

Bitter Sweet Symphony: the impact of sugars on autoimmunity

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Surface immunoglobulin variable domain glycosylation affects autoantigen binding and acts as threshold for human autoreactive B-cell activation

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Abstract

The hallmarking autoantibodies in Rheumatoid Arthritis (RA) are characterized by variable domain glycans (VDGs). Their abundant occurrence results from the selective introduction of N-linked glycosylation sites during somatic hypermutation, and their presence is predictive for disease development. However, the functional consequences of VDGs on autoreactive B cells remain elusive. Combining crystallography, glycobiology, and functional B-cell assays allowed us to dissect key characteristics of VDGs on human B-cell biology. Crystal structures showed that VDGs are positioned in the vicinity of the antigen-binding pocket, and dynamic modeling combined with binding assays elucidated their impact on binding. We found that VDG-expressing B-cell receptors stay longer on the B-cell surface and that VDGs enhance B-cell activation. These results provide a rationale on how the acquisition of VDGs might contribute to the breach of tolerance of autoreactive B cells in a major human autoimmune disease.

Teaser

The molecular underpinnings and the functional impact of variable domain glycans on human autoreactive B cells.

Introduction

A pathogenic role of B cells in autoimmunity is evidenced by the effective treatment of multiple autoimmune diseases, such as rheumatoid arthritis (RA), multiple sclerosis, antineutrophil cytoplasmic antibody (ANCA)-associated vasculitis, and systemic lupus erythematosus (SLE) with B-cell targeted therapies¹. Together with the observation that disease-specific autoantibody responses often characterize autoimmune diseases, these findings point to the notion that autoreactive B cells are involved in disease pathogenesis. Autoreactive B cells could contribute to disease via diverse mechanisms, such as antigen presentation to T cells, secretion of cytokines, or the production of pathogenic antibodies². To mediate these functions, an autoreactive B-cell response needs to be induced, which involves the breach of tolerance mechanisms that generally prevent the activation of pathogenic B cells. However, the mechanisms leading to the induction and survival of autoreactive B cells that contribute to autoimmune diseases remain unclear. In mouse models, it has been shown that B cells harboring autoreactive B-cell receptors (BCRs) are regulated in the bone marrow and peripheral tissues by clonal deletion, anergy induction (short-lived inactivated cells), or receptor editing³⁻⁷. This multistep process eliminates high-affinity and potentially autoreactive clones⁸. During immune responses against non-self or modified proteins carried by unwanted intruders, some initially autoreactive B cells might somatically mutate to be more efficient defense cells, thereby potentially losing affinity to self-antigens^{9,10}. This might include a role for variable domain glycans (VDGs) that could regulate self/non- self discrimination of B cells as has been described in a B-cell receptor-transgenic mouse model¹¹. Notably, the survival of autoreactive B cells involves costimulatory receptors and negative regulators, such as the sialic acid-binding lectin CD22, which will, together with the BCR, determine the outcome of the activation, the timing of T-cell help and the accessibility of antigens^{12,13}. At present, although we have some alimpses through the study of autoreactive B-cell biology in mice, the processes controlling human autoreactive B cells are largely unexplored.

The hallmarking disease-specific autoantibodies in RA, a common human autoimmune disease affecting synovial joints and cartilage, are anti-citrullinated protein antibodies (ACPAs)¹⁴. ACPAs specifically recognize a citrulline side chain and show limited interactions with the side-chains of surrounding amino acids on the protein surface, explaining their promiscuity in their protein or peptide specificity¹⁵. Recently, we have found that more than 90% of ACPA are glycosylated in their variable domains^{16,17}. The VDGs displayed on ACPAs are acquired through the introduction of N-linked glycosylation sites following somatic hypermutation¹⁸. VDGs are mainly complex-type carbohydrates containing a high percentage of sialic acids¹⁶. Intriguingly, the presence of VDGs in ACPA-positive healthy subjects is associated with the transition toward RA¹⁹. This observation, together with the selective introduction of N-linked glycosylation sites and the finding that this introduction has taken place in more than 90% of ACPA in patients with

RA^{16,18}, underlines the importance of this glycan tag and suggests that VDGs play a regulatory role for ACPA-expressing B cells. Intriguingly, variable domain glycosylation of antibodies is a characteristic not only of RA but also for some other human responses, such as ANCA-associated vasculitis, Sjögren's syndrome, anti-Hinge, and antidrug antibodies²⁰⁻²².

By combining crystallography, glycobiology, and antigen-binding studies, we show that ACPA VDGs can mask binding toward potential self-antigens. Functionally, we demonstrate that BCR VDGs enhances B-cell activation and decreases BCR downmodulation. This occurs independently of the negative regulator CD22. Together, the data demonstrate an important role of VDGs in changing the threshold of B cells to respond to self-antigens and to be activated. The generation of B cells with receptors modified by VDGs could lead to a loss of control of the self-reactive response and may play a role in the pathogenesis of RA.

Results

Generation of patient-derived monoclonal antibodies with and without VDGs

To investigate the putative effects of autoantibody VDGs on antigen binding, we generated six monoclonal antibodies (mAbs) based on full-length immunoglobulin G (lgG) BCR sequences isolated from ACPA-positive patients with RA (Figure 1A and Table 1)^{23,24}.

Figure 1. Generation of patient-derived IgG1 monoclonal anti-citrullinated protein (auto)antibodies (ACPAs) with (WT) and without (NG) variable domain glycans (VDGs). (A) Schematic representation of the generated ACPA IgG (7E4, 2E4, 1F2, 2G9, 3F3, and 2D11) carrying various amounts of N-linked glycans in the variable regions of their HC (turquoise) and LC (light gray). (B) Size shift between WT and NG ACPA under non-reduced (IgG) and reduced (HC and LC) conditions on a 4 to 15% gradient SDS protein gel (Bio Rad). The size was determined using the PageRulerTM Plus Prestained Protein Ladder (Thermo Fisher Scientific). (C) Size shift between WT and NG ACPA IgG identified via analytical size exclusion chromatography (SEC). ACPA IgG⁺ fractions were determined via the absorbance at 280 nm. (D) Schematic depiction of the four ACPA variants (WT, NG, WT+neu and NG+neu) analyzed: variable domain glycosylated (VDG⁺), Fc glycosylated (FcG⁺), sialylated (SA⁺) and non sialylated (SA⁺) after neuraminidase (neu) treatment. (E) MALDI-TOF MS analysis of N-linked variable domain and Fc glycans from WT, WT+neu, NG, and NG+neu ACPA IgG 7E4 and 3F3. Glycan structures of the most abundant N-linked glycan peaks are depicted. Blue square, *N*-acetylglucosamine (N); green circle, mannose/ hexose (H); yellow circle, galactose/ hexose (H); red triangle, fucose (F); purple diamond, α 2,6-linked *N*-acetylneuraminic (sialic) acid (S). a.u., arbitrary units. **>**



We produced the IgG1 mAbs as wild-type (WT) variants including their respective N-linked glycosylation sites, which are required for the acquisition of VDGs. Next, non-glycosylated (NG) mAbs were generated by mutating the N-linked glycosylation sites in the variable domains back to the respective germline sequence (Table 1). Both mAb variants express the typical asparagine (N) 297-linked Fc glycans. The recombinant proteins were expressed in human embryonic kidney cells with transiently overexpressed glycosyltransferases (i.e., for glycoengineering)²⁵ to achieve glyco-variants similar to those found on ACPA IgG isolated from patients with RA¹⁶. We analyzed the mAb integrity and the occupancy of the N-linked glycosylation sites by SDS-polyacrylamide gel electrophoresis (PAGE) and size exclusion chromatography (SEC). Our results confirmed an increased apparent molecular weight of ACPA IgG due to the presence of VDGs¹⁷, as evidenced by a shift of the WT versus the NG variants in gel electrophoresis and SEC (Figures 1, B and C). Furthermore, our data indicated that all N-linked glycosylation sites were fully occupied by VDGs, except for the glycosylation sites in the light chain (LC) of the mAbs 1F2 and 2G9, which appeared to be partially occupied (Figure 1B). We analyzed the variable domain- and Fc-glycan profile of the mAb variants (Figure 1D) in detail using mass spectrometry (MS) [matrix-

assisted laser desorption/ ionization-time-of-flight (MALDI-TOF)] (Figure 1E and Figure S1). The exemplarily illustrated results of 7E4 and 3F3 (Figure 1E) showed a highly sialylated glycan profile for the WT variant with a dominant complex type, bisected, and α -2,6-linked sialic acid comprising biantennary glycan peak H5N5F1S2 (*m/z=2651*) (Figure 1E), similar to the most prominent VDGs found on ACPA IgG directly isolated from patients with RA¹⁶. As we wished to determine the effect of negatively charged sialic acids (specifically *N*-acetylneuraminic acids) on antigen-binding functionalities, we treated the WT and NG mAb variants with neuraminidase (+neu). As depicted in figure 1E, this resulted in an effective cleavage of sialic acids. Likewise, the NG variant showed no sialylated glycan peaks but only the three most abundant Fc glycan forms comprising different amounts of galactose (H3N4F1, H4N4F1 and H5N4F1) (Figure 1E). Together, these results demonstrate the successful generation of patient-derived IgG mAb with engineered VDGs presenting the expected size and glycan composition as well as their NG counterparts.

Spatial orientation of VDG and antigen-binding pockets

Crystal structures of two ACPA IgG Fab fragments (3F3_{rab} and 1F2_{rab}) were generated to identify the structural composition of the antigen-binding pockets and the spatial orientation of the VDGs. The ACPA Fab fragments were complexed with distinct citrullinated peptides [3F3_{E3}:citvim 59-74 [Protein Data Bank (PDB): 6YXK] and 1F2_{Fab};cit-CII-C-39 (PDB: 6YXM)] selected on the basis of high reactivities within in vitro binding multiplex assays (as described in¹⁵) (Figure 2A). Additionally, 1F2 $_{\rm Fab}$ was crystallized in an unbound state (PDB: 6YXN) (Figure 2A). The crystallographic data collection and refinement parameters are summarized in supplementary table 3. The results showed that $3F3_{_{Fab}}$ and $1F2_{_{Fab}}$ adopted characteristic Ig folds with well-defined binding grooves indicated by electron densities (Figure 2B). The core binding regions of the citrullinated peptides are well-defined, except for the terminal parts, which extend beyond the binding pocket (Figure 2B). Furthermore, the complexed citrullinated (CIT) peptides are visible in the antigen-binding sites and residues SSAV(CIR)L(CIT)SS of cit-vim 59-74 and LPGQ(CIT) GERG of cit-CII-C-39 could be modelled, while the other peptide residues were invisible (Figure 2C). The buried molecular surface areas of the Fab fragment and its respective peptide upon binding are 503 Å for 1F2_{rab} [178 Å heavy chain (HC) and 325 Å LC] and 622 Å for cit-CII-C-39 as well as 659 Å for 3F3 $_{\rm Fab}$ (323 Å HC and 336 Å LC) and 789 Å for cit-vim 59-74, respectively. The results indicate that the LC, especially for 1F2_{Fab}:cit-CII-C-39, contributed considerably to antigen binding. Superposition of the Apo-form of 1F2 (non-ligated) and its peptide-bound complex gave a backbone atom root mean square deviation (RMSD) of 0.4 Å, indicating no major changes in Ig folding upon antigen binding. When analyzing the structures of the unbound state, we found two deep pockets located within the antigen-binding region (Figure 2B), similar to pockets found in a third ACPA, 7E4_{Fab}, that has been described previously (PDB:5OCY)¹⁵. Noteworthy, these pockets, composed of polar and hydrophobic residues from HC and LC, are sufficient to accommodate a citrulline side-chain (Figure 2C). The flanking residues around the citrulline

were, in both cases, partially embedded in the binding groove, while the terminal residues of the peptides were directed towards the solvent (Figure 2C). The two peptides adopt a β -sheet like conformation atop the complementary determining regions (CDRs). To determine the structural basis required to accommodate citrullinated peptides, we performed a structural comparison between the determined ACPA 3F3_{Fab} and 1F2_{Fab} crystal structures and 7E4_{Fab}. The superposition of the three Fab structures revealed distinct structural features of paratopes formed by the CDRs including different conformations, suggesting heterogenous antigen-binding regions among different ACPAs complexed with various peptides (Figure 2D). Moreover, there was a major difference in the electrostatic potentials that might determine the recognition of flanking residues of citrulline (Figure 2E). Noteworthy, the binding pocket of 3F3_{Fab} is positively charged, different from the negative or neutral antigen-binding pocket characteristics of the other ACPA Fab fragments described (Figure 2E).

Hence, all ACPA Fab fragments exhibit an open-ended polar/hydrophobic binding groove, fitting the binding of the citrulline side chain, which enables them to promiscuously accommodate different peptide variants, while excluding others due to steric repulsion with other amino acid side chains. This and the fact that interactions are solely formed between the terminal nitrogen and oxygen atom of the citrulline side chain could explain the low affinity toward many of their antigens and the broad cross-reactivity toward various antigens and post-translational modifications harbouring these atoms (acetylated and carbamylated antigens)²³.

Next, we characterized the spatial localization of the VDGs as this could provide insights into the potential impact of ACPA VDGs on antigen binding at the molecular level. Although N-linked glycans are highly flexible structures, the starting monosaccharides (core domain) were cocrystallized with the respective antigen-binding region of the generated monoclonal ACPA 7E4_{Fab}:cit-CII-C-48 (PDB: 6ZJG), 3F3_{Fab}:cit-vim 59-74 and 1F2_{Fab}:cit-CII-C-39 (Figure 3A). The data illustrate that the glycans are positioned outside, but in close vicinity to the peptide-binding domain. To examine whether VDGs could potentially reach the antigen-binding pocket and thereby interfere with antigenic interactions, we used the crystals as a basic framework and modeled the complete disialylated glycan compositions on top of these structures. Fab crystal structures with modeled VDGs were subsequently used for molecular dynamics (MD) simulations (Figure 3B and Figure S2A). The number of hydrogen bonds formed during simulation time was analyzed for every Fab domain and the respective HC and LC N-glycans. The results showed that up to six hydrogen-bonds were formed between the LC N-glycan and the mAb structure. The HC glycan formed fewer interactions over time, probably because it is more embedded into the protein structure and therefore more rigid (Figure 3C).

