Studies of a Murine Monoclonal Antibody Directed against DARC: Reappraisal of Its Specificity

Dorota Smolarek1,4,5,10, Claude Hattab4,5,8,9, Anna Buczkowska1,6, Radoslaw Kaczmarek1, Anna Jarzab3, Sylvie Cochet4,5,8,9, Alexandre G. de Brevern4,5,8,9, Jolanta Lukasiewicz2, Wojciech Jachymek2, Tomasz Niedziela2, Magdalena Grodecka1, Kazimiera Wasniowska1,7, Yves Colin Aronovici4,5,8,9, Olivier Bertrand4,5,8,9, Marcin Czerwinski1,7,*

1 Laboratory of Glycoconjugate Immunochemistry Ludwik Hirszfeld Institute of Immunology and Experimental Therapy, Wroclaw, Poland, 2 Laboratory of Microbial Immunochemistry and Vaccines Ludwik Hirszfeld Institute of Immunology and Experimental Therapy, Wroclaw, Poland, 3 Laboratory of Medical Microbiology, Ludwik Hirszfeld Institute of Immunology and Experimental Therapy, Wroclaw, Poland, 4 INSERM, UMR_S1134, F-75015 Paris, France, 5 Institut National de la Transfusion Sanguine, F-75015 Paris, France, 6 Institute of Biochemistry and Biophysics, Warsaw, Poland, 7 Institute of Physiotherapy, Faculty of Physical Education and Physiotherapy, Opole University of Technology, Opole, Poland, 8 Université Paris-Diderot, Sorbonne Paris Cité, F-15013, Paris, France, 9 Laboratory of Excellence GR-Ex, Paris, France, 10 Division of Transplantation Immunology and Mucosal Biology, MRC Centre for Transplantation, King’s College London, Guy’s Hospital, London, United Kingdom

* czerwinski@iitd.pan.wroc.pl

Abstract

Duffy Antigen Receptor for Chemokines (DARC) plays multiple roles in human health as a blood group antigen, a receptor for chemokines and the only known receptor for Plasmodium vivax merozoites. It is the target of the murine anti-Fy6 monoclonal antibody 2C3 which binds to the first extracellular domain (ECD1), but exact nature of the recognized epitope was a subject of contradictory reports. Here, using a set of complex experiments which include expression of DARC with amino acid substitutions within the Fy6 epitope in E. coli and K562 cells, ELISA, surface plasmon resonance (SPR) and flow cytometry, we have resolved discrepancies between previously published reports and show that the basic epitope recognized by 2C3 antibody is 22FEDVW26, with 22F and 26W being the most important residues. In addition, we demonstrated that 30Y plays an auxiliary role in binding, particularly when the residue is sulfated. The STD-NMR studies performed using 2C3-derived Fab and synthetic peptide corroborated most of these results, and together with the molecular modelling suggested that 25V is not involved in direct interactions with the antibody, but determines folding of the epitope backbone.
Introduction

Duffy Antigen Receptor for Chemokines (DARC) is a seven-transmembrane domain glycoprotein present on red blood cells (RBC) and post-capillary endothelial cells or Purkinje cells. DARC plays an important role in human health, being a blood group antigen, the only known receptor for *Plasmodium vivax* parasite that causes malaria and a receptor for chemokines.

DARC glycoprotein carries blood group antigens Fya and Fyb, which are encoded by two allelic genes, designated FY^a^01 and FY^b^02 respectively. The FY^a^/FY^b^ polymorphism depends on the presence of Gly or Asp residue at position 42 in the polypeptide chain of DARC (Gly in Fya, Asp in Fyb). The third common allele called FY^o2N.01 occurs mostly in Africans or African-American people and results in the lack of DARC expression on erythrocytes, but not on other tissues. That phenotype, called Fy(a-b-) is caused by a single nucleotide polymorphism at position -67T>C in the GATA box of DARC gene promoter [1] and it was thought for decades that it was associated with resistance of red blood cells to infection by *Plasmodium vivax* merozoites [2]. However, recent data cast some doubt on this dogma, suggesting that another receptor for *P. vivax* may exist [3]. *P. vivax* is the most widespread species of parasite causing malaria with about 2.85 billion of people living in endemic regions. DARC is recognized by *P. vivax* Duffy Binding Protein (PvDBP), and thus serves as a receptor for *P. vivax* merozoites [4, 5]. The crystal structure of DARC-PvDBP complex has been recently resolved [5].

DARC is also a chemokine receptor belonging to G-Protein Coupled Receptor (GPCR) family [6, 7], and this function most probably determines DARC’s involvement in malignancies [8–10], inflammation [11], HIV infection or AIDS progression [12–14]. DARC binds chemokines [15], but lacks the DRY motif which is typical for this family of receptors and required for signal transduction, hence it was considered a “silent chemokine receptor” [16, 17]. Recently, it was proposed to call it Atypical Chemokine Receptor 1 (ACKR1) [18]. It was shown that DARC plays an important role in regulation of the chemokine level, but in a different manner depending on its localization: on the erythrocytes it serves as a reservoir of chemokines, while on the endothelial cells it facilitates chemokines translocation through the vessel walls.

DARC contains three N-linked oligosaccharide chains located at Asn-16, 27, and 33 [19]. These moieties are mostly of triantennary complex type, terminated with sialic acid residues [19–21]. In addition, Tyr-30 and 41 are sulfated [22], and sulfation of Tyr-41 increases binding of PvDBP to DARC approximately thousand times [22].

There are several epitopes recognized by human and murine antibodies on DARC: two of them determine the Fya/Fyb blood group status, the other is Fy3, a conformational epitope consisting of fragments of extracellular domains 1 and 3, and a linear epitope called Fy6, present on the ECD1 of the protein.

Specificities of several monoclonal anti-Fy6 antibodies have been precisely defined [23–25]. It was found that all tested antibodies with anti-Fy6 specificity recognize linear epitopes containing ^19QLDFEDV^25 sequence located between two potential N-glycosylation sites of Duffy protein, Asn-16 and Asn-27. One of the widely used anti-DARC antibodies, monoclonal antibody 2C3 with anti-Fy6 specificity, was obtained after immunization of mice with CHO cells transfected with the vector encoding DARC. Its epitope was evaluated by Wasniowska et al. using Pepscan analysis, and determined to be a linear sequence ^22FEDVW^26, in which only Asp-24 may be replaced by other amino acids without influencing binding of the antibody [24]. In contrast, Tournamille et al., using recombinant forms of DARC expressed in K562 cells suggested that Trp-26 can be substituted by other amino acids without losing reactivity to 2C3 antibody. In addition, it was claimed that the antibody does not bind to the recombinant form of Duffy antigen, in which Tyr-30 is substituted by another amino acid [25].
Since antibodies that recognize Fy6 epitope can inhibit interaction between chemokines [26] or PvDBP [27] and DARC, the exact determination of epitope recognized by 2C3 antibody is of great importance. The aim of our study was to precisely characterize the epitope recognized by 2C3 antibody and to resolve discrepancies between the two abovementioned reports. To this end, we prepared two sets of constructs: one set comprised the full-length consensus DARC or mutants with substitutions introduced in the first extracellular domain (ECD1); these constructs were expressed in K562 cells. The other set comprised the consensus or mutant ECD1 (with the same substitutions as in the full-length DARC mutants) fused in-frame to nuclease from *Staphylococcus aureus* and expressed in *E. coli*. The proteins obtained in *E. coli* or K562 cells were used to evaluate the binding of 2C3 antibody using ELISA, SPR and flow cytometry. In addition, we employed saturation transfer difference spectroscopy (STD-NMR) of a 2C3-derived Fab fragment with a synthetic peptide encompassing the Fy6 epitope in order to precisely define how the side chains of amino acids interact with the Fab.

