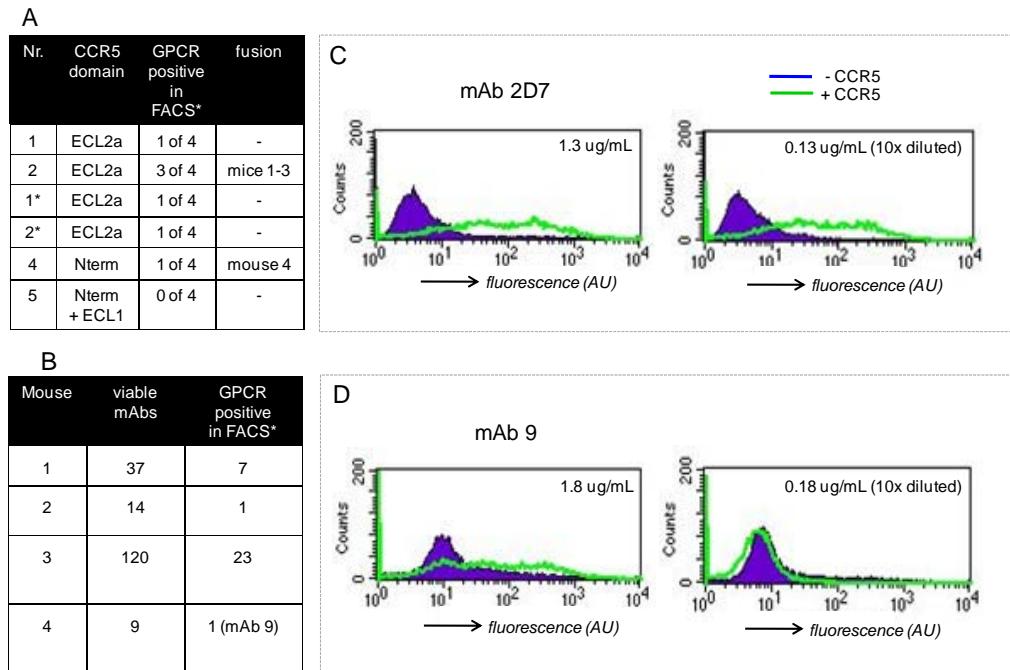


Fig. (12). Overview of results from flow cytometric analysis on CCR5-expressing cells for **A**) antisera elicited with six different CCR5-mimics, **B**) 180 viable hybridoma clones generated with splenocytes from mice with CCR5-positive antisera. Flow cytometry data for **C**) mAb 2D7 (positive control) and **D**) mAb 9. Binding of mAb 2D7 and mAb 9 to CCR5-expression cells (+ CCR5) was detected by an anti-mouse IgG1-FITC conjugate. As a control, background binding was assessed on CCR5 negative non-transfected cells (- CCR5).

structurally more complicated membrane-bound proteins (e.g. ion-channels).

CONCLUSIONS AND FUTURE PERSPECTIVES

In conclusion, **CLIPSTM** technology proves to be useful in protein binding site reconstruction, not only for epitope mapping purposes, but more importantly also to raise neutralizing antibodies. The technology is particularly attractive for its ease of application, the extremely fast and clean conversions of the peptides involved, and the mild reaction conditions combined with the chemical robustness of the cyclised peptides. Application of this technology to map mAbs against therapeutically relevant targets (hFSH, hCG, CD20, and CCR5) illustrates clearly how the technology can be used to define binding sites at the single amino acid level. The work presented holds great promise for application to other protein targets, in particular those with considerable structural complexity (ion-channels, viruses).

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