Table 1. ACPA IgG BCR variable region sequence details. ACPA IgG BCR sequences isolated from single B cells
of 3 RA patients ^{23,24} . Immunoglobulin (IG) heavy (H) and kappa (K) or lambda (L) light chain CDR3 amino-acid
sequences are depicted and the amount of nucleotide mutations compared to germline. The N-linked glycan
motifs presented in the variable heavy (V_{\tiny H}) or light (V_{\tiny L}) chain are visualized together with their respective
location. The germline motifs are depicted based on the IMGT database.

ACPA lgG	IGH-CDR3 AA	IGH nt-mutations	IGK/L-CDR3 AA	IGK/L nt-mutations
7E4 ²⁴	CVRIRGGSSNW	28	CAAWNGRLSAFVF	28
2E4 ²³	CARGSFLERPESVPFHPW	71	CLQYHAEPYTF	61
1F2 ²³	CVRGGSLGIFGGSVGYW	44	CQSYYRGDWVL	46
2G9 ²³	CVRWGEDRTEGLW	61	CMQRLRFPLTF	31
3F323	CARGTYLPVDESAAFDVW	56	CQQYYEAPYTF	37
2D11 ²³	CARRGGKDNVWGDW	21	CQQYNDWPVTF	11

* Amino-acid sequence and location.

⁺ Determined by IMGT/ V-QUEST.

These results suggest a possibly higher impact of the LC glycan on antigen binding. We next visualized the amino acids and N-glycan monosaccharides involved in the hydrogen bond interactions as donors or acceptors within the respective three-dimensional (3D) variable domain structures based on IMGT (the international ImMunoGeneTics information system; Figure 3D and Figure S2 based on Table S4 to S6). The results show that hydrogen bonds between the core fucose or N-acetylglucosamine and the antibody structure were mainly formed between amino acids within or close to the N-linked glycan sites. Notably, N-glycan galactoses or terminal sialic acids also interacted with aa in the CDR1/CDR2 or even in the CDR3 (Figure 3D and Figure S2C). The negatively charged sialic acids of the 3F3 HC N-glycan interacted, for instance, with Y100, an amino acid located in the positively charged binding pocket. As amino acids located around Y100 in the HC are mainly involved in citrulline binding, an interaction with the N-linked glycan will likely affect antigen recognition (Figure 2D). Together with the structural composition of the antigen-binding pockets, these results show that carbohydrates are positioned in close vicinity to the antigen-binding pocket and are likely capable of disrupting antibody-antigen interactions by forming hydrogen bond interactions with the CDR structures directly involved in binding.

Disialylated complex-type VDG modulates binding to citrullinated antigens

As the data described above indicate that VDGs can modulate antigen binding, we next sought to validate the impact of carbohydrates attached to the variable domain on binding using the different monoclonal ACPA IgG variants. As shown by crystallography, the minor amino acid substitutions (germline back mutation) used to generate non variable domain glycosylated variants did not change CDR conformations, and hence, a potential impact on antigen-binding ability can be

N-linked glycan motifs $V_{_{H}}^{^{*}}$	N-linked glycan	Germline motifs $V_{\rm H}{}^{\rm +}$	Germline
	motifs V_{L}^{*}		motifs V_{L}^{\dagger}
NES (CDR1)	NVT (FR1)	SES (CDR1)	KVT (FR1)
NIT (FR3) and NST (FR3)	х	TMT (FR3) and STA (FR3)	х
NFS (CDR1) and NLT (CDR2-FR3)	NLT (FR1)	TFT (CDR1) and NPT (CDR2-FR3)	NFM (FR1)
NGS (CDR1), NTS (FR3) and NMT (FR3)	NIS (FR3)	GGS (CDR1), NPS (FR3) and KLS (FR3)	KIS (FR3)
NMT (FR3) and NTS (FR3)	NLT (FR3)	TMT (FR3) and STA (FR3)	TLT (FR3)
NFT (CDR1)	х	SFT (CDR1)	х

ruled out (Figure S2B). We identified the effects of various glycan compositions (WT, WT+neu, NG, and NG+neu) (Figure 1D) within six different mAbs (Table 1) on multiple antigenic interactions. Because of their heterogenous binding behaviour, different antigens were tested among different ACPA mAbs. First, binding toward several citrullinated peptides [cyclic citrullinated peptide 2 (CCP2), cit-fibrinogen α 27-43, cit-fibrinogen β 36-52, CCP1, cit-vimentin 59-74 and cit-enolase 5-20] was investigated, and our results showed an overall reduced binding capacity of the VDGexpressing WT variants (Figure 4 and Figures S3 and S4). Cleaving off negatively charged sialic acids increased binding to most antigens. The total absence of VDGs, as in the NG variants, resulted in considerably higher binding capacities (up to 90%) (Figure 4). In particular, VDGs affected the binding of ACPAs toward antigens recognized with low equilibrium K_p values, such as citrullinated enolase for the mAbs 7E4 and 1F2 or CCP1 for the mAb 3F3 (Figure 4A, Figure S4 and Table S2). Interactions with citrullinated antigens that show high maximal binding rates already at low concentrations, as observed for the CCP2 or citrullinated fibrinogen, were less affected by the presence of the carbohydrates (Figure 4A and Figure S4). Although no direct affinities could be calculated from the biphasic binding curves (Table S1), surface plasmon resonance (SPR) measurements showed a similar interference of the 7E4 VDG with citrullinated antigens. The lack of sialic acids resulted in higher response units, which raised even further, when the entire glycan structure was absent (Figure S5). Similar observations as made using citrullinated peptides as the antigenic target for ACPAs were obtained when analyzing binding toward citrullinated proteins (native autoantigens) such as citrullinated fibrinogen, vinculin, ovalbumin (OVA) and myelin basic protein (MBP). Alike the peptide binding results, the absence of VDGs raised the binding to most protein targets (Figures 4, C and D, and Figure S4), although 1F2 binding remained similar, an observation also made for some peptide antigens (Figure 4B).



◄ Figure 2. Crystal structures of ACPA Fab fragments and their peptide-binding modes. (A) Crystal structures of ACPA IgG Fab domains complexed with their respective peptides ($3F3_{Fab}$:cit-vim 59-74 and $1F2_{Fab}$:cit-CII-C-39) and the unbound (APO) form of $1F2_{Fab}$ in a cartoon representation. The peptides bound to the Fab are shown as sticks with carbon (green), oxygen (red), and nitrogen (blue) atoms. Citrulline residues are colored in magenta. (B) Top view of the ACPA paratope and the bound peptides for $3F3_{Fab}$:cit-vim 59-74 and $1F2_{Fab}$:cit-CII-C-39 and for $1F2_{Fab}$ in an unbound state. The peptides are shown as sticks with carbon (green), oxygen (red), and nitrogen (blue) atoms. (C) Detailed paratope/epitope interactions between Fab fragments and their bound ligands shown for $3F3_{Fab}$:cit-vim 59-74 and $1F2_{Fab}$:cit-CII-C-39. Fab HC and LC are depicted as cartoon or sticks. All residues involved in the polar/hydrophobic interactions to the ligands are marked. Water molecules are depicted as red spheres, hydrogen bonds are depicted as black dashed lines, and the zinc ion is depicted as a dark sphere. Citrullinated residues are indicated as CIT. (D) Superposition of the CDR loops (CDR1, CDR2, and CDR3) of $3F3_{Fab}$. $1F2_{Fab}$ and $7E4_{Fab}$. (E) Electrostatic surface potentials of $3F3_{Fab}$ and $1F2_{Fab}$ are depicted. Positively charged regions are colored in red and negatively charged regions in blue. LC is depicted in light gray and HC in steel blue.

Thus, the enzyme-linked immunosorbent assay (ELISA) based binding assays of six different patient-derived monoclonal autoantibodies showed an overall negative impact of VDGs and sialic acids on binding to several potential (auto)antigens (Figure 4D and Figures S3 and S4). Thus, these data are in line with our prediction from the crystal structure analyses and indicate that VDGs can interact with and mask antigen-binding pockets of ACPAs.

VDGs expressed by autoreactive BCRs masks antigen binding on the B-cell surface

The data described above provide novel insights into the potential influence of autoantibody VDGs on antigen binding. To address the question of whether VDG also affects autoantigen binding on the B-cell surface, we investigated the involvement of VDG on antigen binding at the cellular level. To this end, the human B-cell line Ramos, in which the endogenous IGHM, IGHD and IGLC and activation-induced cytidine deaminase were knocked out (MDL-AID KO)²³, was used to express citrullinated protein (CP)-directed BCRs. Two Ramos B-cell transfectants carrying variable domain glycosylated 3F3 and 7E4 membrane IgG (mIgG) BCRs²³ and their NG counterparts were generated (Figure 5A). NG variants were produced as mentioned above by mutating the N-linked glycan sites back to the respective germline amino acid sequences. Transduced B cells were sorted on the basis of green fluorescence protein (GFP) and mlgG expression to ensure an identical amount of WT and NG BCRs on the surface of both cell lines (Figures 5, B and C). The human Ramos B cells transduced with patient-derived autoreactive BCRs were highly GFP and mIgG BCR positive, while the non-transduced control MDL-AID KO cell line showed no BCR and GFP expression (Figure 5B). The occupancy of the N-linked glycan sites and thus the expression of VDGs on the WT mlgGs were verified by gel electrophoresis after B cell lysis and BCR capturing (Figure 5D). A size shift was present between the WT and NG BCRs for both cell lines. This mass shift was higher when comparing the 3F3 to the 7E4 BCR variants due to six N-linked glycans attached to the 3F3 BCR HC and LC chain compared to four glycans attached to the 7E4 mlgG (Figure 5D). Furthermore, we analyzed the specific IgG BCR VDG composition after an in-gel total glycan release (PNGaseF treatment) followed by liquid chromatography-MS (LC-MS). Although this method does not allow a site-specific glycan analysis, the results obtained show that WT BCRs isolated from human Ramos B cells harbor fully processed complex type H5N5F1S2 glycans as opposed to their NG counterparts (Figures 5, F and G and Figure S6, A and B). Likewise, we observed more bisecting N-acetylglucosamine and sialic acid moieties on the VDG (WT) BCR compared to the non-VDG (NG) BCR, which only contained Fc glycans, while galactosylation and fucosylation were comparable between the variants (Figure 5E). We confirmed the expression of complex-type glycans on surface IgG BCRs by cell surface biotinylating and western blot analysis of the captured biotinylated BCRs. To analyze the nature of surface IgG specific glycans, we treated the biotinylated BCR with Endo H, an enzyme that is unable to cleave complex-type N-glycans and only able to remove mannosylated glycans, or PNGase F, which cleaves all N-glycan structures as also elegantly described in previous publications²⁶. The results showed a size shift toward the NG BCR after PNGase F treatment and almost no susceptibility of the BCR VDGs to Endo H treatment (Figure S6C). Thus, these data indicate that surface IgG BCRs on Ramos B cells predominantly express fully processed complex-type H5N5F1S2 glycans in their variable domains, such as the VDGs found on isolated ACPA IgG from patients, and no high-mannose structures.

Next, we used the autoreactive 7E4 and 3F3 BCR-expressing cell lines to identify the impact of mIgG H5N5F1S2 VDGs on autoantigen binding by flow cytometry analyses using labeled citrullinated peptide tetramers²⁷. The results obtained are consistent with the ELISA observations and indicate a reduction in antigen binding for B cells that express VDG BCRs (Figure 6). Similarly to the mAb data, binding of low-affinity antigens by a given ACPA, such as CCP1 for 7E4 or cit-vimentin 59-74 for 3F3, was more affected compared to binding of antigens that display higher equilibrium K_p values, such as cit-fibrinogen α 27-43 for 7E4 and CCP2 for 3F3 (Table S2 and Figure 6).

Figure 3. Fab structures crystallized with the starting (core) monosaccharides of the VDG and MD simulations to predict VDG-antibody interactions. (A) Fab structures of $7E4_{Fab}$;cit-CII-C-48, $3F3_{Fab}$;cit-Vim 59-74, and $1F2_{Fab}$;cit-CII-C-39 crystallized with the first two *N*-acetylglucosamines (GlcNAcs) of the VDG. LC is depicted in light gray and HC in steel blue. The peptides bound to the respective Fab are depicted as sticks with citrulline residues (yellow), carbon (green), oxygen (red), and nitrogen (blue) atoms. The GlcNAcs are presented as sticks in magenta. (**B**) MD simulation, two time points, of $7E4_{Fab}$, $3F3_{Fab}$ and $1F2_{Fab}$ crystal structures modeled with full length disialylated VDGs represented as sticks in magenta. (**C**) Hydrogen bond (H-bond) interactions between the VDG (HC or LC) and the antibody structure are visualized over simulation time. (**D**) ACPA $3F3_{Fab}$ HC and LC variable gene amino acid sequence is depicted on the basis of IMGT (2D). Acceptors/-donors of H-bond interactions between the antibody structure and the HC (#1 and #2)/LC N-glycans are visualized. High occupancy is depicted in dark blue (HC: 13%, LC:16%), and low occupancy is depicted in light blue (0.01 to 0.2%). Amino acids and their respective interaction partners (N-glycan monosaccharides) are shown. Blue square, GlcNAc; green circle, mannose; yellow circle, galactose; red triangle, fucose; purple diamond, α 2,6-linked *N*-acetylneuraminic acid (sialic acid). **•**



Overall, our results demonstrate the first human B-cell model that can be used to study the impact of BCR glycans on autoantigen recognition, B-cell functions, or B-cell fate. Our findings show that VDGs are not only able to modulate antigen interactions on secreted autoantibodies but also capable of affecting the interplay between BCRs and their autoantigens.