**Materials and Methods**

**Monoclonal antibody**

MAb 2C3 (clone name NaM185-2C3) was obtained previously by immunizing mice with CHO cells transfected with the vector encoding DARC protein [24].

**Recombinant proteins used as reagents for 2C3 MAb studies**

Constructs with substitutions of amino acids located within or in the vicinity of the Fy6 epitope were prepared by mutagenesis using plasmid containing first extracellular domain of DARC followed by nuclease from *Staphylococcus aureus* as a template [26].

The protein (called thereafter ECD1-nuc) was expressed in *E. coli* BL21 strain (New England Biolabs, Ipswich, MA, USA) grown in autoinduction medium, and purified from whole cell extract using ion exchange Capto-S column (GE Healthcare, Little Chalfont, UK) followed by reversed-phase liquid chromatography on a Vydac TCP 104 column (Grace, Columbia, MD, USA).

The full-length DARC was purified from K562 cells expressing DARC, essentially as described [26] by immunochromatography on an immobilized camel antibody fragment (nanobody), but using a column format. Briefly, 16 × 10^8 cells were lysed in 300 ml of lysis buffer (20 mM Tris-HCl pH 8.0 containing 0.15 M NaCl, 5 mM EDTA, 1% Triton X100 and protease inhibitors (Complete, Roche, Basel, Switzerland). After centrifugation, supernatant was loaded at 15 ml/hour onto a 1 ml affinity column (prepared using NHS-activated Sepharose and purified anti-DARC nanobody). After washing the column with lysis buffer followed by 20 mM Tris-HCl pH 8.0 containing 0.15 M NaCl and 0.3% C12E8 detergent (Sigma), DARC was eluted by passing onto the column 5 ml of peptide DFEDVW (dissolved in the latter buffer at 4 mg/ml concentration). Column was further rinsed with 0.1 M Glycine pH 2.8. DARC was freed of peptide by dialysis. The DARC purity was evaluated by SDS-PAGE (S1 Fig.).

**Cloning and expression of 2C3 Fab fragment**

cDNA encoding variable regions of the 2C3 Fab fragment was cloned by PCR using total RNA extracted from 2C3-producing hybridoma cells. The amplified DNA was cloned into pComb3H vector, expressed in *E. coli* and purified as described previously [28, 29]. The Fab purity was evaluated by SDS-PAGE (S2 Fig.).
ELISA

Microtiter 96-well plate (MaxiSorb, Nunc, Roskilde, Denmark) was coated with wild-type ECD1-nuc 0.1 μg of protein per well in 50 μl of carbonate buffer pH 9.6, overnight at 4°C. Non-specific binding was blocked by incubation for 1 hour at room temperature with TBS (Tris-buffered saline: 20 mM Tris-HCl, and 150 mM NaCl pH 7.4) containing 5% non-fat dry milk. After washing the wells once with TBS, 50 μl of 2C3 MAb or 2C3 Fab fragments appropriately diluted in TBS were added to each well, and the mixture was incubated for 1 h at room temperature. After five washes with TBS, 50 μl of a 1000 x dilution in TBS of rabbit anti-Fab antiserum conjugated with alkaline phosphatase or avidin-alkaline phosphatase 1000 x diluted (Pierce, Rockford, IL, USA) were added to each well and the plates were incubated for 1 h at room temperature. To develop the reaction, p-nitrophenylphosphate was used (1 mg/ml in 60 mM carbonate buffer, pH 9.5, containing 1 mM MgCl2); the OD$_{405}$ was determined using Perkin Elmer EnSpire 2300 Multilabel Reader (Waltham, MA, USA). All experiments were performed in triplicates.

Inhibition of the antibody binding was evaluated by incubating appropriately diluted MAb or Fab with serially diluted ECD1-nuc mutants for 1h. The samples were then applied on the coated plate and incubated for 10 min, and the bound antibody was detected as described above. The percentage of inhibition was calculated as: $[1- (OD_{405} \text{ of sample})/(OD_{405} \text{ of control})] \times 100$.

Surface plasmon resonance (SPR)

All experiments were performed in the Biacore X100 machine using CM5 type sensor chips (GE Healthcare), the running buffer was HBS-EP (GE Healthcare). To immobilize proteins, amine coupling procedure was used according to the manufacturer’s instructions. 2C3 MAb or 2C3 Fab was immobilized in the flow cell, and murine monoclonal anti-Kx2 antibody [30] was immobilized in the reference cell. Approximately 400 RU of protein was immobilized in each flow cell. ECD1-nuc with wild type or mutated sequences were used as analytes, and each construct was evaluated in 5 different concentrations. 50 mM HCl was used for regeneration of the chip. The association phase (binding of ligand) was run for 100 sec., and then dissociation was continued for 600 sec. The results were processed using BIAevaluation software and 1:1 Langmuir binding model.

For SPR inhibition assay, 2000 RU of WT ECD1-nuc construct was immobilized on Fc2 channel of a sensor chip as described above, with S. aureus nuclease in the Fc1 reference cell. 2C3 MAb (culture supernatant diluted 40 times) was incubated with the ECD1-nuc constructs at room temperature for one hour, the samples were then applied to the chip, the association phase was continued for 100 sec., and the RU value was read at 90 sec. The obtained values were used to calculate the concentration of each ECD1 mutant causing 50% inhibition of binding of 2C3 to ECD1-nuc. The synthetic peptides with sequences DFEDVWNSSYG and DFEDVWNSSYG(SO$_3$H)G were obtained from Cambridge Research Biochemicals, Cleveland, UK.