Figure 4. Impact of disialylated ACPA VDGs on citrullinated (auto)antigen binding. (A) ELISA titration binding curves of the ACPA 7E4 (0 to 10 μ g/ml) variants (WT, WT+neu, NG, and NG+neu) toward citrullinated peptides (CCP2, CCP1, cit-fibrinogen α 27-43, cit-fibrinogen β 36-52, cit-vimentin 59-74 and cit-enolase 5-20). Binding to the arginine control peptide was subtracted. Reactivity was determined via the optical density (OD) at 415 nm. Each data point represents the mean of two technical replicates, and each binding experiment was repeated two to three times. (B) Relative binding of the ACPA mAb 7E4, 1F2, and 3F3 (2 to 40 µg/ml) variants (WT, WT+neu, NG and NG+neu) to citrullinated peptides. N = 2 to 6. Unpaired two-tailed t- tests assuming the same SD. 7E4 WT-NG: ****p (CCP2, CCP1, cit-vimentin 59-74) < 0.0001 and **p (cit-enolase 5-20) = 0.0005; 7E4 WT-WT+neu: **p (CCP1) = 0.029, *p (CCP2) = 0.0227 and *p (cit-enolase 5-20) = 0.0419; 1F2 WT-NG: *p (cit-vimentin 59-74) = 0.0104 and *p (cit-enolase 5-20) = 0.0101: 1F2 WT-WT+neu: **p (cit-enolase 5-20) = 0.0096 and *p (cit-vimentin 59-74) = 0.0449; 3F3 WT-NG: ***p (CCP1) = 0.0009 and **p (cit-vimentin 59-74) = 0.0010; 3F3 WT-WT+neu: **p (CCP1) = 0.0017 and **p (cit-vimentin 59-74) = 0.0082. (C) Relative binding of the ACPA mAb 7E4, 1F2, and 3F3 (5 to 20 µg/ml) variants (WT, WT+neu, NG and NG+neu) to citrullinated proteins. N = 2. Unpaired two-tailed t-tests assuming the same SD. 7E4 WT-NG: **p (cit-vinculin) = 0.0022, **p (cit-MBP) = 0.0014 and *p (cit-OVA) = 0.0032; 3F3 WT-NG: ***p (cit-MBP) = 0.0003; 3F3 WT-WT+neu: *p (cit-MBP) = 0.0213. (D) Heat- map of relative binding (0%, blue, 100%, red) of all monoclonal ACPA (7E4, 2E4, 1F2, 2G9, 3F3, and 2D11) to citrullinated peptides and proteins. Non determined reactivities are illustrated with a cross. N = 2 to 3. (F) Relative binding of ACPA 7E4, 1F2, and 3F3 (5 to 20 μ g/ml) variants (WT, WT+neu, NG, NG+neu) to citrullinated proteins. N = 2 to 6.

Functional impact of sialylated BCR VDG on B-cell activation

Next, we wished to delineate whether disialylated VDGs expressed on mlgG BCRs influence B-cell activation independently of the "masking" effect on antigen binding. For this, we performed calcium flux experiments and investigated the phosphorylation of the spleen tyrosine kinase (Svk), a central kinase in BCR signal initiation and amplification^{23,28}. Ramos WT and NG B cell lines, displaying an identical BCR surface expression (Figure 5C), were either stimulated with anti-IgG F(ab)'2 or with a citrullinated antigen (Figure 7A). For these experiments, the highaffinity antigen CCP2-streptavidin (strep.) tetramer was used, which exhibits similar binding strengths to both WT and NG BCRs (Figure 6B). Both stimuli triggered calcium flux within a few seconds, as measured by the ratio of Indo-1 in a calcium-bound and unbound state (Figures 7, B and C). No calcium flux was detected after stimulating the cells with the non-citrullinated, arginine-containing control tetramers (CArgP2-strep.) (Figure S7B). Likewise, activation with antigens that are recognized with lower affinity, such as cit-fibrinogen β 36-52, did not result in sufficient B-cell activation in this assay (Figure S7B). Intriguingly, when stimulated with either anti-IgG F(ab)'2 or the citrullinated antigen (CCP2), both WT cell lines (3F3 and 7E4) exhibited higher calcium flux peaks compared to the cells expressing NG BCRs (Figures 7, C and D). B cells expressing VDG BCRs displayed not only a higher maximal calcium flux but also a faster calcium release as indicated by the slope of the curve (Figure 7C and Figure S7A). In addition, we analyzed the phosphorylation of the BCR signal-transducing kinase Syk in these cell lines by phospho-flow and western blotting. Notably, and in line with the calcium release, both WT cell lines showed a higher pSyk level than the NG BCR carrying cells when stimulated with either anti-IgG F(ab)'2 or CCP2-strep. (Figures 7, D, F and G, and Figure S7C). In addition, depending on the stimulus given, our data indicated not only a higher activation signal but also a prolonged Syk phosphorylation after stimulation for different time periods (2 to 20 min) (Figure 7E). Thus, these data indicate that human B-cell lines expressing a variable domain glycosylated autoreactive BCR depict a higher activation after stimulation, potentially explaining why autoreactive B cells with VDG modified BCRs have a selection advantage in RA.



Figure 5. Generation of human Ramos B-cell lines carrying disialylated VDG BCRs. (A) Schematic depiction of generated human Ramos B-cell transfectants with mlgG VDG⁺ BCRs (WT) and VDG⁻ BCRs (NG). The untransduced MDL-AID KO cell line shows no GFP and endogenous BCR expression (GFP⁻, BCR⁻, VDG⁻). (B) GFP and BCR surface expression of the MDL-AID KO control cell line and Ramos cells expressing 3F3 mlgG with (WT) and without (NG) VDGs. (C) Histograms depicting an identical WT and NG mlgG BCR expression of the 3F3 and 7E4 Ramos B-cell lines and no BCR expression of the MDL-AID KO cell line. (D) Size shift between VDG⁺ and VDG⁻ 3F3 and 7E4 BCRs visualized on a 4 to 15% SDS polyacrylamide gel (Bio Rad). No mlgG BCR protein band was detected for the MDL-AID KO cell line. The size was determined using the PageRuler[™] Plus Prestained Protein Ladder (Thermo Fisher Scientific). (E) Bar graphs of the relative abundance of galactosylation, sialylation, bisecting GlcNAc, and fucosylation per N-glycan and sialylation per galactosylation on the Ramos 3F3/7E4 WT, NG, and MDL IgG BCRs after passing quality control (QC) settings. N = 3 (biological replicates). Multiple non paired t-tests (Bonferroni-Dunn method): ****p < 0.0001, ***(3F3) p = 0.0008 and ***(7E4) p = 0.0004. (F) Absolute intensity of glycan traits expressed on the MDL and 3F3 WT and NG BCR after passing QC settings.</p>

Effect of VDG on BCR downmodulation and antigen internalization

To determine whether variable domain glycosylation also regulates BCR downmodulation after antigen exposure, we stimulated the Ramos B cell lines for several minutes with the CCP2-strep. or the CArgP2-strep. control tetramer at 4 °C. We then let the cells incubate at 37 °C to allow internalization and measured mIgG BCR downmodulation using a Fab anti-human IgG detection antibody (Figure 8A). The transduced and GFP-positive Ramos cells showed no changes in cell surface BCR expression after phosphate-buffered saline (PBS) treatment or when incubated on ice. In contrast, downmodulation could be readily detected after 15 min of incubation at 37 °C (Figure 8B), indicating rapid antigen-induced BCR internalization. We identified a timedependent decrease in BCR surface expression for both the WT and the NG B-cell line (Figure 8C). Notably, the signal of surface-bound BCRs after antigenic stimulation was significantly decreased for the B-cell lines carrying NG BCRs compared to their variable domain glycosylated counterparts (Figures 8, D to F), despite the equivalent binding intensity of CCP2 at baseline. Loss of cell surface BCR expression was faster for the 3F3 expressing cell lines as most of the mlgG was lost within 30 min, whereas no downmodulation could be detected after stimulating the cells with the control peptide (Figures 8, E and F).



Figure 6. Binding of Ramos B cell lines carrying a VDG⁺ or VDG⁻ BCRs to citrullinated antigens. (A) Histograms of Ramos 3F3 and 7E4 B cells carrying VDG⁺ or VDG⁻ BCRs and their binding to CCP2/cit-vimentin 59-74 or CCP2/CCP1-strep. APC labeled tetramers, respectively. Binding to the arginine control peptides is shown in dark (WT) and light (NG) gray. (**B**) Binding titration curves of 7E4/ 3F3 WT and NG BCRs to citrullinated peptide-strep. tetramers (CCP2, CCP1, cit-fibrinogen α 27-43, cit-fibrinogen β 36-52, and cit-vimentin 59-74) and their respective control peptides. N = 2. The y-axis depicts the MFI ratio between antigen binding and mlgG BCR expression. (**C**) Binding of 3F3/ 7E4 WT and NG Ramos B cells to citrullinated peptide-strep. tetramers (1 to 5 μ g/ ml). The y axis depicts the MFI ratio between antigen binding and mlgG BCR expression. (**D**) Relative binding of 7E4/ 3F3 WT and NG BCRs towards five citrullinated peptides (0%, blue; 100%, red).

Furthermore, confocal microscopy confirmed the flow cytometry-based BCR downmodulation data. We assessed antigen internalization by stimulating the cells with AF568-labeled CCP2-strep. tetramers. BCR modulation was evaluated by staining the remaining BCRs on the fixed cell surface after antigenic stimulation. mlgG BCRs were detected using a Fab anti-human lgG antibody coupled to AF647. All antigen-bound mlgG BCRs were still located on the B-cell surface after incubation on ice. Incubating the stimulated B cells at 37 °C for several minutes induced BCR clustering followed by antigen internalization and a reduced amount of surface mlgGs (Figure 8G). The quantified results showed that especially NG BCRs were markedly downmodulated after antigenic stimulation and incubation for 5 min at 37 °C, while WT BCRs were still stably expressed on the B cell surface (Figure 8H). Despite a similar binding toward the CCP2-strep. tetramer (Figure 8I), more antigen-bound BCRs were internalized, when no VDGs were present on the 3F3 and 7E4 mlgGs (Figure 8J and Figure S7D). These results, together with the flow cytometry based BCR expression data, indicate that human BCRs harboring VDGs stay longer on the B-cell surface in an antigen-bound manner.

The role of the sialic acid-binding lectin CD22 on BCR VDG-mediated effects

To identify whether interactions with the α 2-6-linked sialic acid-binding lectin (Siglec) CD22 can explain the effect of VDGs on BCR signaling and cell surface expression, we performed CRISPR Cas9 KO in the 3F3 WT and NG BCR-expressing Ramos B cell lines. The KO was validated by flow cytometry, showing that the surface expression of CD22 was clearly absent in both the 3F3 WT and NG CD22KO cell lines compared to their non-KO counterparts (Figures 9, A to C). Furthermore, we verified the KO of the CD22 target gene by bulk sequencing. Both KO cell lines showed a gene KO efficiency of more than 90% as evidenced by TIDE (tracking of indels by decomposition) analysis (Figure S8A). Mostly nucleotide deletions (>70%) were observed and, to a lower extent, +1 nucleotide insertions (< 20%) with the highest frequency of guanine insertions (> 85%) (Figures S8, A and B). Both CD22KO cell lines, expressing a VDG (WT) or NG BCR, as well as their CD22⁺ counterparts, were assessed in functional assays. Flow cytometry based phospho-flow depicted, in line with our previous experiments, an increased pSyk expression of stimulated 3F3 cells carrying a WT compared to a NG BCR (Figure 9D). Furthermore, we identified an increased expression of pSyk in stimulated CD22KO cell lines compared to their CD22-expressing counterparts (Figure 9D). The increased signaling capacity after the CD22KO was similar for the WT and NG variants, indicating that VDGmediated differences in BCR signaling are independent of CD22. These results were confirmed by calcium flux (Figure 9E) and western blot analyses, which showed that the absence of CD22 did not affect an increased pSyk expression in the presence of the VDGs (Figure 9F). In addition, we could not observe an impact of the Siglec CD22 on BCR downmodulation as evidenced by the BCR surface expression data of the CD22KO cell lines after activation with a citrullinated antigen (Figure 9G). However, similar to our previously presented results, the NG BCR was downmodulated quicker compared to its VDG counterpart (Figure 9G). Together, these data point out that the highly sialylated VDGs alters BCR downmodulation and signaling via a mechanism that does not involve the sialic acid-binding lectin CD22.



Figure 7. Impact of mIgG BCR VDGs on human Ramos B-cell activation. (A) Ramos B cells (WT and NG) were stimulated with antigen (CCP2-strep.) or anti-IgG F(ab)'2. B-cell activation was analyzed via Ca²⁺ release or the phosphorylation of Syk. (B) Ca²⁺ flux (Ca²⁺ -bound Indo-1/ unbound Indo-1) of 3F3 WT and NG Ramos B cells after stimulation with a-IgG F(ab)'2. (C) Ca²⁺ flux overlays of 7E4/ 3F3 WT and NG Ramos B cells after stimulation [a-IgG F(ab)'2 or CCP2-strep.]. (D) Paired analysis of Ca²⁺ flux and pSyk(Y348) expression after 5 min of stimulation [a-IgG F(ab)'2 or CCP2-strep.] for 3F3/7E4 WT and NG Ramos B cells. Paired two-tailed t-test. N = 5 to 7. 3F3 Ca²⁺ flux: **p = 0.005 and *p = 0.0316; 3F3 pSyk: *p = 0.0109; 7E4 Ca²⁺ flux: *p = 0.0309 and *p = 0.0441; 7E4 pSyk: **p = 0.0089 and *p = 0.0276. (E) pSyk (Y348) time-point analysis of 3F3/7E4 WT and NG Ramos B cells after adding no stimulus or 2, 5, 10, 15, and 20 min of stimulation [a-IgG F(ab)'2 or CCP2-strep.]. pSyk(Y348) MFI ratio (stimulated cells) is depicted. (F) pSyk (Y348) histograms of unstimulated, CCP2-strep. or a-IgG F(ab)'2 stimulated 3F3/7E4 WT and NG Ramos B cells. CD22, Syk, pCD22 (Y822) and pSyk(Y352) expression are shown. Cell Iysates of 1 million (unstimulated and stimulated first slot), 0.5 million (stimulated second slot) and 0.25 million (stimulated third slot) cells were blotted. Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) was used as a loading control, and 1 million MDL-AID KO cells were added as an additional control.