To evaluate how the presence of a negative charge on tyrosine residues influences the antigen binding, the ECD1-nuc construct was phosphorylated by Src kinase (Cell Signalling Technology, Danvers, MA, USA) and binding of 2C3 Fab to the phosphorylated ligand was studied. Phosphorylation of proteins already immobilized on the sensor chip was performed in the Biacore apparatus. Constructs of ECD1-nuc were used as a ligand and nuclease was immobilized in the reference cell (the phosphorylation site on Tyr-54 of nuclease was removed by mutagenesis from both ECD1-nuc and nuclease constructs). The phosphorylation reaction mixture was prepared by mixing Src Kinase (25 ng/μl final concentration), calcium-calmodulin (Sigma, 58 ng/μl final concentration), and ATP (73 μM final concentration) in the buffer supplied by...
the enzyme manufacturer. The mixture was injected at 5 μl/minute in two pulses of 1060 and 470 seconds onto the chip equilibrated at 37°C.

DARC purified from DARC-expressing K562 cells by immunochromatography on immobilized anti-DARC VHH (S1 Fig.) was immobilized at ca. 400 RU on the analysis flow cell, the reference cell was only activated and deactivated.

Analysis of the influence of glycosylation on binding to DARC was performed by in situ deglycosylation of purified DARC immobilized on the chip. Reaction mixture prepared by diluting N-glycosidase (New England Biolabs, Ipswich, MA, USA) to 50 UI/μl in the buffer recommended by the manufacturer was injected three times for 1000 sec. each time, the temperature was set to 37°C and flow rate to 5 μl/min. To evaluate efficiency of the deglycosylation process, *Sambucus nigra* lectin (Vector Labs, Burlingame, CA, USA) in concentration 70 nM and 7 nM was injected.

**Transfection of K562 cells**

The mutants were prepared using the Quickchange kit according to the manufacturer’s instruction (Stratagene, La Jolla, CA, USA). 2 × 10⁶ of K562 cells (obtained from ATCC, Manassas, VA, USA, clone number CCL-243) were transfected with vector encoding DARC using Amixa Nucleofector system (Lonza, Basel, Switzerland). The DARC-expressing cells were selected for resistance to geneticin, as described [19].

**Flow Cytometry analysis**

Approximately 10⁶ of K562 cells transfected with vector encoding DARC were used in each analysis. After washing with PBS, the cells were incubated for 1 hour with either one of three antibody solutions in PBS-0.2% BSA: i) anti-mouse IgG (Sigma, St. Louis, MO, USA) as a negative control, ii) anti-Fya human antibody (clone 665, kindly provided by Dr. F. Buffiere, Bordeaux, France) as a positive control [25], iii) 2C3 MAb culture supernatant. The cells were then washed 3 times with PBS, and incubated in the dark with secondary antibody conjugated to PE (anti-mouse or anti-human, Becton-Dickinson, Franklin Lakes, NJ, USA) for 40 min. After three washes the cells were resuspended in 300 μl of PBS and evaluated by flow cytometry (FacsCalibur, Becton-Dickinson).

**NMR spectroscopy**

The NMR spectra of the synthetic peptide (DFEDVWNSSYG) were obtained for 2H2O solutions at 25°C with a Bruker AVANCE III 600 MHz spectrometer (Bremen, Germany), using acetone (δH 2.225 and δC 31.05) as the internal reference. The main groups of signals were assigned by one- and two-dimensional experiments (COSY, TOCSY, ROESY, HMBC, and HSQC-DEPT). In the TOCSY experiments the mixing times were 30, 60, and 100 ms. The delay in the HMBC experiments and the mixing time in the ROESY experiments were 60 and 200 ms, respectively. The data were acquired and processed using standard Bruker software. The processed spectra were assigned with help of the SPARKY program [31].

One-dimensional STD NMR experiments were performed at 25°C with a 5-mm inverse-detection QCI cryoprobe equipped with z-gradient. Spectra were recorded using the pulse programs bundled with the Topspin software. The excitation sculpting pulse sequences were used to suppress water signals in the spectra. The broad resonances of a protein were suppressed with a 20-ms spin-lock pulse. The protein was irradiated at δH –0.5 ppm and 9.5 ppm (on-resonance) and δH30 ppm (off-resonance) with a train of Gaussian shaped pulses (50 ms) and the data were collected as a pseudo2D, split into 1D on-resonance and off-resonance spectra and followed by calculation of the difference spectra.
The setup of the STD NMR experiments was optimized in a series of experiments with ligand-protein and ligand-only samples to ensure that the irradiation at the selected frequency did not affect the ligand. The saturation time used in the experiments was 2 s. Samples (total volume, 160 μl) were prepared in 3 mm NMR tubes, using PBS made with $^2$H$_2$O, pH 7.5 (not corrected for $^2$H$_2$O). Fab/peptide ratios of 1:46 (21 μM Fab with ~1 mM peptide) were used.

Analysis of Peptides Structure and Folding

The secondary structure and folding of DFEDVWNSSYG and peptides with Val-25 substitutions were predicted and calculated by PEP-FOLD, an online resource for de novo peptide structure prediction in aqueous solutions [32]. The PEP-FOLD provides pdb (Protein Data Bank) files with representation of different peptide models combined in clusters. The models obtained by molecular modeling of peptides with the lowest sOPEP energy (sOPEP—Optimized Potential for Efficient Structure Prediction) were used for further analysis. The pdb files were visualized and analyzed by Chimera software and the distances between Trp-26, Tyr-30 and Phe-22, residues involved in the peptide-antibody interaction were calculated [33].

Results

Evaluation of the epitope recognized by 2C3 monoclonal antibody

Soluble ECD1-nuc constructs with consensus-type sequence or the sequence containing amino acid substitutions were tested by ELISA for ability to inhibit binding of the antibody to ECD1-nuc immobilized on a polystyrene plate. Profiles of inhibition of 2C3 MAb by different ECD1-nuc constructs are shown in S3 Fig. The binding strength of a given construct to antibody was calculated as the concentration of protein required for 50% inhibition of binding to the immobilized wild type construct (Table 1). The results demonstrate that only constructs in which Trp-26 is present in the epitope can inhibit, at low concentration, binding of the 2C3 antibody to coated ECD1-Nuc (Table 1, compare rows 1-3 with rows 4-8). Substitution of this residue by another aromatic amino acid or Ala results in a decrease of inhibition (Table 1, rows 6-8). However, this effect was somewhat compensated for when a negative charge was introduced at position 30 by replacing Tyr with Glu (Table 1, rows 4 and 5).