Figure 8. Effect of BCR VDG on antigen internalization and BCR downmodulation. (A) Ramos B cells were incubated at 4 °C, stimulated with antigen or PBS and incubated at 4 °C or 37 °C to allow BCR downmodulation. The remaining surface BCRs were stained (Fab anti-human IgG-AF647). (B) GFP and mIgG expression of MDL-AID KO, 3F3 WT and NG cells after PBS or CCP2-strep. treatment (4 °C or 15 min at 37 °C). (C) Histograms (mIgG) of CCP2-strep. stimulated 3F3 WT or NG Ramos B cells (4 °C or 5, 15, 30 min at 37 °C). (D) Histogram (mIgG) overlay of 3F3 WT and NG Ramos B cells after PBS or CCP2-strep. treatment (4 °C or 15 min at 37 °C). (E) BCR downmodulation of 3F3 WT and NG Ramos B cells after PBS, CCP2-strep. or CArgP2-strep. stimulation (4 °C or 5, 15, 30 min at 37 °C). (E) BCR downmodulation of 3F3 WT and NG (F) 7E4 WT and NG after PBS, CCP2-strep. or CArgP2-strep. stimulation (4 °C or 5, 15, 30 min at 37 °C). Paired two tailed t- test. N = 3 to 4. 3F3:*(5 min) p = 0.0146 and **p = 0.0054; 7E4 **p = 0.0054. (G) Spinning disk confocal microscopy of GFP+ 3F3 WT and NG cells after CCP2-strep.-AF568 stimulation (incubation at 4 °C or 5, 15 min at 37 °C), 2% PFA fixation and a-IgG-AF647 surface staining. (H) 3F3 WT and NG BCR expression after CCP2-strep. stimulation (4 °C or 5, 15 min at 37 °C). N = 114, 213, 474, 224, 495 and 595 cell slices respectively. Ordinary one-way ANOVA, ****p < 0.0001. (I) CCP2-strep. binding of 3F3 WT and NG at 4 °C. (J) CCP2-strep. internalization of 3F3/TE4 WT and NG BCRs at 4 °C or after 5, 15 min incubation at 37 °C. N(3F3) = 114, 213, 474, 224, 495 and 595 cell slices respectively. N(7E4) = 619, 459, 645, 433, 738 and 302 cell slices respectively. Ordinary one-way ANOVA, ****p < 0.0001.</p>



Figure 9. Impact of CD22 CRISPR Cas9 KO on BCR VDG-mediated effects. (**A**) GFP and CD22 (anti-CD22 APC) expression of 3F3WT, WT CD22KO, NG and NG CD22KO Ramos B cells. (**B**) Histogram overlay (CD22) of unstained cells, 3F3WT, and WT CD22KO or 3F3NG and NG CD22KO Ramos B cells. (**C**) CD22 expression of 3F3WT, WT CD22KO, NG, NG CD22KO and MDL-AID KO Ramos B cells. (**D**) Percentage of pSyk(Y348) positive cells of 3F3 WT, WT CD22KO, NG, and NG CD22KO Ramos B cells after 5, 10, and 15 min of a-IgG F(ab)'2 stimulation. Paired two-tailed t- test. N = 4. 5 min: *(3F3 WT vs. NG) p = 0.0318; 10 min: *(3F3 WT vs. NG) p = 0.0106, *(3F3WT vs. WT CD22KO) p = 0.0298; 15 min: *(3F3WT vs. NG) p = 0.0166, *(3F3NG vs. NG CD22KO) p=0.0315. (**E**) Ca2+ flux of 3F3 WT, WT CD22KO, NG, and NG CD22KO Ramos B cells after a-IgG F(ab)'2 stimulation. Paired two-tailed t- test. N = 4. *(3F3 WT vs. NG) p = 0.0342, *(3F3 NG vs. NG CD22KO) p = 0.0171. (**F**) Western blot analyses of 3F3 WT, WT CD22KO, NG and NG CD22KO Ramos B cell ysates after 5 min of a-IgG F(ab)'2 stimulation. CD22 and pSyk(Y352) expression are shown. β-Actin was used as loading control and MDL-AID KO cell Iysates as additional control. (**G**) BCR downmodulation of 3F3 WT, WT CD22KO, NG, and NG CD22KO Ramos B cells after PBS or CCP2-strep. stimulation and incubation at 4 °C or for 5, 15 or 30 min at 37 °C. Paired two tailed t-test. N = 3. *(4 °C) p = 0.0474, * (5 min 37 °C) p = 0.0249, * (30 min 37 °C) p = 0.0463.

Discussion

We here investigated the impact of VDGs on human B cell biology using citrullinated proteindirected BCR sequences isolated from ACPA-positive patients with RA. These molecules display naturally occurring N-linked glycan sites within the variable regions that are introduced during somatic hypermutation (SHM)¹⁸ and allow the incorporation of VDGs. More than 90% of ACPA IgG harbor VDGs, which makes them a hallmarking feature of the RA-specific ACPA immune response¹⁶. Although readily found on ACPA from patients with RA, VDGs are not abundantly present in healthy individuals¹⁹. Furthermore, it has become evident that the transition to ACPApositive disease is associated with a rise in VDG levels before disease¹⁹, which is intimately connected to an increase in autoantibody levels and a "maturation" of the ACPA response as defined by an increased citrullinated antigen recognition profile and isotype usage. Notably, this maturation is not coupled to an increase in avidity as the ACPA response displays only limited avidity maturation across different (pre)disease stages²⁹. Together, these observations are intriguing as they imply that affinity maturation does not drive the selection of autoreactive ACPA-expressing memory B cells, but rather their propensity to introduce glycosylation sites into their variable domains that potentially support their expansion and/or survival³⁰. BCR analyses of ACPA-expressing B cells from patients with RA indicate that the introduction of glycosylation sites is a consequence of a selective process not explained by the number of SHM. In line with this concept, the most prominent genetic risk factor for RA, the human leukocyte antigen (HLA) class II region, does not associate with the presence of ACPA as such but mainly with ACPA-expressing VDGs^{31,32}. Hence, the acquisition of VDG represents an important event connected to the expansion of the B-cell response occurring before RAonset that is also predictive for disease development in healthy first-degree family members of patients with RA¹⁹. Furthermore, accumulating evidence suggests that N-linked VDGs are also at play in several other human autoimmune diseases including ANCA-associated vasculitis and Sjögren's syndrome^{20,21}.

The data presented in this manuscript support a concept in which VDGs act as a "threshold" for human autoreactive B-cell activation, by showing that CP-directed B cells increase BCR signaling upon the presence of VDG. The introduction of VDGs during SHM is thus likely involved in the selection and activation process of autoreactive B cells in RA and possibly other autoimmune diseases. Next to their ability to increase B-cell activation, VDGs are able to delay BCR downmodulation after antigenic stimulation and "mask" binding to especially "low-affinity" antigens. Whether or how these functional events are interconnected and contribute in setting the "threshold" for autoreactive B-cell activation and survival is unclear. However, it is conceivable that VDGs reduce the breadth of self-antigen recognition in the GC and thereby promote B-cell survival. In case of a ubiquitous expression of self-antigens in the GC environment, self-reactive B cells might be eliminated as demonstrated in mouse

studies³³. A differential impact of VDGs on antigen binding, although not self-antigens, has been reported for high-affinity anti-adalimumab and anti-infliximab clones²² as well as for follicular lymphoma-derived VDGs artificially introduced into anti-NIP or anti-HEL BCRs and expressed in a murine B cell-derived reporter cell line³⁴. Our crystallographic data indicate that naturally occurring VDGs can interact with the antibody paratope and thereby compete with antigens for binding. This might provide a mechanism for autoreactive B cells to escape elimination through recognition of (abundantly) expressed autoantigens, while allowing activation via cross-reactivity to (higher-affinity) foreign-epitopes. Structural data together with modeling studies demonstrate that VDGs undergo hydrogen bond formations with CDR domains and can interfere with antigen binding. Our results also show that ACPAs display an open-ended binding groove and interact specifically with the modified (citrullinated) residue and, to a minor extent, with backbone structures surrounding the modification. This might explain the overall low affinity of ACPA and indicates that already marginal disturbances, by, e.g., carbohydrates, can interrupt binding.

One limitation of our study is that NG mAb variants, used for the binding assays, were generated by mutating N-linked glycan sites back into the predicted germline sequence. This prediction is based on the highest identity with the respective germline sequence but might cause altered antibody functions. However, the crystal structures of, e.g., 7E4_{Fab} showed no structural differences in CDR conformations when expressed with or without VDGs indicating no potential impact of the germline backmutation on antigen-binding ability. In line with the crystallographic results, our ELISA data illustrate that VDG of autoantibodies in RA can diminish binding to several citrullinated peptides and proteins. The interactions between VDGs and the binding pockets can be more easily disrupted by high-affinity antigens, whereas low-affinity antigens are more strongly affected by the masking carbohydrates. We assessed the impact of VDG on antigen binding not only with germline reverted variants, but also confirmed them with neuraminidase treated mAbs harboring glycan traits without the terminal sialic acids. Furthermore, the results obtained for the mAbs 7E4 and 3F3 were in line with the binding BCRs 7E4 and 3F3 expressed on human Ramos B cells.

Although BCR surface expression levels and glycan compositions might differ from the in vivo situation, we show that the Ramos model B cell lines are able to express disialylated and bisected VDG structures, similar to glycan traits observed on secreted autoantibodies from patients with RA. Despite the limitation of testing only two recombinant BCRs, our data provide the first and consistent experimental evidence that VDGs are able to boost BCR signaling, as evidenced by an increase in Syk phosphorylation and calcium release, which presumably provides an advantage to these autoreactive B cells. Enhanced variable domain glycosylation is also thought to provide a survival advantage in follicular lymphoma B cells,

although this has, so far, not been investigated experimentally^{35,36}. A limitation of our study is that we could not confirm our BCR signaling results with citrullinated protein-reactive B cells directly obtained from patients as these cells represent a rare population in peripheral blood of patients with RA (approximately 0.05% of CD19+ B cells). Moreover, at present, it is not feasible to separate these citrullinated protein-reactive B cells in subsets of cells that do or do not express VDGs.

Currently, it is unclear how VDGs affect BCR triggering and thereby set the threshold for activation. It has been suggested that sialylated VDGs interact with sialic acid-binding lectins, such as CD22³⁷, potentially leading to an altered co localization of the IgG BCR and the Siglec in a resting or activated state. Our CD22 KO data show however, no impact of CD22 on the VDGmediated effects on BCR downmodulation and signaling. Furthermore, interactions with other Siglecs or (soluble secreted) lectins, such as galectin 9, may be involved in altered signaling, similar to the role of galectin 9 described in IgM BCR signaling³⁸. Moreover, VDG might alter the molecular organization of B-cell surface structures by forming distinct clusters that are absent on B cells expressing BCRs without VDGs. A diverse BCR organization on B cells expressing variable domain glycosylated versus non variable domain glycosylated BCRs could explain the significant differences in B-cell activation after BCR cross-linking as could differences in BCR internalization, degradation, and modulation³⁹. The presence of VDGs modulates the dynamics of the BCR surface expression after antigenic stimulation. We found a decreased BCR downmodulation and uptake after antigen exposure for variable domain glycosylated BCRs compared to NG BCRs. Thus, BCRs containing N-linked glycans show a decreased ability to be downmodulated from the cell surface and are consequently retained longer on the B-cell surface after triggering with antigen. The prolonged surface expression might modulate BCR signaling strength and duration as observed by our B-cell activation studies similar to previous studies describing increased BCR signaling after inhibition of BCR uptake pathways⁴⁰. Further studies are needed to identify whether signal attenuation and BCR internalization are directly linked and, if so, how VDG affects BCR uptake. In this respect, it will be important to delineate the temporal dynamics and spatial organization of variable domain glycosylated mlgG BCRs at the nanoscale level also in relation to its association with lipid rafts, the actin cytoskeleton, and clathrin-coated pits.

Together, we show that the presence of N-linked glycans on the variable domain affects fundamental mechanisms involved in the functionality of B cells (antigen binding, BCR signaling, and downmodulation) and thus might critically determine the outcome of the B-cell responses. In this respect, the abundance of VDG on the hallmarking autoantibody response in a prominent human autoimmune disease is noteworthy. Thus, we provide experimental evidence for the selective and abundant presence of VDG on autoreactive B cells in RA that likely conveys a signaling advantage, potentially explaining the outgrowth of these B cells and the increase of autoantibody levels towards disease-onset. Our findings are highly relevant for a better

understanding of B-cell driven autoimmune diseases as they provide a rationale on how the acquisition of VDGs might contribute to the escape of critical immune checkpoints in humans and thus the breach of B-cell tolerance.

Materials and Methods

Blood samples from patients diagnosed with RA – Peripheral blood samples from anticitrullinated protein (auto)antibody-positive patients with RA visiting the outpatient clinic of the Rheumatology Department at the Leiden University Medical Center (LUMC) were included in this study. All patients fulfilled the EULAR/ACR 2010 criteria for the classification of RA⁴¹. None of the patients were previously treated with B cell-depletion therapies. Blood samples were taken upon obtaining written informed consent before inclusion and with approval from the local ethics committee of the LUMC, The Netherlands.

Recombinant monoclonal (auto)antibody production - Monoclonal (auto)antibodies were generated on the basis of full-length BCR sequences from APCA⁺ patients with RA^{23,24}. CCP2strep. and CArgP2-strep. tetramers were used to isolate autoreactive B cells as described previously²⁷. Single sorted B cells were cultured on irradiated CD40L cells and a cytokine mixture in IMDM (Gibco) medium for 10 to 12 days. RNA isolation, complementary DNA (cDNA) synthesis, ARTISAN polymerase chain reaction (PCR), and BCR sequencing were performed as described earlier⁴². The 7E4 sequence was provided by T.R., Sanguin, The Netherlands²⁴. For the generation of NG variants, the N-linked glycosylation sites (N-X-S/T, X \neq P) were specifically back-mutated into the germline sequence (based on IMGT) at the respective position (Table 1). WT sequences, including the N-glycan sites, and NG sequences were codon-optimized and the HC/LC variable genes together with 5'-BamHI and 3'-Xhol restriction sites, the Kozak sequence, and the IGHV1-18*01 leader sequence ordered from GeneArt (Life Technologies). The constructs were ligated into a pcDNA3.1 (+) expression vector (Invitrogen) carrying the IGHG1 or the IGLC3/IGKC constant regions (UniProt), respectively, flanking the 3'-Xhol site. The recombinant mAbs were produced in Freestyle[™] 293-F cells (Gibco) as previously stated²³. Glycoengineering was performed by adding D-galactose substrate to the medium before transfection (Sigma Aldrich, G0750-5G). Furthermore, to generate complex-type N-glycans on the antibody variable domains, the recombinant IgGs were co-expressed with 1% β -1,4-*N*-acetylglucosaminyltransferase III (GnTIII), 2.5% α 2,6-sialyltransferase 1 (ST6gaIT) and 1% β -1,4-galactosyltransferase 1 (B4GaIT1). The supernatants were harvested 5 to 6 days after transfection.

IgG purification – IgG1 antibodies were purified using a 1 ml HiTrap Protein G HP affinity column (GE Healthcare) followed by a direct buffer exchange using a 53 ml HiPrep[™] 26/10 Desalting column (GE Healthcare) according to the manufacturer's instructions. The recombinant mAbs were concentrated with Amicon Ultra-15 50K filter devices (Merck) to a final concentration of 1 mg/ml and used for further experiments.