The influence of Tyr-30 sulfation was tested using sulfated and non-sulfated synthetic peptides encompassing amino acid residues 21-31 of ECD1 and evaluating ability to inhibit binding to ECD1-Nuc: results of the inhibition of 2C3 MAb binding to ECD1-Nuc by these peptides evaluated by ELISA is shown in Fig. 1. The SPR analysis revealed that concentration

| No. | Sequence     | Concentration required for 50% inhibition [μg/ml] |
|-----|--------------|--------------------------------------------------|
| 1   | FEDVWNSSY    | 0.17±0.19                                       |
| 2   | FEDVWNSE     | 0.22±0.12                                       |
| 3   | FEDVWNSSA    | 0.3±0.34                                        |
| 4   | FEDVFNSSA    | 28±1.23                                         |
| 5   | FEDVANSSA    | 48±1.45                                         |
| 6   | FEDVFNSSA    | >100                                             |
| 7   | FEDVANSSY    | >100                                             |
| 8   | FEDVANSSA    | >100                                             |

Concentrations of soluble ECD1-nuc constructs required for 50% inhibition of antibody binding to coated wild type ECD1-nuc are shown.

doi:10.1371/journal.pone.0116472.t001
required for 50% inhibition of 2C3 MAb binding is 410 nM (±40) for DFEDVWNSSYG and 252 nM (±20) for DFEDVWNSSY(SO3)G. Thus, sulfation of Tyr-30 is associated with a higher binding ability of the peptide to the antibody. In addition, to further investigate the role of sulfation of Tyr-30, we modified this residue in ECD1-nuc immobilized on a Biacore chip. Because a sulfotransferase is not commercially available, we used Src kinase to introduce a phosphate to the Tyr-30 residue to mimic a sulfate group. As a control, we used an anti-phosphotyrosine antibody which bound to ECD1 only after Src kinase treatment (data not shown). The kinetic data showed that phosphorylation of Tyr-30 caused an increase of 2C3 affinity, mainly by increasing \( k_a \) (Table 2 rows 1 and 4).

This set of experiments was completed with the studies performed using transfected K562 cells expressing wild type or mutated DARC. The binding of 2C3 antibody was evaluated by flow cytometry, and the strength of binding was estimated as a ratio between the percentages of cells bound by 2C3 MAb to the percentage of cells bound by anti-Fya antibody (clone 655), which recognizes the YGANLE epitope [25]. We found that only the cells transfected with constructs encoding Trp at position 26 were bound by the antibody with high affinity (Table 3 rows 1-5), whereas substitution of that residue with alanine resulted in a dramatic decrease of

**Figure 1. Inhibition of 2C3 MAb binding to ECD1-nuc.** The antibody was incubated with serially diluted DFEDVWNSSYG (circles) or DFEDVWNSSY(SO3)G (squares) peptides. Subsequent binding to ECD1-nuc was evaluated by ELISA as described in Materials and Methods. The difference in % of inhibition was statistically significant in the range of concentrations 3-60 nM (\( P < 0.05 \), two-tailed Mann-Whitney U test).

**Table 2. Kinetic constants of 2C3 Fab binding to ECD1-nuc and purified DARC measured by SPR.**

| Ligand                          | \( k_a (M^{-1}s^{-1}) \) | \( k_d(s^{-1}) \) | \( K_D(M) \) |
|--------------------------------|--------------------------|-------------------|--------------|
| 1 ECD1-nuc                      | 5.39 \( \times 10^4 \)   | 7.66 \( \times 10^{-4} \) | 1.42 \( \times 10^{-8} \) |
| 2 DARC from K562 cells          | 2.076 \( \times 10^6 \)   | 5.472 \( \times 10^{-4} \) | 2.626 \( \times 10^{-10} \) |
| 3 DARC from K562 cells after deglycosylation | 9.754 \( \times 10^6 \)   | 2.757 \( \times 10^{-4} \) | 2.826 \( \times 10^{-11} \) |
| 4 ECD1-nuc after phosphorylation | 1.692 \( \times 10^6 \)   | 7.769 \( \times 10^{-4} \) | 4.592 \( \times 10^{-10} \) |

doi:10.1371/journal.pone.0116472.t002
When Trp-26 was substituted with Phe, the binding was reduced five times, but was not completely abolished (Table 3, row 9), suggesting that an aromatic ring at position 26 may to some extent substitute for the indole present in Trp residues. The flow cytometry experiments also confirmed the role of Tyr-30 and tyrosine sulfation, since substitution of Tyr-30 by Ala caused a decrease of 2C3 MAb binding (Table 3, row 2), while substitution of Tyr-30 by Glu increased the binding to some extent (Table 3, row 3). These data were consistent in ELISA and SPR experiments.

To evaluate the influence of glycosylation on 2C3 MAb binding, the recombinant DARC purified from K562 cells (S1 Fig.) was immobilized on the chip and the oligosaccharides were removed by PNG-ase F. The *Sambucus nigra* agglutinin (SNA) lectin which recognizes terminal residues of sialic acids was used to evaluate if the reaction was complete. It was shown before that terminal sialic acid residues are present in oligosaccharide chains of DARC [20, 21]. We found that binding of SNA was largely decreased after treatment of the immobilized protein by PNG-ase F (data not shown), while affinity of the antibody increased (Table 2, rows 2 and 3), suggesting that oligosaccharide chains, which are not part of the epitope, may in fact hinder partially access of the antibody to the peptide backbone.

In contrast, when binding of 2C3 MAb was evaluated using K562 cells transfected with the vector encoding DARC without glycosylation site at Asn-27, no difference in binding was observed between such construct and consensus DARC (Table 3, compare rows 1 and 4). It may be argued that the affinity increase observed in the SPR experiment could not be detected when the less sensitive flow cytometry technique was employed.

Cloning and expression of 2C3 Fab

The 2C3 Fab fragment was cloned and expressed in *E. coli* and purified (S2 Fig.), and evaluated by ELISA and surface plasmon resonance (Table 4). The accession numbers for variable regions of the light and heavy chains are KP179400 and KP179401, respectively.

It was found that the affinity of 2C3 Fab is about two times lower (statistically significant at concentration 20 nM, \( P < 0.05 \), two-tailed Mann-Whitney U test) than the affinity of 2C3 parent antibody (Fig. 2).

The lower affinity is an asset to STD-NMR studies as discussed below.
STD-NMR studies of interaction between 2C3 Fab and its epitope

The role of structural elements within the peptide that contribute to the Fab-binding epitope were investigated by STD NMR [34, 35] using synthetic peptide DFEDVWNSSYG. We found that the enhanced resonances in the STD NMR spectrum belonged predominantly to protons ε3, ζ2, ζ3, η2 and δ1 of the Trp-26 residue (the enhanced resonances are indicated in Fig. 3). Minor enhancements of the resonances of protons ε1/ε2 and δ1/δ2 of the Phe residue were also observed. The enhanced resonance at ~7.12 ppm could not be assigned unequivocally as protons δ1/δ2 of Tyr residue and proton ζ3 of the Trp were not resolved. Minute enhancements of Tyr protons ε1/ε2 were only noticeable when the on-resonance irradiation frequency was set to 9.5 ppm. Additionally, no substantial enhancements were observed in the region of signals from aliphatic side chains. Interestingly, no enhancements of valine (V) CH3 proton resonance were observed. No enhancement of Val-35 STD signal gives evidence of either the lack of binding or too strong binding and low off rate value or insufficient energy transfer from the protein to this residue due to the unfavorable conformation causing the distance too large for STD signal observation. The STD-NMR experiments confirmed that the immunodominant

|       | ka      | kd      | KD       |
|-------|---------|---------|----------|
| 2C3 Fab | 0.998 × 10⁴ | 9.259 × 10⁴ | 9.305 × 10⁻⁸ |
| 2C3 MAb | 4.62 × 10⁴   | 5.11 × 10⁻⁵ | 1.1 × 10⁻⁹ |

doi:10.1371/journal.pone.0116472.t004

STD-NMR studies of interaction between 2C3 Fab and its epitope

The binding of 2C3 MAb (squares) and 2C3 Fab (circles) was evaluated by ELISA as described in Materials and Methods. The difference in binding is statistically significant at concentration 20 nM (P < 0.05, two-tailed Mann-Whitney U test).

doi:10.1371/journal.pone.0116472.g002
epitope recognized by the Fab is limited predominantly to the Trp residue within the peptide. However, the intensities of the enhanced signals observed in the STD NMR spectrum were low. This could be explained by the tight binding of the Fab to the peptide and presumably small off rates (k_{off}) resulting in an inefficient saturation transfer to ligand.

**Modeling of the peptide structure**

In order to figure out why Val-25, which seemed to be indispensable part of the epitope, does not interact directly with the antibody, structures of peptide DFEDYWNSSYG and peptides with Val-25 substituted with other amino acids were evaluated using PEP-FOLD webserver. The most relevant peptides conformations, i.e. associated with lowest sOPEP energies, were
selected for further analysis (S1 Table). Based on a structural alphabet approach [36] and a coarse-grained protein force field for structure fold prediction [37], PEP-FOLD proposed various local peptide structures near to their native conformations. Experimental 3D structure of the Gln-19—Tyr-30 portion of DARC was extracted from published DARC-PvDBP complex (PDB 4NUU) [5], it was compared with peptide models obtained in our in silico studies. Structures of peptide models based on sOPEP energy show striking similarities to the experimental one. The degree of similarity of experimental and predicted structures was evaluated by computing backbone RMSD (root-mean square deviation): RMSD values range between 0.69 and 0.83 Å. This strongly supports the pertinence of our modeling approach. Moreover the distances between Trp-26, Tyr-30 and Phe-22 in the peptide DFED[V, I, E, L]WNSSYG, were calculated (S1 Table), and show some significant differences underlining the influence on the peptide structure of the amino acid present at position 25.

Indeed the peptide models shown in Fig. 4 and S1 Table demonstrate how Val-25 substitution alter the epitope 3D structure and may influence binding to 2C3 MAb. The distances between residues involved in binding of the antibody are slightly different for those models. The substitution of Val (HI = 4.2) with Ile residue, an amino acid with higher hydrophobic index (HI = 4.5) but longer aliphatic chain does not cause significant structure deformation (0.45 to 0.66 Å more than for Val-25 model, S1 Table). Val-25 substitutions with amino acids such as Ala (HI = 1.8) and Leu (HI = 3.8) change the epitope conformation and extend the distances between amino acids involved in the interaction with antibody 2C3. Leu and Ala substitutions imply 0.09-2.019 and 0.15-2.29 Å shifts compared to Val-25 model, respectively.

Moreover, conformational changes of particular amino acids side-chains, which may impact the epitope formation, were also observed (Fig. 4).

Discussion
As the role of monoclonal antibodies in biology and medicine becomes increasingly important, so does the precise evaluation of the epitopes being recognized by the antibodies. There are
many methods allowing to precisely define amino acid residues that are involved in antibody binding, such as evaluation of binding of the antibody to synthetic peptides (Pepscan analysis) or to recombinant peptides expressed either in bacteria on in eukaryotic cells [26, 38–43]. Recently, saturation transfer difference spectroscopy NMR (STD-NMR) has been increasingly used in epitope-antibody studies, although majority of antibodies evaluated by this method bind carbohydrate antigens [44–46]. In this paper we used a variety of methods, including STD-NMR, to precisely define the peptidic epitope of 2C3 MAb.

The murine monoclonal antibody 2C3 is one of the widely used anti-DARC antibodies. Since its binding interferes with some of the DARC functions such as chemokine binding [24], and invasion of erythrocytes by *P. vivax* merozoites [27], precise knowledge on the recognized epitope seems necessary for its potential applications.

Our results unequivocally re-establish the important role of Trp-26 in DARC recognition by the antibody confirming the data published by Wasiowska [24], as it cannot be replaced by any residue without significant decrease of affinity. However, when Trp-26 is substituted by Phe, binding of 2C3 MAb was not abolished completely, suggesting that the aromatic ring may substitute the indole ring. The role of Tyr-30 has also been clarified: it can be substituted with Ala without strongly decreasing the affinity, which stands in contrast with the results published previously [25]. However, our results suggest that sulfation of Tyr-30 causes an increase of the antibody affinity to some extent. Moreover, when ECD1-nuc construct contained both Phe at position 26 and Glu at position 30, the binding of the antibody increased in comparison to the construct with Phe-26 only. Finally, phosphorylation of Tyr-30 on ECD1-nuc used to mimic sulfation, caused and increase of the 2C3 MAb affinity. These data suggest that the negative charge at Tyr-30 possibly increases the antibody—antigen interaction.

Results of STD-NMR presented in this paper support these conclusions. Saturation transfer difference (STD)-NMR technique can be used to identify the proton resonances of a ligand in close contact with the binding protein: side chains within the ligand that interact most strongly show a strong STD effect. There are several reports in which interaction between receptor and peptide ligand [47, 48] or monoclonal antibody and oligosaccharide epitope [44, 49, 50] is evaluated. However, only a handful of papers exist in which interactions between monoclonal antibody and peptide epitope are studied [45, 51]. The main limitation of the technique is the dissociation constant, which must be in range 10⁻³ to 10⁻⁸M [34, 35]. In the case of 2C3 antibody, the *K*₀ is too low (1.1 × 10⁻⁸M) so we decided to use Fab fragment, whose *K*₀ is 9.3 × 10⁻⁸M. Evaluation of interaction of peptide DFEDVWNSSYG with 2C3 Fab by STD-NMR technique confirmed that Trp-26 plays a major role in the antibody binding. It was found that interaction between Phe-22 and the antibody is weaker, whereas Tyr-30 shows the weakest interaction. Thus, only aromatic residues have been identified by STD-NMR, while other residues (such as Val-25) cannot be resolved in the spectrum. The Val-25 seems to play an important role in the epitope, because it was shown before that it cannot be replaced by another residue (with exception of Ile) without losing ability to bind 2C3 antibody [24].