Monoclonal antibody variable domain glycosylation analysis - Human monoclonal (auto) antibodies were analyzed for levels of variable domain glycosylation via SEC, gel electrophoresis, and MALDI-TOF MS analysis. SEC was performed using a Superose6Increase 10/300 GL column (GE Healthcare) according to the manufacturer's instructions. The recombinant proteins were monitored by ultraviolet (UV) absorption at 280 nm. For SDS-PAGE, 1.5 µg of the mAbs was diluted in 4× Laemmli buffer (Bio Rad) with (reduced) or without (non-reduced) 2% B-mercaptoethanol (Sigma Aldrich) and incubated for 5 min at 95 °C. Sample (10 µl) and 3 µl of PageRuler[™] Plus Prestained Protein Ladder (Thermo Fisher Scientific) were loaded on 4 to 15% SDS-polyacrylamide gels (Bio Rad). For protein detection, gels were stained with Coomassie Brilliant Blue G-250 Dve (Thermo Fisher Scientific). MS analysis of released N-glycans was performed as previously described⁴³. In brief, the IgG samples (10 μ g in 5 μ l) were reduced in 10 μ I of 2% SDS and incubated with 0.5 U of PNGase F (Roche Diagnostics, Germany), in 10 μ I 1:1 5x PBS/4% NP-40, overnight at 37 °C. The total released glycan mixture was subjected to sialic acid stabilization by adding 100 μ l ethylation reagent [0.5 M 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride and 0.5 M 1-hydroxybenzotriazole hydrate] and incubating for 1 hour at 37 °C. For glycan purification, the samples were brought to 85% acetonitrile (ACN) (Biosolve, Valkenswaard, The Netherlands) and purified with 15 µl cotton hydrophilic interaction liquid chromatography (HILIC) tips using 85% ACN and 85% ACN+1% trifluoroacetic acid for washing. Released and purified N-glycans were eluted from the HILIC tips using 10 μ I MQ. Three microliters of the purified and ethyl-esterified glycans were spotted on a MALDI target (MTP AnchorChip 800/384 TF, Bruker Daltonics) together with 1 µl of super-DHB (5 mg/ml) in 50% ACN and 1 mM NaOH. Spots were dried at room temperature (RT) and analyzed on an UltrafleXtreme (Bruker Daltonics) operated under flexControl 3.3 (Build 108, Bruker Daltonics). A mass spectrum from m/z 1000 to 3000 was recorded, combining 10000 shots in a random walk pattern at 1000 Hz. The instrument was calibrated with a peptide calibration standard (Bruker Daltonics). MS data were analyzed using flexControl 3.3 and glycan peaks above S/N of nine were included into the analysis.

Fab fragment generation, crystallization and structure determination – For Fab fragment preparation, chimeric 3F3 and 1F2 antibodies were produced on the basis of human variable regions and mouse constant (Fc) domains, expressed in Expi393F[™] cells (Life Technologies), purified as previously described¹⁵ and dialyzed against PBS. The 3F3 and 1F2 Fab fragments were prepared using the ImmunoPure Fab Preparation Kit (Pierce) following the manufacturer's

instructions. Cleavage was evaluated by SDS-PAGE. Fab fragments were further purified by SEC on a HiLoad 16/600 Superdex 200 column. The peptides cit-vimentin 59-74, cit-CII-C-39, and cit-CII-48 used for co-crystallization are described in previous studies^{15,23}. The crystals used for data collection were grown as follows: For 3F3_{rab}:cit-vimentin 59-74, the sitting drop consisted of 1 µl of 3F3_{Esb} (10 mg/ml) mixed with cit-vimentin 59-74 in twofold molar excess in 20 mM tris (pH 7.5), 20 mM NaCl and 0.5 μ l of reservoir solution [0.2 M ammonium chloride (pH 6.3) and 20% (w/v) PEG 3350]. For 1F2_{Fab}:cit-CII-C-39, the sitting drop consisted of 1 μ I of 1F2_{Fab} solution (10 mg/ ml) in 20 mM tris (pH 7.4), 20 mM NaCl mixed with cit-CII-C-39 in twofold molar excess and 0.5 µl of reservoir solution [20% (w/v) PEG 6000, 0.1 M Hepes 7.0 (pH 7.0) and 0.01 M zinc chloride]. For 1F2_{Fab}, the sitting drop was set up from 1 μ l of 1F2_{Fab} (10 mg/ml) in 20 mM tris (pH 7.4), 20 mM NaCl, and 1 µl of reservoir solution [0.2 M sodium acetate and 20% (w/v) PEG 3350]. For 7E4_{rab}-cit-CII-48, the sitting drop consisted of 1 μ l of 7E4_{rab} solution (10 mg/ml) in 20 mM tris (pH 7.4), 20 mM NaCl mixed with cit-ClI-C-48 in twofold molar excess and 1 μ l of reservoir solution [20% (w/v) PEG 3350, 0.1 M bis-tris propane (pH 6.5), and 0.2 M potassium thiocyanate]. The complexes were incubated overnight before crystallization. All crystals were cryoprotected by briefly soaking in the corresponding reservoir solution containing 30% (v/v) ethylene glycol or glycerol before flash-freezing in liquid nitrogen. Diffraction data were collected at the beamlines and with the statistics given in table S3. The images were processed using XIA2⁴⁴, and scaled by AIMLESS⁴⁵ from the CCP4 program suite⁴⁶. The structures of unbound Fab fragments or Fab-peptide complexes were solved by molecular replacement using PHASER⁴⁷. Iterative cycles of the manual model building were performed using COOT⁴⁸ and TLS as well as restrained refinement with Phenix or REFMAC5⁴⁹ until R-factors converged. Refinement statistics are given in table S3. About 5% of the reflections were randomly selected and set aside for unbiased cross-validation (calculation of R_{frag}). The "Protein Interfaces, Surfaces and Assemblies" service PISA at the European Bioinformatics Institute (www.ebi.ac.uk/pdbe/prot int/pistart.html)⁵⁰ was used to analyze molecular surfaces. Structure comparisons and RMSD calculations were performed with SSM⁵¹ as implemented in PyMOL or COOT. Figures were prepared with PyMOL. The crystallographic coordinates and structure factors have been deposited in the PDB with the accession codes 6YXK, 6YXM, 6YXN and 6ZJG.

Molecular-dynamics (MD) simulations - The crystal structures of 7E4_{Fab}, 3F3_{Fab} and 1F2_{Fab} were used as initial structures for the MD simulation. Quick MD Simulator integrated with a module of Glycan Reader at the CHARMM-GUI website (<u>www.charmm-gui.org</u>) was used for preparing the input files for simulation⁵². The N-glycans were predicted by NetNGlyc1.0 Server (<u>www.cbs.</u> <u>dtu.dk/services/NetNGlyc/</u>)⁵³. For simulation, we adopted the default setting from CHARMM-GUI for preparing the input files. Briefly, the TIP3P model was used for explicit water molecules. A distance of 15 Å between the protein atoms and the cubic system boundary was introduced and 150 mM NaCI was added to the system. The CHARMM36 force field was used for proteins and carbohydrates. All calculations were performed at 300 K. The particle mesh Ewald algorithm

was applied to calculate electrostatic forces. Each system was equilibrated in constant particle number, volume, and temperature condition with restraints using CHARMM36. An additional short period of constant particle number, pressure, and temperature equilibration was also applied without restraints for each system. The simulation was performed in Amber 18⁵⁴. All analyses were performed in Visual Molecular Dynamics⁵⁵.

Protein citrullination and citrullinated peptide synthesis - Vinculin, fibrinogen, myelin basic protein (MBP) and OVA (Sigma Aldrich) protein modification (citrullination) was performed as previously described⁵⁶. Four linear N-terminal biotinylated peptides fibrinogen α 27-42, fibrinogen β 36-52, vimentin 59-74 and enolase 5-20 and two cyclic N-terminal biotinylated peptides CCP1 and CCP2 (patent EP2071335) were synthesized including the native arginine and the altered citrullinated residue(s) at the same positions within the peptide sequence²³. The integrity of the synthesized peptides after purification was examined by ultra-performance liquid chromatography (UPLC) on an Acquity instrument (Waters), and the exact mass was measured via MS on a Microflex instrument (Bruker) and crosschecked with the calculated masses.

Antigen-binding assays – To identify the impact of VDGs on antigen binding, ELISAs were performed as described earlier²³. Briefly, 1 to 10 μ g/ml of biotinylated citrulline or arginine containing peptides were coupled to pre-coated streptavidin plates (Microcoat, #65001) for 1 hour at RT. mAbs were added in PBS/1% bovine serum albumin (BSA)/0.05% Tween (PBT) (Sigma Aldrich) and incubated for 1 hour at 37 °C. For protein ELISAs citrullinated or native proteins (10 μ g/ml) were directly coupled to Nunc Maxisorp plates (Thermo Fisher Scientific) and incubated overnight on ice at 4 °C. Following blocking with PBS/2% BSA for 6 hours, mAbs were added in PBT and incubated overnight on ice at 4 °C. Antibody interactions were detected using a horseradish peroxidase (HRP) conjugated monoclonal mouse anti-human IgG (Fc) C_H2 antibody (Bio Rad, MK1A6) to prevent VDG influences on secondary antibody binding. ELISA readout was performed using ABTS and H₂O₂. For SPR measurements, biotinylated citrullinated peptides were mounted on a streptavidin iSPR chip (Ssenc BV), and mAb analyte binding was assessed. An SPR imaging system for multiplexing 96 biomolecular interactions (IBISMX96, IBIS Technologies, Enschede, The Netherlands) was used.

Human Ramos B-cell transfectants – Human Ramos B-cell transfectants carrying 7E4 and 3F3 mlgG BCRs were generated as described earlier²³. In brief, HC and LC containing single vector constructs were created with the In-Fusion HD Cloning Kit (Clontech) using the pMIG-IRES-GFP-2AP vector backbone including the IGHG1 transmembrane domain. Variable gene sequences were cloned with and without N-linked glycan sites to generate WT and NG constructs. Inserts were verified by Sanger sequencing performed on Applied Biosystems 96-capillary (ABI3730) systems (LGTC facility, Macrogen). Retroviral transductions were performed as previously described⁵⁷. Briefly, Phoenix-ECO (ATCC CRL-3212[™]) cells were transfected with PolyJet DNA transfection reagent (SignaGen Laboratories). Retrovirus containing supernatants were collected 72 hours after transfection and used for the transduction of MDL-AID (IGHM, IGHD, IGLC and AID KO lymphoma Ramos B cells carrying *slc7a1*. Ramos cell lines were cultured in RPMI1640/GlutaMAX[™]/10% fetal calf serum (FCS)/10 mM Hepes medium (Thermo Fisher Scientific) with penicillin/streptomycin (100 U/ml; P/S) (Lonza).

Ramos BCR glycan analysis - For BCR glycan analysis, 20 million human Ramos B cells were lysed in PBS+1% Triton- X100 for 60 min at 37 °C, followed by total IgG BCR capturing using CaptureSelect[™] FcXL Affinity Matrix (Thermo Fisher Scientific) and an overnight incubation at 4 °C. Laemmli buffer $(4\times)$ was added to the IaG BCR/FcXL bead slurry, boiled for 5 min at 95 °C, and loaded on a 4 to 15% SDS gel (Bio Rad). Proteins were detected with Coomassie Brilliant Blue G-250 Dye (Thermo Fisher Scientific). The visualized IgG BCR bands were extracted from the gel, to exclude glycosylated contaminants, and transferred to Eppendorf tubes. The gel pieces were washed first with 25 mM ammonium bicarbonate (ABC), followed by ACN and, for protein reduction, incubated for 30 min at 56 °C in 10 mM dithiothreitol (DTT)/25 mM ABC. After the reduction step, the gel pieces were washed with ACN and, for alkylation, incubated for 30 min at RT in the dark in 55 mM iodoacetamide/25 mM ABC to block reactive cysteine groups. After washing steps with 25 mM ABC and ACN, the gel bands were dried in a centrifugal vacuum concentrator for 5 min. N-linked glycans were released with 30 U of PNGase F (Roche Diagnostics, Germany) in 2%- NP-40/2.5× PBS by an overnight incubation at 37 °C. Released glycans were subsequently labeled with 2-aminobenzoic acid (Sigma Aldrich) and 2-picoline borane (Sigma Aldrich) and purified with 15 µl cotton HILIC tips as described above. Glycans were separated on an Ultimate 3000 UHPLC system (Dionex/ Thermo Fisher Scientific, Breda, The Netherlands) coupled to a MaXis Impact HD quadrupole-TOF mass spectrometer (MaXis HD, Bruker Daltonics, Bremen, Germany) equipped with a CaptiveSpray NanoBooster source (Bruker Daltonics, Bremen, Germany) as previously described⁵⁸. In short, 20% of the released, purified, and labeled glycans were loaded onto a C18 trap column (Acclaim PepMap 100; 100 μ m by 2 cm, particle size of 5 μ m, pore size 100 Å; Dionex/Thermo Fisher Scientific) and washed for 2 min with 15 μ l/min of 0.1% formic acid (FA)/1% ACN. Glycans were subsequently separated on a C18 analytical column (Acclaim PepMap 100; 75 μ m by 15 cm, particle size of 3 μ m, pore size of 100 Å; Dionex/Thermo Fisher Scientific), and elution was performed at a flow rate of 700 nl/min with buffer A [0.1% FA (v/v)] and buffer B [95% ACN/ 0.1% FA (v/v)]. A gradient of 1% to 70% buffer B in 70 min was applied (t = 0 min, B = 1%; t = 5 min, B = 1%; t = 30 min, B = 50%; t = 31 min, B = 70%; t = 35 min, B = 70%; t = 36 min, B = 1%; t = 70 min, B = 1%). The CaptiveSpray NanoBooster was operated with ACN-enriched gas (0.2 bar) and dry gas (3 liters/min) at 180 °C and a capillary voltage of 1150 V. Mass spectra were acquired within a mass range of m/z 550 to 1800. LC-MS data were first examined manually using DataAnalysis (Bruker Daltonics). Data processing, including peak integration, was performed using LaCyTools v1.1.0. Glycans with a S/N above nine, a mass accuracy of +/-20, and an isotopic peak quality of 0.2 were included into the analysis⁵⁹.