In order to explain the possible role of Val-25 in the epitope, we used *in silico* modeling of DFEDVWNSSYG peptide, and of peptides with Val-25 substituted by Ile, Leu or Ala. Our models suggest that Val-25 substitution by Leu or Ala implies changes in peptide conformation and moves slightly the hydrophobic amino acids away from each other. However, it must be noted that the Ile side chain (which is longer than Val) might have an influence on epitope formation. The modeling suggests that Val-25 might cramp with other hydrophobic amino acids and control the polypeptide structure. Thus, substitution of Val-25 with other amino acid residues with different hydrophobicity may alter this structure by rendering Tyr-30 further from the epitope (S4B Fig.). Modeling studies showed that the essential residues forming the epitope (Phe-22, Trp-26 and Tyr-30) are exposed one face of the helix while Val-25 located on the
opposite face of the helix might not participate to contacts with antibody. We hypothesize though that Ile, which has a longer side chain than Val, might in some way make some steric hindrance to the antibody epitope interaction since modeling suggests that it does not modify grossly epitope structure. In summary, the peptide structure of the epitope seems to be compact, with aromatic residues forming a single block on one side of an helix (Fig. 4).

Taken together, our results confirm that the minimal DARC epitope recognized by 2C3 MAb is linear and consists of five amino acids: 22FEDVW26, but there is a certain interaction between Tyr-30 and the antibody as shown by STD-NMR. In addition, we found that when Tyr-30 was substituted by a residue with a negative charge, such as glutamic acid, binding of 2C3 antibody was slightly increased despite other crucial amino acid substitutions, suggesting that the negative charge at amino acid 30 may play an auxiliary role in recognition. In vivo, the negative charge is provided by the Tyr-30 due to its sulfation.

Our results were obtained using a set of methods that are applicable for establishing epitope of any monoclonal antibody. We have demonstrated that using eukaryotic and prokaryotic expressed constructs, any doubts regarding antibody specificity can be clarified. We also showed usefulness of SPR analysis to demonstrate influence of post-translational modifications by carrying out enzymatic reactions “in situ” on the protein previously immobilized on the chip. In addition, we showed that STD-NMR technique can be used to evaluate binding of an antibody to a peptide, which to our knowledge is one of the first experiments of this kind described in literature.

Supporting Information

S1 Fig. Analysis of immunoaffinity chromatography on immobilized anti DARC VHH. Lanes 1: molecular weight standard (molecular weight of bands is noted on the left), supplemented with purified anti DARC nanobody. Lane 2 and 4 20 μl of fractions collected during wash of the column with Tris saline 0.3% C12E8 buffer. Lane 3 and 5: 20 μl of material eluted from the column with DFEDVW peptide. Lane 6 contains 20 μl of material eluted from column with 0.1 M glycine buffer. Lanes 1-3: silver nitrate staining, lanes 4-5: Western blotting with 2C3 MAb.

S2 Fig. SDS-PAGE analysis of purified soluble 2C3 Fab. Lane 1: molecular weight standards. The purified protein (4 μg) were loaded onto 10% polyacrylamide gel in the absence (A) or presence (B) of 2-mercaptoethanol, and visualized with Coomasie Brilliant Blue.

S3 Fig. Inhibition of binding of 2C3 MAb to ECD1-nuc constructs evaluated by ELISA. 2C3 MAb was incubated with ECD1-nuc constructs in different concentration as described in Materials and Methods, and subsequent binding to ECD1-nuc—covered plates was evaluated by ELISA.

S4 Fig. The 3-D structure of DARC bound to PvDBP (PDB 4NUU) as visualized by Chime-ra software (A) and compared to peptide models obtained in in silico studies (B). DBP-RII molecules are shown in blue and green, DARC molecule is shown in red [5]. DARC was enlarged on the right and the crucial residues were highlighted using different colors: Phe-22 (purple), Val-25 (red), Trp-26 (green), Tyr-30 (blue). Q19-Y30 peptide from DARC (red) was enlarged and compared to peptide models (black) obtained in in silico studies: (a) DFEDVWNSSSYG (RMSD = 0,75), (b) DFEDSWNSSYG (RMSD = 0,83), (c) DFEDLWNSSYG (RMSD = 0,70), and (d) DFEDAWNSSYG (RMSD = 0,67); B. 3-D structure of peptides...
obtained in modelling studies. Amino acids exposed on the same face of the helix, involved in antibody binding, are shown in purple (Phe-22), green (Trp-26) and blue (Tyr-30). Residues located on the other surface of the helix (Val-25, Ile-25, Leu-25 and Ala-25) are shown in red.

(S1 Table. The representative structure models of the native DFEDVWNSSYG peptide and peptides with Val-25 substitution. Three-dimensional structure of sOPEP energy model was calculated by PEP-FOLD server. The distances between Trp-26, Tyr-30 and Phe-22 were calculated by Chimera software. The colors show hydrophobicity of surface amino acids in the Kyte-Doolittle scale (blue for the most hydrophilic to white at 0.0 and to orange red for the most hydrophobic). Peptide models underline the influence of hydrophobic/hydrophilic amino acids on position 25 (Val, Ile, Leu, Ala) on peptide folding.

(ACKNOWLEDGMENTS)

We thank Corine Sandström of the Department of Chemistry, Swedish University of Agricultural Sciences, Uppsala BioCenter, for her assistance regarding the STD NMR experiments.

(AUTHOR CONTRIBUTIONS)

Conceived and designed the experiments: DS JL YCA KW OB MC. Performed the experiments: DS CH AB RK AJ SC ADB JL WJ TN MG OB MC. Analyzed the data: DS WJ TN OB MC. Contributed reagents/materials/analysis tools: KW YCA OB MC. Wrote the paper: DS JL OB MC. Performed modeling of the peptide: AJ.

(REFERENCES)