Cell surface biotinylating and surface IgG Ramos BCR glycan analysis - Ramos cell surface biotinylating was performed according to the manufacturer's instructions of the Pierce Cell Surface Protein Biotinylating and Isolation Kit (#A44390). In brief, 24 million cells were harvested and biotinylated using Sulfo-NHS-SS-Biotin for 10 min at RT. The label was removed by washing with tris-buffered saline following cell lysis for 30 min on ice. Biotinylated surface proteins were captured using NeutrAvidin[™] Agarose and eluted in elution buffer containing 10 mM DTT. Subsequently, 2% SDS was added, and the samples were incubated for 30 min at 60 °C. The denatured, reduced and biotinylated surface proteins were treated with 2 U of Endo H (Roche Diagnostics, Germany) in 50 mM sodium acetate buffer (pH 5.5) or 2 U of PNGase F (Roche Diagnostics, Germany) in 1:1 5XPBS/4% NP-40 overnight at 37 °C. The presence of high-mannose (Endo H treatment) or complex-type (PNGase F treatment) N-glycans on the surface IgG BCRs was identified via western blot analysis as described below.

Assembly of ribonucleoprotein complexes – Assembly of ribonucleoproteins (RNPs) was performed in a similar way as described in the CRISPR genome editing user guide from integrated DNA technologies (IDT). Briefly, pre-annealed CD22 exon 1 specific complexes of CRISPR-targeting RNA (crRNA) and transactivating crRNA (tracrRNA) were designed via IDT. The single guide RNA (sgRNA) Hs.Cas9.CD22.1.AB (sequence: 5'-TGTCATTGAGGTGCACCGGG-3'; position: 35332832; on-target score: 52; off-target score: 76) was diluted to 44 μ M in IDTE buffer (IDT). Alt-R S.p. Cas9 Nuclease V3 (IDT) was diluted to 36 μ M in resuspension buffer R (IDT). 90 pmol sgRNA was combined with 30 pmol of Cas9 to form the RNP complex.

Ramos cell electroporation with RNP complexes - Two million 3F3 WT and NG mlgGexpressing Ramos B cells were used per transfection reaction and washed with PBS to remove culture medium FCS. Ramos cell pellet was homogenously resuspended with 100 μ l of 11 μ M RNP and 11 μ M Cas9 Electroporation Enhancer (IDT) diluted in resuspension buffer R (Invitrogen). Electrolyte buffer (3 ml; Invitrogen) was added to the transfection tube of the Neon transfection machine (Invitrogen). Cell/RNP mixture was pipetted into the Neon tip and transferred to the transfection tube. Electroporated cells were transferred using the following settings: V = 1350, W = 30 ms and P = 1. Electroporated cells were transferred into a 24-well plate and supplemented with 1 ml of prewarmed medium. Fluorescence activated cell sorting (FACS) or flow cytometry validation experiments were performed after 1 week of culturing using an anti-CD22 allophycocyanin (APC) labeled antibody (BD, #562850) on a BD 4 laser Aria1 FACS or a BD LSRII-flow cytometry instrument, respectively.

Genomic DNA isolation, bulk sequencing and TIDE analysis – Genomic DNA was isolated from 5 million 3F3 WT/NG Ramos B cells with and without CD22 CRISPR Cas9 KO using the Isolate II Genomic DNA Kit (Bioline, BIO-52066) according to the manufacturer's instructions. Isolated genomic DNA (50 ng) was PCR amplified 500 to 1500 base pairs enclosing the designed editing

site using the pre-designed forward and reverse primers (10 μ M) and myTaq mix (Bioline, BIO-25041). PCR gel bands were purified using the PCR Clean-up Gel Extraction Kit (Takara), and 25 to 50 ng/ μ l were prepared for Sanger sequencing performed on an Applied Biosystems 96-capillary (ABI3730) system (LGTC facility, Macrogen). Sequences were analyzed using SnapGene software V5.0.4. The efficiency of the CD22 KO was quantified using TIDE analysis version 3.3.0 (alignment window: 100 to 217, decomposition window: 242 to 481, indel size: 10, and p threshold: 0.001)⁶⁰.

Flow cytometer experiments - Binding of the generated 7E4/ 3F3 WT and NG mlgG and GFP-expressing B-cell lines to citrullinated antigens was analyzed by flow cytometry. Ramos B cells were stained with APC-labeled cit-peptide SA-tetramers (0 to 5 μ g/ml) (CCP2, CCP1, cit-fibrinogen α 27-43, cit-fibrinogen β 38-52 and cit-vimentin 59-74) and 0.5 μ g/ml goat anti-human lgG-Fc phycoerythrin (PE) (#12-4998-82) in staining solution (PBS/0.5%BSA/ 0.02% NaN3) via incubation for 30 min on ice. Tetramer formation was performed as previously described²⁷. Unmodified peptides including arginine residues were measured as negative controls in the same concentration range (0 to 5 μ g/ml) to analyze antigen-specificity.

B-cell activation was measured via calcium release of the generated B-cell lines after stimulation. Therefore, 1 million mlgG and GFP-expressing B cells were collected and stained with 200 µl of calcium-indicator loading dye medium, 2 µM Indo-1 AM (#ab142778) and 0.05% pluronic acid (#P6866) in 1 ml of stimulation medium (RPMI1640/100 U/ml P/S/GlutaMAXTM/10 mM Hepes/2%FCS), for 35 min at 37 °C in the dark. After washing with stimulation medium only, B cells were incubated with 500 µl stimulation medium plus 2 mM calcium and incubated on ice, in the dark until usage. Fifteen min before the analysis, B cells were prewarmed in a water bath at 37 °C to decrease baseline activation upon measurement. The analysis was performed on a BD LSRII-flow cytometry or a Cytek Aurora 5L instrument including a UV laser and acquiring 500 cells/s at a high speed. After 2 min of baseline measurement, 100 µl of stimulus [CCP2/CArgP2 (5 µg/ml) or anti-human IgG F(ab)'2 (80 µg/ml)] was added to the samples and mixed adequately, and the measurement continued for another 5 min until the signal reached almost baseline again. The calcium flux was measured as the ratio of calcium-bound Indo-1 to unbound Indo-1. MDL-AID KO B cells, without an endogenous BCR, were used as negative controls in all experiments. The maximal calcium flux (peak-baseline signal) and its speed (slope of the curve) were analyzed.

BCR signaling was analyzed via the intracellular expression of pSyk after activation. Therefore, 1 million mIgG and GFP-expressing B cells were collected and stimulated with CCP2-strep. tetramer (5 µg/ml) or 15 µg of anti-human IgG F(ab)'2 for 0, 2, 5, 10, 15, or 20 min at 37 °C in stimulation medium (RPMI/100 U/ml P/S/GlutaMAX[™]/10 mM Hepes/1% FCS). Afterward, cells were fixed (BioLegend Fixation Buffer, #420801) and permeabilized (True-Phos[™] Perm Buffer, #425401). After washing, the intracellular expression of phosphorylated Syk was analyzed with a mouse anti-human pSyk(Y348)-PE mAb (#moch1ct, eBioscience[™]) diluted 1:20 in staining solution. The rate of pSyk expression in Ramos cells was calculated as the percentage of pSyk(Y348)⁺ cells or the pSyk median fluorescence intensity (MFI) ratio between stimulated and unstimulated cells. Gating was based on the MDL-AID KO control cell line stimulated with CCP2 or anti-IgG F(ab)'2 respectively.

To assess BCR modulation, 0,2 million mlgG and GFP-expressing Ramos B cells were first incubated for 30 min on ice followed by a 15 min stimulation at 4 °C with either control PBS or CCP2-/CArgP2-strep. tetramers (5 μ g/ml) in PBS/2% FCS. The stimulated B cells were then incubated for 0, 5, 15 or 30 min at 37 °C to allow BCR downmodulation. The remaining surface BCRs were stained with a AF647 NHS (*N*-hydroxysuccinimide) ester (Thermo Fisher Scientific, #A20006) labeled Fab goat anti-human IgG (Jackson ImmunoResearch, #109007003) diluted 1:2000 in staining solution. Stained cells were analyzed on a BD LSR-II flow cytometry instrument. Data were analyzed with FlowJo_V10.

Western blot analysis - CD22, Syk, pCD22, pSyk and glyceraldehyde-3-phosphate dehydrogenase (GAPDH)-expression of Ramos cells before and after stimulation were analyzed via western blot analyses. Therefore, 4 million Ramos cells were lysed in 20 µl NP40 cell lysis buffer (Thermo Fisher Scientific) including 1× Protease/ Phosphatase Inhibitor Cocktail (100×; Cell Signaling Technology) via incubation for 30 min on ice. Before cell lysis, cells were either stimulated with 20 ug/ml CCP2 or anti-human IgG F(ab)'2 or incubated with stimulation medium (RPMI1640/100 U/ml P/S/GlutaMAX[™]/10 mM Hepes/2% FCS) only for 5 min at 37 °C. The supernatant of the lysed cells was stored at 20 °C for further usage. Lysates (10 μ l; 1 million cells) were subjected to 4 to 15% SDS-polyacrylamide gels (Bio Rad) together with $4 \times$ Laemmli buffer (Bio Rad) after incubation for 5 min at 95 °C. Subsequently, immunoblotting was performed on a nitrocellulose membrane (Bio Rad). Blots were incubated in PTE (3% skim milk powder/PBS/0.05% Tween) for 2 hours at RT. Following washing with PBS/0.05% Tween (PT), the blots were incubated at 4 °C overnight with 1:20,000 mouse anti-human GADPH (#MAD374, Bio Rad) or 1:3000 rabbit anti-human β-actin (D6A8, #8457S), 1:15,000 rabbit anti-human CD22 (#ab207727, BioLegend), and 1:300 rabbit anti-human Syk (#2712S, Cell Signaling Technology) or 1:200 rabbit anti-human pCD22 (Y822, #ab32123, BioLegend) and 1:1000 rabbit anti-human pSyk (p-Zap-70, Y319, #2717S, Cell Signaling Technology), respectively diluted in 5 ml of PTE. After washing with PT, blots were incubated for 1 hour at RT with 5 ml of PTE containing 1:1000 goat anti-rabbit- Ig HRP (#P0448, DAKO) and 1:5000 goat anti-mouse-Ig HRP (#P0447, DAKO) secondary antibodies. Blots were washed, and bound antibodies were visualized using enhanced chemiluminescence (GE Healthcare, RPN2109). The readout was performed on a Bio Rad Chemidoc Touch Imaging system. For biotinylated surface IgG BCR detection after Endo H or PNGase F treatment, samples were mixed with 4× Laemmli buffer, incubated for 5 min at 95 °C and loaded on 4 to 15% gels (Bio Rad). Western blot analysis was performed as described above using a goat anti-human IgG (#31410, Invitrogen) HRP labeled antibody diluted 1:1000 in PTE.

Spinning disk confocal microscopy – BCR modulation and antigen internalization were identified using spinning disk confocal microscopy. One million Ramos B cells were incubated for 30 min on ice followed by a 10 min stimulation at 4 °C with 5 ug/ml CCP2-strep. AF568 tetramers in PBS/2% FCS. The stimulated B cells were then incubated for 0, 5, or 15 min at 37 °C to allow BCR downmodulation and antigen internalization. After fixation with 2% paraformaldehyde (PFA), the cells were stained for 15 min with an AF647 NHS ester (Thermo Fisher Scientific, #A20006) labeled Fab goat anti-human IgG (Jackson ImmunoResearch, #109007003) antibody diluted 1:2000 in staining solution to detect the remaining BCRs on the fixed cell surface. After washing, the cells were settled to poly D-lysine precoated coverslips (Ibidi) for at least 40 min. Imaging was performed on an Andor Dragonfly 500 spinning disk confocal microscope at the Light and Electron Microscopy Facility of the LUMC, The Netherlands. Z-stacks of 20 slices per cell were acquired for three laser lines (488, 561, and 637), and the mean signal intensity on the B-cell membrane or inside the cell was assessed using Fiji/Image J.

Statistical analysis – Antigen-binding data of mAbs were analyzed using two-tailed unpaired t-tests assuming that all samples show the same scatter (SD). LC-MS mlgG BCR glycan data were analyzed using multiple t-tests assuming that all samples show the same SD and corrected for multiple comparisons using the Bonferroni-Dunn method. Paired two-tailed t-tests were performed for B-cell activation and BCR downmodulation assays with a number of 3 to 5 pairs. Ordinary one-way analysis of variance (ANOVA) tests were performed for parametric non-matched confocal microscopy data. Significant differences are denoted as follows: not significant (ns; p > 0.05), *(p < 0.05), **(p < 0.01), ***(p < 0.001) or ****(p < 0.001).

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Author contributions

All authors were involved in drafting the article or revising it critically for important intellectual content, and all authors approved the final version to be published. Conceptualization: T.K., C.G., L.H., T.W.J.H., H.U.S., M.R., R.H., and R.E.M.T. Methodology: T.K., C.G., L.M.V., L.H., L.M.S., M.C., G.J.M.P., and M.W. Software: C.G., L.M.V., and LH. Investigation: T.K., C.G., L.H., J.C.K., L.M.S., Y.H., K.A.J.V.S., R.D.V., A.S.B.K., S.R., G.S.-R., and B.X. Visualization: T.K., C.G., and L.H. Supervision: M.C., R.H., T.W.J.H., H.U.S., and R.E.M.T. Writing–original draft: T.K. and R.E.M.T. Writing–review and editing: C.G., L.D., J.C.K., L.M.S., M.C., Y.H., K.A.J.V.S., R.D.V., A.S.B.K., S.R., G.S.-R., L.M.V., H., B.X., G.J.M.P., M.W., T.R., T.W.J.H., H.U.S., M.R., and R.H.

Conflict of interest

C.G., B.X., T.R., and R.H. are named as coinventors of a patent application (PCT/EP2018/082236) regarding the arthritis-protective effects of ACPA. H.U.S., T.W.J.H., and R.E.M.T. are mentioned as inventors on a patent on ACPA IgG V-domain glycosylation. G.J.M.P. is mentioned as inventor on the CCP2-related patent EP2071335. The other authors declare that they have no competing interests.

Data and materials availability

We confirm that the data supporting the findings of this study are available within the article and/or the Supplementary Materials. The CCP2 peptide can be provided by G.J.M.P. (Radboud University, The Netherlands) pending scientific review and a completed material transfer agreement. Requests for the CCP2 peptide should be submitted to G.Pruijn@ncmls.ru.nl. The 7E4 sequence and antibodies can be provided by R.H. (Karolinska Institutet, Sweden) or T.R. (Sanquin Research, The Netherlands) under the protection of a completed material transfer agreement. Requests for the 7E4 sequence should be submitted to Rikard.Holmdahl@ki.se.

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Supplemental Tables

Table S1. SPR Kd1 and Kd2 values of monoclonal ACPA IgG with VDGs (WT) and w/o VDGs (NG) calculated based on biphasic curve fits.

Kd1	CCP2	cit-fibrinogen α 27-43	cit-fibrinogen β 36-52	
7E4 WT	8.80E-05	nd*	7.14E-05	
7E4 WT+neu	2.89E-04	6.10E-04	1.19E-04	
7E4 NG	9.36E-05	7.33E-05	nd*	
7E4 NG+neu	nd*	7.93E-04	nd*	
Kd2	CCP2	cit-fibrinogen α 27-43	cit-fibrinogen β 36-52	
7E4 WT	3.74E-05	2.22E-04	nd*	
7E4 WT+neu	nd*	1.39E-04	nd*	
7E4 NG	1.94E-04	4.36E-05	nd*	
7E4 NG+neu	5.39E-05	6.06E-05	nd*	

*nd = not-determined, negative Kon values.

Table S2. Equilibrium $\rm K_{\rm p}$ values of monoclonal ACPA IgG with VDGs (WT) and without VDGs (NG).

Equilibrium $K_{\rm D}$ and $B_{\rm max}$ values were calculated based on ELISA curves and shown including their standard errors.

peptide	7E4 WT		7E4 WT + neu		
	K _p [M]	B _{max}	К _р [М]	B _{max}	
CCP2	1.04E-08 ± 6.56E-10	2.54 ± 0.048	6.78E-09 ± 5.02E-10	2.81 ± 0.051	
Cit-fibrinogen a 27-43	6.64E-09 ± 5.67E-10	3.20 ± 0.066	5.26E-09 ± 6.84E-10	3.48 ± 0.097	
CCP1	NM*	NM*	NM*	NM*	
Cit-vimentin 59-74	NM*	NM*	NM^*	NM*	
Cit-enolase 5-20	NM*	NM*	NM^*	NM*	
Cit-fibrinogen β 36-52	1.30E-08 ± 3.67E-10	2.11 ± 0.022	9.24E-09 ± 3.54E-10	2.31 ± 0.029	

*NM = not-measurable, ELISA curves not at saturation.

cit-vimentin 59-74	cit-enolase 5-20	Kd1	CCP2
1.78E-05	3.72E-05	2D11 WT	1.21E-04
2.90E-05	5.18E-05	2D11 WT+neu	7.82E-05
4.83E-04	3.99E-03	2D11 NG	5.94E-05
3.96E-05	nd*	2D11 NG+neu	7.53E-05
cit-vimentin 59-74	cit-enolase 5-20	Kd2	CCP2
1.41E-04	3.20E-04	2D11 WT	nd*
nd*	4.60E-03	2D11 WT+neu	5.75E-05
nd*	nd*	2D11 NG	7.96E-05
nd*	nd*	2D11 NG+neu	nd*

7E4 M	١G	7E4 NG + neu		
K _D [M]	B _{max}	K _p [M]	B _{max}	
2.66E-09 ± 4.77E-10	3.25 ± 0.082	2.88E-09 ± 2.29E-10	3.16 ± 0.037	
2.83E-09 ± 3.47E-10	3.43 ± 0.062	2.32E-09 ± 3.47E-10	3.29 ± 0.063	
7.16E-08 ± 6.38E-09	4.83 ± 0.280	5.82E-08 ± 6.91E-09	4.11 ± 0.298	
4.91E-08 ± 9.56E-09	4.44 ± 0.497	4.76E-08 ± 1.03E-08	4.51 ± 0.553	
1.73E-07 ± 2.10E-08	5.39 ± 0.453	1.81E-07 ± 1.92E-08	5.54 ± 0.413	
5.53E-09 ± 6.03E-11	2.70 ± 0.009	5.12E-09 ± 1.81E-10	2.61 ± 0.028	

	3F3:cit-vimentin 59-74	
Wavelength [Å]	0.91840	
Resolution range [Å]	52.28 - 2.0 (2.071 - 2.0)	
Space group	P 21 21 21	
Unit cell a, b, c [Å]	53.379, 82.094, 135.613	
Unit cell a, b, c [Å]	90, 90, 90	
Total reflections	325932 (33061)	
Unique reflections	40868 (4035)	
Multiplicity	8.0 (8.2)	
Completeness [%]	99.37 (99.51)	
Mean I/sigma [I]	9.93 (2.16)	
Wilson B-factor	32.61	
R-merge	0.1256 (0.886)	
R-meas	0.1344 (0.9457)	
R-pim	0.04728 (0.3275)	
CC1/2	0.996 (0.682)	
CC*	0.999 (0.901)	
Reflections used in refinement	40845 (4025)	
Reflections used for R-free	2023 (193)	
R-work	0.2011	
R-free	0.2414	
Number of non-hydrogen atoms	3762	
macromolecules	3395	
ligands	64	
solvent	303	
Protein residues	447	
RMS [bonds]	0.019	
RMS [angles]	2.39	
Ramachandran favored [%]	97.00	
Ramachandran allowed [%]	3.00	
Ramachandran outliers [%]	0.00	
Number of TLS groups	3	

Table S3. Data collection and refinement statistics. Statistics for the highest-resolution shell are shown in parentheses.

1F2:cit-CII-C-39	1F2	7E4:cit-CII-C-48
0.97622	0.97622	0.91840
36.19 - 2.85 (2.952 - 2.85)	35.01 - 3.05 (3.159 - 3.05)	59.78 - 2.45 (2.538 - 2.45)
P 21 21 21	C121	C 2 2 21
53.01, 89.65, 118.68	152.43, 88.8, 88.02	77.183, 151.774, 97.066
90, 90, 90	90, 113.26, 90	90, 90, 90
121917 (11635)	95969 (9446)	195694 (19513)
13784 (1351)	20678 (2032)	21351 (2104)
8.8 (8.5)	4.6 (4.6)	9.2 (9.3)
98.10 (98.69)	99.63 (99.61)	99.96 (99.86)
9.97 (3.66)	9.92 (3.31)	22.47 (4.80)
44.66	77.47	57.62
0.1524 (0.5557)	0.09467 (0.3793)	0.05246 (0.4131)
0.1623 (0.5933)	0.1073 (0.4305)	0.05567 (0.438)
0.05479 (0.2044)	0.04964 (0.2002)	0.01827 (0.1431)
0.992 (0.901)	0.993 (0.938)	0.999 (0.951)
0.998 (0.974)	0.998 (0.984)	1 (0.987)
13532 (1351)	20638 (2032)	21348 (2104)
638 (74)	1054 (102)	1129 (106)
0.2117	0.2349	0.2384
0.2610	0.2663	0.2746
3372	6444	3363
3275	6392	3276
64	52	39
33		48
434	848	440
0.013	0.012	0.013
1.77	1.84	1.89
95.01	96.12	96.69
4.75	3.03	3.31
0.24	0.85	0.00
3	4	3

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Table S4. H-bond interaction details between the ACPA 7E4 HC and LC Fab fragments and the respective N-linked glycan. Shown are the H-bond donor and acceptor pairs (three letter code of the respective amino acids and monosaccharides) and the occupancy of the interaction in percentage. GlcNAc: *N*-acetyl glucosamine, Neu5Ac: *N*-acetylneuraminic acid, Fuc: fucose, Gal: galactose, Man: mannose.

	7E4 HC			7E4 LC	
donor	acceptor	oc. [%]	donor	acceptor	oc. [%]
Asn 29	GlcNAc 221	5.85	Asn 16	GlcNAc 214	10.76
GlcNAc 221	Pro 73	1.15	Fuc 226	Gly 14	8.84
Ser 74	GlcNAc 222	0.94	Fuc 226	Asn 16	2.44
GlcNAc 222	Ser 74	0.84	Fuc 226	GIn 15	1.56
Ser 74	GlcNAc 222	0.49	GIn 15	Fuc 226	1.36
Fuc 233	Asn 29	0.49	Arg 52	Neu5Ac 220	1.04
His 53	Fuc 233	0.36	Man 217	Ala 60	0.9
Fuc 233	His 53	0.35	Arg 52	Neu5Ac 220	0.88
Lys 75	GlcNAc 225	0.35	Neu5Ac 220	Asn 66	0.73
GlcNAc 222	Ser 74	0.35	Neu5Ac 220	Ser 65	0.55
Lys 75	GlcNAc 232	0.32	GIcNAc 214	Asp 74	0.42
Met 28	GlcNAc 221	0.29	Arg 52	GlcNAc 218	0.37
Lys 75	GlcNAc 229	0.22	Arg 54	Man 217	0.35
Asn 76	GlcNAc 221	0.22	Gly 77	Fuc 226	0.31
Asn 76	GIcNAc 222	0.19	Man 217	Ser 63	0.28
GlcNAc 222	Asn 76	0.14	Ser 63	Man 217	0.28
Gal 226	Thr 22	0.14	Arg 54	Neu5Ac 220	0.26
GIn 77	Gal 226	0.11	Neu5Ac 220	Ser 63	0.25
Thr 22	Gal 226	0.1	Neu5Ac 220	Ser 63	0.21
Ser 20	Gal 226	0.04	Neu5Ac 220	lle 58	0.21
GlcNAc 232	Lys 75	0.04	Neu5Ac 220	Gly 57	0.17
GlcNAc 232	Ser 74	0.04	Arg 54	Gal 219	0.16
Gal 226	Ser 74	0.02	Ala 60	GlcNAc 218	0.11
Gal 226	Lys 75	0.02	Ser 56	Neu5Ac 220	0.08
Ser 74	GlcNAc 221	0.01	GlcNAc 215	Ala 60	0.08
Fuc 233	His 53	0.01	lle 58	Neu5Ac 220	0.06
Lys 75	Neu5Ac 231	0.01	Neu5Ac 220	Ser 56	0.05
His 53	GlcNAc 221	0.01	Ala 60	Man 217	0.05
GlcNAc 221	Ser 74	0.01	Neu5Ac 220	Ser 56	0.05
Neu5Ac 231	Thr 56	0.01	Arg 61	Man 217	0.04
Gal 230	Thr 54	0.01	Gal 223	GIn 79	0.04
Glu 30	GlcNAc 221	0.01	Arg 54	GlcNAc 218	0.04
His 53	Gal 230	0.01	GlcNAc 215	Ala 76	0.03
			Neu5Ac 224	Gly 14	0.03
			Neu5Ac 224	GIn 15	0.03

	7E4 HC			7E4 LC	
donor	acceptor	oc. [%]	donor	acceptor	oc. [%]
			Ser 56	Gal 219	0.02
			Gal 219	Ser 56	0.01
			GIn 79	Neu5Ac 224	0.01
			Neu5Ac 224	GIn 79	0.01
			Neu5Ac 224	Pro 13	0.01
			Arg 61	Man 217	0.01
			Arg 80	Neu5Ac 224	0.01
			Neu5Ac 224	Pro 13	0.01
			GIn 15	Neu5Ac 224	0.01
			Neu5Ac 220	Gly 57	0.01
			Gly 57	Neu5Ac 220	0.01
			GIcNAc 214	Ala 76	0.01
			Gal 223	Pro 13	0.01

Table S4. Continued

Table S5. H-bond interaction details between the ACPA 3F3 HC and LC Fab fragments and the respective N-linked glycan. Shown are the H-bond donor and acceptor pairs (three letter code of the respective amino acids and monosaccharides) and the occupancy of the interaction in percentage. GlcNAc: *N*-acetylglucosamine, Neu5Ac: *N*-acetylneuraminic acid, Fuc: fucose, Gal: galactose, Man: mannose.

3F3 H	IC #1 N-glycar	n	3F3 H	IC #2 N-glycar	ı		3F3 LC		
donor	acceptor	oc.	donor	acceptor	oc.	donor	acceptor	oc.	
		[%]			[%]			[%]	
Fuc 236	Glu 81	13.02	Fuc 249	Glu 22	10.53	Ser 71	GlcNAc 219	15.43	
Lys 18	Fuc 236	1.68	Fuc 249	Ser 74	4.57	Asn 78	GIcNAc 219	8.69	
Fuc 236	Thr 70	1.21	Gal 246	Gly 25	2.00	GlcNAc 220	Asp 76	5.96	
Fuc 236	GIn 64	0.68	Phe 28	GlcNAc 237	0.47	GIcNAc 219	Asp 76	4.23	
GlcNAc 224	Glu 81	0.65	GlcNAc 237	Ser 24	0.41	Lys 24	GlcNAc 227	1.18	
Thr 70	Fuc 236	0.26	Gal 246	Ser 24	0.21	Gal 228	Ser 26	0.96	
Neu5Ac 230	Gly 65	0.24	Neu5Ac 243	Tyr 100	0.08	Lys 24	GlcNAc 220	0.54	
Asn 68	GlcNAc 224	0.21	Lys 53	Man 240	0.03	Lys 24	Man 226	0.50	
Fuc 236	Asn 68	0.14	Asn 29	GlcNAc 237	0.03	Thr 20	Fuc 231	0.34	
Neu5Ac 230	GIn 64	0.12	Lys 53	GlcNAc 238	0.03	GlcNAc 227	Thr 75	0.31	
GIn 64	Fuc 236	0.06	Neu5Ac 247	Gly 25	0.03	Neu5Ac 229	Asp 9	0.28	
Met 69	GlcNAc 224	0.04	Tyr 100	Neu5Ac 243	0.03	Fuc 231	Thr 20	0.26	
Lys 53	Neu5Ac 234	0.03	Lys 53	Neu5Ac 243	0.02	Neu5Ac 229	Asp 9	0.21	
Lys 53	Gal 229	0.02	lle 27	Gal 246	0.01	GIcNAc 219	Asp 76	0.19	
Neu5Ac 230	Gly 14	0.02	Tyr 100	Neu5Ac 243	0.01	Ser 28	Gal 228	0.17	
Fuc 236	Gly 65	0.02				Fuc 231	Thr 20	0.14	

3F3	HC #1 N-glycan	l	3F3	HC #2 N-glyca	n		3F3 LC	
donor	acceptor	oc.	donor	acceptor	oc.	donor	acceptor	oc.
		[%]			[%]			[%]
Lys 53	Neu5Ac 230	0.01				Neu5Ac 225	Ser 26	0.14
Thr 83	Neu5Ac 230	0.01				GIcNAc 227	Ser 25	0.14
GlcNAc 224	Thr 83	0.01				Thr 75	GIcNAc 227	0.13
Thr 70	Neu5Ac 234	0.01				GIcNAc 227	Asp 76	0.08
						Neu5Ac 225	Ser 73	0.06
						Arg 18	Fuc 231	0.05
						Gal 228	Tyr 7	0.05
						Gal 228	Gln 27	0.05
						Asp 9	Gal 228	0.04
						Neu5Ac 229	Tyr 7	0.03
						Ser 73	Neu5Ac 225	0.03
						Tyr 7	Neu5Ac 229	0.02
						Neu5Ac 225	Gln 27	0.02
						Tyr 7	Gal 228	0.02
						Fuc 231	Asn 22	0.02
						Tyr 7	GlcNAc 227	0.02
						Gal 228	Thr 75	0.02
						Thr 5	Neu5Ac 229	0.01
						Neu5Ac 225	Thr 75	0.01
						Ser 28	Neu5Ac 225	0.01
						Ser 73	Neu5Ac 225	0.01
						Gal 228	Asp 9	0.01
						Gln 27	Gal 228	0.01
						Thr 75	Gal 228	0.01
						Neu5Ac 225	Ser 71	0.01

Table S5. Continued

Table S6. H-bond interaction details between the ACPA 1F2 HC and LC Fab fragments and the respective N-linked glycan. Shown are the H-bond donor and acceptor pairs (three letter code of the respective amino acids and monosaccharides) and the occupancy of the interaction in percentage. GlcNAc: *N*-acetylglucosamine, Neu5Ac: *N*-acetylneuraminic acid, Fuc: fucose, Gal: galactose, Man: mannose.