1. Tournamille C, Colin Y, Cartron JP, Le Van Kim C (1995) Disruption of a GATA motif in the Duffy gene promoter abolishes erythroid gene expression in Duffy-negative individuals. Nat Genet 10: 224–228. doi: 10.1038/ng0695-224
2. Miller LH, Mason SJ, Clyde DF, McGinniss MH (1976) The resistance factor to Plasmodium vivax in blacks. The Duffy-blood-group genotype, FyFy. N Engl J Med 295: 302–304. doi: 10.1056/NEJM197608052950602 PMID: 778616
3. Mercereau-Puijalon O, Menard D (2010) Plasmodium vivax and the Duffy antigen: a paradigm revisited. Transfus Clin Biol 17: 176–183. doi: 10.1016/j.trac.2010.06.005 PMID: 20655790
4. Batchelor JD, Zahm JA, Tolia NH (2011) Dimerization of Plasmodium vivax DBP is induced upon receptor binding and drives recognition of DARC. Nat Struct Mol Biol. doi: 10.1038/nsmb.2088 PMID: 21743458
5. Batchelor JD, Maiped BM, Omattage NS, DeKoster GT, Henzler-Wildman KA, et al. (2014) Red blood cell invasion by Plasmodium vivax: structural basis for DBP engagement of DARC. PLoS Pathog 10: e1003869. doi: 10.1371/journal.ppat.1003869 PMID: 24415938
6. Hadley TJ, Peiper SC (1997) From malaria to chemokine receptor: the emerging physiologic role of the Duffy blood group antigen. Blood 89: 3077–3091. PMID: 9129009
7. Pogo AO, Chaudhuri A (2000) The Duffy protein: a malarial and chemokine receptor. Semin Hematol 37: 122–129. doi: 10.1016/S0037-1963(00)90037-4 PMID: 10791881
8. Shen H, Schuster R, Stringer CF, Waltz SE, Lentsch AB (2006) The Duffy antigen/receptor for chemokines (DARC) regulates prostate tumor growth. FASEB J 20: 59–64. doi: 10.1096/fj.05-4764com PMID: 16934288
9. Lentsch AB (2002) The Duffy antigen/receptor for chemokines (DARC) and prostate cancer. A role as clear as black and white? FASEB J 16: 1093–1095. doi: 10.1096/fj.02-0066hyp PMID: 12087701
10. Sun G, Wang Y, Zhu Y, Huang C, Ji Q (2011) Duffy antigen receptor for chemokines in laryngeal squamous cell carcinoma as a negative regulator. Acta Otolaryngol 131: 197–203. doi: 10.3109/00016489.2010.516012 PMID: 21105847
11. Gardner L, Patterson AM, Ashton BA, Stone MA, Middleton J (2004) The human Duffy antigen binds selected inflammatory but not homeostatic chemokines. Biochem Biophys Res Commun 321: 306–312. doi: 10.1016/j.bbrc.2004.06.146 PMID: 15358176

12. Walton RT, Rowland-Jones SL (2008) HIV and chemokine binding to red blood cells—DARC matters. Cell Host Microbe 4: 3–5. doi: 10.1016/j.chom.2008.06.006 PMID: 18621004

13. Kulkarni H, Marconi VC, He W, Landrum ML, Okulicz JF, et al. (2009) The Duffy-null state is associated with a survival advantage in leukemic HIV-infected persons of African ancestry. Blood 114: 2783–2792. doi: 10.1182/blood-2009-04-215186 PMID: 19620399

14. Lachgar A, Jaureguiberry G, Le Buenac H, Bizzini B, Zagury JF, et al. (1998) Binding of HIV-1 to RBCs involves the Duffy antigen receptors for chemokines (DARC). Biomed Pharmacother 52: 436–439. doi: 10.1016/S0753-3322(99)80021-3 PMID: 9921412

15. Horuk R, Chitnis CE, Darbonne WC, Colby TJ, Rybicki A, et al. (1993) A receptor for the malarial parasite Plasmodium vivax: the erythrocyte chemokine receptor. Science 261: 1182–1184. doi: 10.1126/science.7689250 PMID: 7689250

16. Rot A (2005) Contribution of Duffy antigen to chemokine function. Cytokine Growth Factor Rev 16: 687–694. doi: 10.1016/j.cytoifr.2005.05.011 PMID: 16054417

17. Ribbs R, Graham G, Rot A (2003) Chemokines on the move: control by the chemokine “interceptors” Duffy blood group antigen and D6. Semin Immunol 15: 287–294. doi: 10.1016/j.smim.2003.08.006 PMID: 15001178

18. Bachelerie F, Ben-Baruch A, Burkhardt AM, Combadiere C, Farber JM, et al. (2014) International Union of Basic and Clinical Pharmacology. [corrected]. LXXXIX. Update on the extended family of chemokine receptors and introducing a new nomenclature for atypical chemokine receptors. Pharmacol Rev 66: 1–79. doi: 10.1124/pr.113.007724 PMID: 24218476

19. Czerwinski M, Kern J, Grodecka M, Paprocka M, Krop-Watorek A, et al. (2007) Mutational analysis of the N-glycosylation sites of Duffy antigen/receptor for chemokines. Biochem Biophys Res Commun 356: 816–821. doi: 10.1016/j.bbrc.2007.03.054 PMID: 17382291

20. Grodecka M, Czerwinski M, Duk M, Lisowska E, Wasniowska K (2010) Analysis of recombinant Duffy protein-linked N-glycans using lectins and glycosidases. Acta Biochim Pol 57: 49–53. PMID: 20234884

21. Grodecka M, Bertrand O, Karolak E, Lisowski M, Wasniowska K (2012) One-step immunopurification and lectinochemical characterization of the Duffy atypical chemokine receptor from human erythrocytes. Glycoconj J 29: 93–105. doi: 10.1007/s10719-011-9367-9 PMID: 22246380

22. Choe H, Moore MJ, Owens CM, Wright PL, Vasilieva N, et al. (2005) Sulphated tyrosines mediate association of chemokines and Plasmodium vivax Duffy binding protein with the Duffy antigen/receptor for chemokines (DARC). Mol Microbiol 55: 1413–1422. doi: 10.1111/j.1365-2958.2004.04478.x PMID: 15720550

23. Wasniowska K, Blanchard D, Janvier D, Wang ZX, Peiper SC, et al. (1996) Identification of the Fy6 epitope recognized by two monoclonal antibodies in the N-terminal extracellular portion of the Duffy antigen receptor for chemokines. Mol Immunol 33: 917–923. doi:10.1016/S0161-5890(96)00056-9 PMID: 8960115

24. Wasniowska K, Petit-LeRoux Y, Tournamille C, Le van Kim C, Cartron JP, et al. (2002) Structural characterization of the epitope recognized by the new anti-Fy6 monoclonal antibody NaM 185. Cell Mol Life Sci 69: 205–211. doi: 10.1007/s00018-002-0073-x PMID: 12164140

25. Tournamille C, Filipe A, Wasniowska K, Gane P, Lisowska E, et al. (2003) Structure-function analysis of the extracellular domains of the Duffy antigen/receptor for chemokines: characterization of antibody and chemokine binding sites. Br J Haematol 122: 1014–1023. doi: 10.1046/j.1365-2141.2003.04533.x PMID: 12956774

26. Smolarek D, Hattab C, Hassanzadeh-Ghassabeh G, Cochet S, Gutierrez C, et al. (2010) A recombinant dromedary antibody fragment (VHH or nanobody) directed against human Duffy antigen receptor for chemokines. Cell Mol Life Sci 67: 3371–3387. doi: 10.1007/s00018-010-0387-6 PMID: 20458517

27. Russell B, Suwanarusk R, Borlon C, Costa FT, Chu CS, et al. (2011) A reliable ex vivo invasion assay of human reticulocytes by Plasmodium vivax. Blood 118: e74–81. doi: 10.1182/blood-2011-04-348748 PMID: 21768300

28. Czerwinski M, Siemaszko D, Siegel DL, Spitalnik SL (1998) Only selected light chains combine with a survival advantage in leukopenic HIV-infected persons of African ancestry. Blood 114: 2783–2792. doi: 10.1182/blood-2009-04-215186 PMID: 19620399

29. Czerwinski M, Krop-Watorek A, Lisowska E, Spitalnik SL (2002) Construction of dimeric F(ab) useful in blood group serology. Transfusion 42: 257–264. doi: 10.1046/j.1537-2995.2002.00018.x PMID: 11896344
30. Carbonnet F, Blanchard D, Hattab C, Cochet S, Petit-Leroux Y, et al. (2000) A murine monoclonal antibody against Kx protein which reacts also with beta-spectrin. Transfus Med 10: 145–154. doi: 10.1046/j.1365-3148.2000.00245.x PMID: 10849386

31. Goddard TD, Kneller G (2001) SPARKY 3. University of California, San Francisco.

32. Thevenet P, Shen Y, Maupetit J, Guyon F, Derreumaux P, et al. (2012) PEP-FOLD: an updated de novo structure prediction server for both linear and disulfide bonded cyclic peptides. Nucleic Acids Res 40: W288–293. doi: 10.1093/nar/gks419 PMID: 22581768

33. Pettersen EF, Goddard TD, Huang CC, Couch GS, Greenblatt DM, et al. (2004) UCSF Chimera—a visualization system for exploratory research and analysis. J Comput Chem 25: 1605–1612. doi: 10.1002/jcc.20084 PMID: 15264254

34. Mayer M, Meyer B (1999) Characterization of ligand binding by saturation transfer difference NMR spectroscopy. Angewandte Chemie-International Edition 38: 1784–1788. doi: 10.1002/(SICI)1521-3773(19990614)38:12<1784::AID-ANIE1784>3.3.CO;2-H

35. Meyer B, Peters T (2003) NMR Spectroscopy techniques for screening and identifying ligand binding to protein receptors. Angewandte Chemie-International Edition 42: 864–890. doi: 10.1002/anie.200390233

36. Offmann B, Tyagi M, de Brevern AG (2007) Local protein structures. Current Bioinformatics 2: 165–202. doi: 10.2174/157489030771862105

37. Maupetit J, Tuffery P, Derreumaux P (2007) A coarse-grained protein force field for folding and structure prediction. Proteins 69: 394–408. doi: 10.1002/prot.21505 PMID: 17660832

38. Chen HW, Zhang Z, Huber E, Chao CC, Wang H, et al. (2009) Identification of cross-reactive epitopes on the conserved 47-kilodalton antigen of Orientia tsutsugamushi and human serine protease. Infect Immun 77: 2311–2319. doi: 10.1128/IAI.01298-08 PMID: 19289508

39. Chen Y, Zhang J, Qiao C, Wang J, Yang H, et al. (2012) Identification of a linear epitope on the hagemagglutinin protein of pandemic A/H1N1 2009 influenza virus using monoclonal antibodies. Arch Virol 159: 1413–1419. doi: 10.1007/s00705-013-1955-5 PMID: 24385157

40. Yermakova A, Vance DJ, Mantis NJ (2012) Sub-domains of ricin’s B subunit as targets of toxin neutralizing and non-neutralizing monoclonal antibodies. PLoS One 7: e44317. doi: 10.1371/journal.pone.0044317 PMID: 22984492

41. Huang L, Lu Y, Wei Y, Guo L, Liu C (2012) Identification of three new type-specific antigen epitopes in the capsid protein of porcine circovirus type 1. Arch Virol 157: 1339–1344. doi: 10.1007/s00705-012-1268-0 PMID: 22437253

42. Bushnell RV, Tobin JK, Long J, Schultz-Cherry S, Chaudhuri AR, et al. (2010) Serological characterization of guinea pigs infected with H3N2 human influenza or immunized with hemagglutinin protein. Virol J 7; 200. doi: 10.1186/1423-0410-7-200 PMID: 20738549

43. Rasamoeilolo M, Czerwinski M, Bruneau V, Lisowska E, Blanchard D (1997) Fine characterization of a series of new monoclonal antibodies directed against glycoporphin A. Vox Sang 72: 185–191. doi: 10.1046/j.1423-0410.1997.7230185.x PMID: 9145941

44. Plum M, Michel Y, Wallach T, Raiber T, Blank S, et al. (2011) Close-up of the immunogenic alpha1,3-galactose epitope as defined by a monoclonal chimeric immunoglobulin E and human serum using saturation transfer difference (STD) NMR. J Biol Chem 286: 43103–43111. doi: 10.1074/jbc.M609816200 PMID: 17192263

45. Szczepina MG, Bleile DW, Pinto BM (2011) Investigation of the binding of a carbohydrate-mimetic peptide to its complementary anticyanogenic antibody by STD-NMR spectroscopy and molecular dynamics simulations. Chemistry 17: 11446–11455. doi: 10.1002/chem.201100222 PMID: 21953925

46. Enriquez-Navas PM, Chiodo F, Marradi M, Angulo J, Penades S (2012) STD NMR study of the interactions between antibody 2G12 and synthetic oligomannosides that mimic selected branches of gp120 glycans. Chembiochem 13: 1357–1365. doi: 10.1002/cbic.201200119 PMID: 22628288

47. Mesleh MF, Shirley WA, Heise CE, Ling N, Maki RA, et al. (2007) NMR structural characterization of a minimal peptide antagonist bound to the extracellular domain of the corticotropin-releasing factor1 receptor. J Biol Chem 282: 6338–6346. doi: 10.1074/jbc.M609816200 PMID: 17192263

48. Pons J, Evrard-Todeschi N, Bertho G, Gharbi-Benarous J, Tanchou V, et al. (2008) Transfer-NMR and docking studies identify the binding of the peptide derived from activating transcription factor 4 to protein ubiquitin ligase beta-TrCP. Competition STD-NMR with beta-catenin. Biochemistry 47: 14–29. doi: 10.1021/bi7014212 PMID: 18052253

49. Niedziela T, Letowska I, Lukasiewicz J, Kaszowska M, Czarnecka A, et al. (2005) Epitope of the vaccine-type Bordetella pertussis strain 186 lipooligosaccharide and antiendotoxin activity of antibodies directed against the terminal pentasaccharide-tetanus toxoid conjugate. Infect Immun 73: 7381–7389. doi: 10.1128/IAI.73.11.7381-7389.2005 PMID: 16239537
50. Enriquez-Navas PM, Marradi M, Padro D, Angulo J, Penades S (2011) A solution NMR study of the interactions of oligomannosides and the anti-HIV-1 2G12 antibody reveals distinct binding modes for branched ligands. Chemistry 17: 1547–1560. doi:10.1002/chem.201002519 PMID: 21268157

51. Johnson MA, Pinto BM (2004) Saturation-transfer difference NMR studies for the epitope mapping of a carbohydrate-mimetic peptide recognized by an anti-carbohydrate antibody. Bioorg Med Chem 12: 295–300. doi:10.1016/j.bmc.2003.09.041 PMID: 14697795