······································										
1F2 H	IC #1 N-glycan	1	1F2 H	IC #2 N-glycan	1F2 LC					
donor	nor acceptor oc.		donor	acceptor	oc.	donor	acceptor	oc.		
		[%]			[%]			[%]		
Ser 30	GlcNAc 223	11.08	GlcNAc 236	Thr 55	4.27	Gal 222	Asp 151	21.55		
His 31	Fuc 235	9.5	Lys 54	Nau5Ac 246	2.7	Neu5Ac 223	Gly 156	10.44		
Fuc 235	His 31	2.4	Fuc 248	Ala 56	1.65	GlcNAc 224	Thr 160	10.39		

1F2 H	C #1 N-glycan		1F2 H	C #2 N-glycan		1F2 LC		
donor	acceptor	oc.	donor	acceptor	oc.	donor	acceptor	oc.
		[%]			[%]			[%]
Lys 54	Fuc 235	1.98	Thr 55	GlcNAc 236	0.93	Neu5Ac 219	Gln 161	9.88
Fuc 235	Ser 30	1.32	Fuc 248	Thr 55	0.86	Lys 153	Gal 222	7.29
Neu5Ac 233	Leu 102	1.14	Neu5Ac 246	Lys 54	0.74	Gal 218	Thr 157	5.81
Ser 30	GlcNAc 223	1.13	Asn 57	GlcNAc 236	0.22	Gln 161	GlcNAc 224	4.83
Phe 29	GlcNAc 223	0.89	Gal 245	Asn 57	0.2	Neu5Ac 219	Thr 183	3.81
Thr 74	GlcNAc 224	0.25	GlcNAc 236	Ala 56	0.15	GlcNAc 213	Thr 3	2.83
lle 104	Fuc 235	0.13	Fuc 248	Asn 57	0.12	Thr 157	Neu5Ac 223	2.75
Fuc 235	Asn 28	0.09	Lys 54	Neu5Ac 246	0.1	Gal 218	Gln 161	2.48
Asn 28	GlcNAc 223	0.05	Thr 55	Fuc 248	0.1	Fuc 225	Asn 25	2.15
GlcNAc 231	Lys 54	0.04	Thr 55	Gal 245	0.07	Neu5Ac 223	Gly 156	1.85
GlcNAc 224	Thr 74	0.03	Neu5Ac 246	Thr 55	0.03	Neu5Ac 223	Thr 157	1.74
Fuc 235	Leu 102	0.02	Gal 245	Lys 54	0.02	Asn 1	GlcNAc 213	1.59
Lys 54	Gal 232	0.02	Leu 58	Fuc 248	0.01	Gln 161	Neu5Ac 219	1.42
His 31	GlcNAc 223	0.01	Thr 55	GlcNAc 236	0.01	Gal 218	Gln 161	1.34
GlcNAc 224	Ser 75	0.01				Gln 161	GlcNAc 217	1.16
Neu5Ac 233	Gly 103	0.01				GlcNAc 224	Gln 161	1.04
						Thr 160	GlcNAc 224	1.02
						GlcNAc 217	Gln 161	0,85
						Gal 222	Gly 156	0.84
						Fuc 225	Asn 25	0.81
						Gln 161	GlcNAc 224	0.62
						Lys 153	Neu5Ac 223	0.55
						Lys 133	Neu5Ac 219	0.53
						Neu5Ac 223	Asp 155	0.49
						Neu5Ac 219	Gln 161	0.46
						Fuc 225	Asn 1	0.42
						Neu5Ac 219	Thr 183	0.39
						Thr 205	Neu5Ac 223	0.39
						GlcNAc 217	Gln 161	0.34
						Fuc 225	Thr 3	0.27
						Thr 160	GlcNAc 224	0.27
						Thr 157	Gal 218	0.26
						Gal 218	Thr 160	0.23
						Gln 161	Gal 218	0.22
						GlcNAc 217	Thr 160	0.22
						Thr 160	Gal 218	0.2

Table S6. Continued

	1F2 HC #1 N-gl	lycan		1	F2 HC	#2 N-glya	an		1F2 LC		
donor	accept	or	oc.	dono	r	acceptor		oc.	donor	acceptor	oc.
			[%]					[%]			[%]
									Thr 3	Fuc 225	0.18
									GlcNAc 213	Leu 2	0.17
									Fuc 225	Asn 1	0.16
									Gly 162	Neu5Ac 219	0.15
									GIcNAc 224	Pro 158	0.15
									Neu5Ac 223	Gly 203	0.15
									Neu5Ac 223	Asp 155	0.14
									Gal 222	Thr 157	0.13
									GlcNAc 221	Asp 151	0.13
									Thr 185	Neu5Ac 219	0.13
									Asn 25	Fuc 225	0.1
									Gal 222	Thr 205	0.08
									Gln 198	Gal 222	0.07
									Neu5Ac 223	Thr 157	0.05
									Arg 191	Neu5Ac 219	0.05
									GlcNAc 221	Thr 160	0.05
									Thr 160	GlcNAc 217	0.04
									Thr 183	Neu5Ac 219	0.03
									GlcNAc 217	Thr 160	0.03
									Neu5Ac 223	Glu 207	0.03
									Gln 161	Man 216	0.02
									Gln 161	GlcNAc 217	0.02
									Thr 205	Neu5Ac 223	0.02
									Thr 185	Neu5Ac 219	0.02
									Gal 222	Glu 207	0.02
									Thr 183	Neu5Ac 219	0.01
									Thr 160	GlcNAc 221	0.01
									Gal 218	Pro 158	0.01
									Gal 218	His 192	0.01
									Neu5Ac 219	Thr 185	0.01
									Gal 218	Val 154	0.01
									Thr 157	GlcNAc 224	0.01
									Fuc 225	Gly 26	0.01
									Thr 157	Neu5Ac 223	0.01

Table S6. Continued



Supplemental Figures

Figure S1. MALDI-TOF analysis of released and stabilized N-linked variable domain and Fc glycans from WT (blue), WT+neu (light blue), NG (red) and NG+neu (light red) ACPA IgG 2E4, 1F2, 2G9 and 2D11. Glycan structures of the most abundant N-linked glycan peaks are depicted. Blue square: *N*-acetylglucosamine (GlcNAc), green circle: mannose, yellow circle: galactose, red triangle: fucose, purple diamond: α2,6-linked *N*-acetylneuraminic acid.



Figure S2. Molecular dynamics (MD)-simulations to predict VDG-antibody interactions and 7E4_{Fab} crystal structure comparison with and without N-linked glycan sites. (A) MD-simulations, 8 time-points, of 7E4_{Fab}, 3F3_{Fab} and 1F2_{Fab} crystal structures modelled with full length disialylated VDG (sticks in magenta). (B) Superposition of 7E4_{Fab} crystal structures expressed with and without N-linked glycan sites. 7E4_{Fab} WT including N-linked glycan sites was crystallized together with the first two GlcNAcs (sticks in magenta) of the LC VDG. LC is depicted in light gray, HC in steel blue, the peptides bound to the respective Fab as sticks with carbon (green), oxygen (red) and nitrogen (blue) atoms. (C) ACPA 7E4_{Fab} and 1F2_{Fab} HC and LC variable gene aa-sequences, based on IMGT (2D), are depicted. Acceptor/ donor pairs of H-bond interactions between the antibody and HC (#1 and #2)/LC N-glycans are visualized. A high occupancy is depicted in dark blue (HC: 6%; 12%, LC: 11%; 22%) and a low occupancy is depicted in light blue (0.01 to 0.2%). Amino acids and their respective interaction partners (N-glycan monosaccharides) are shown. Blue square: *N*-acetylglucosamine, green circle: mannose, yellow circle: galactose, red triangle: fucose, purple diamond: α 2,6-linked *N*-acetylneuraminic acid (sialic acid).



Figure S3. Impact of disialylated ACPA VDG on citrullinated peptide binding. ELISA titration binding curves of the ACPA 2E4, 1F2, 2D11, 2G9 and 3F3 (0 to 40 μ g/ml) variants (WT, WT+neu, NG and NG+neu) towards citrullinated peptides (CCP2, CCP1, cit-fibrinogen α 27-43, cit-fibrinogen β 36-52, cit-vimentin 59-74, cit-enolase 5-20). Binding to the arginine control peptide was subtracted. Reactivity was determined via the OD at 415 nm. Each data point represents the mean of two technical replicates. N = 2 to 3.



Figure S4. Impact of disialylated ACPA VDG on citrullinated protein binding. (A) ELISA titration binding curves of the ACPA 2E4, IF2, 2D11, 2G9 and 3F3 (0 to 40 μ g/ml) variants (WT, WT+neu, NG and NG+neu) towards citrullinated proteins (cit-fibrinogen, cit-vinculin, cit-MBP and cit-OVA). Binding to the arginine control protein was subtracted. Reactivity was determined via the OD at 415 nm. N = 2 to 3. (B) Relative binding of the ACPA mAb 2E4, 2G9 and 2D11 (10 to 40 μ g/ml) variants (WT, WT+neu, NG, NG+neu) towards citrullinated peptides and proteins. N = 2 to 3. Unpaired two-tailed t-tests assuming the same SD. 2E4WT-NG: **p (CCP1) = 0.0028; 2E4WT-WT+neu: *p (CCP1) = 0.0143, *p (cit-fibrinogen) = 0.0226; 2G9WT-NG: ***p (cit-fibrinogen) = 0.0009, **p (cit-OVA) = 0.0047; 2G9WT-WT+neu: **p (cit-fibrinogen) = 0.0004, *p (cit-OVA) = 0.0297; 2D11WT-NG: *p (CCP2) = 0.0406.



Figure S5. SPR sensorgrams of ACPA IgG 7E4 and 2D11 binding to citrullinated peptides. Antigen association and dissociation are represented as response units (RU) over time (s). The experiment represents 2 technical replicates. Four different monoclonal antibody variants were assessed: WT, WT+neu, NG and NG+neu.



Figure S6. Human Ramos IgG B-cell receptor glycan analysis. (A) Absolute intensity of glycan traits expressed on the 7E4 WT, NG and MDL IgG BCR after passing QC settings. (B) LC-chromatogram of glycan traits expressed on the 7E4 WT and NG (IgG BCR after passing QC settings. The respective glycan traits are schematically illustrated. (C) Western blot analyses of PBS-treated (- Biotin) or biotinylated (+ Biotin) surface IgG 3F3 WT and NG BCRs after NeutrAvidin capturing. Biotinylated mIgG with and without VDG were treated with 2 U Endo H (cleavage of high-mannose structures) or 2 U PNGase F (cleavage of all N-glycans).



Figure S7. Impact of mIgG VDGs on human Ramos B-cell activation and BCR downmodulation. (**A**) Paired analysis of Ca²⁺-flux speed (slope) for mIgG 3F3/ 7E4 WT and NG Ramos B cells after a-IgG F(ab)'2 and antigen (CCP2-strep.) stimulation. Paired two-tailed t-test. N = 5 to 7. 3F3: *p = 0.0377; 7E4: *p = 0.0206. (**B**) Overlays of WT and NG Ca²⁺-flux (Ca2+ bound Indo-1/ unbound Indo-1) of mIgG 3F3/ 7E4 WT and NG Ramos B cells after stimulation with CArgP2-SA or cit-fibrinogen β 36-52. (**C**) Western blot analyses of mIgG 3F3/ 7E4 WT and NG Ramos B cells after 5 min of CCP2 stimulation or unstimulated (us). CD22, Syk, pCD22 (Y822) and pSyk(Y352) expression are shown. Cell lysis of 1 million (unstimulated and stimulated 1st slot), 0.5 million (stimulated 2nd slot) and 0.25 million (stimulated 3rd slot) were blotted. GAPDH was used as loading control. Cell lysates of 1 million MDL-AID KO cells were added as additional control. (**D**) Remaining mIgG 7E4 WT and NG expression after CCP2-strep. stimulation and incubation at 4°C or 5, 15 min at 37 °C. N = 619, 459, 645, 433, 738 and 302 cell slices respectively. Ordinary one-way ANOVA, ***p = 0.0002.



Figure S8. TIDE analysis 3F3 CD22KO Ramos B cells. (**A**) Comprehensive profile of all insertions and deletions (indels) in the CD22 KO edited sample according to the TIDE analysis. The total KO efficiency is depicted. (**B**) An estimate of the +1 insertion base composition according to the TIDE analysis. (**C**) pSyk(Y348) expression of mlgG 3F3 WT, WT CD22KO, NG and NG CD22KO BCRs after 5, 10 and 15 min of a-IgG F(ab)'2 stimulation. Gating is based on the unstimulated cells. The pSyk expression is depicted on the y-axis and the GFP expression on the x-axis. One experiment is exemplarily shown.