Stereotyped antibody responses target posttranslationally modified gluten in celiac disease

Omri Snir,1 Xi Chen,1 Moriah Gidoni,2 M. Fleur du Pré,1 Yuguang Zhao,3 Øyvind Steinsbø,1 Knut E.A. Lundin,4,5 Gur Yaari,2 and Ludvig M. Sollid1,5

1Centre for Immune Regulation and Department of Immunology, University of Oslo and Oslo University Hospital–Rikshospitalet, Oslo, Norway. 2Bioengineering Faculty of Engineering, Bar-Ilan University, Ramt Gan, Israel. 3Division of Structural Biology, University of Oxford, Wellcome Trust Centre for Human Genetics, Headington, Oxford, United Kingdom. 4Department of Gastroenterology, Oslo University Hospital–Rikshospitalet, Oslo, Norway. 5KG Jebsen Coeliac Disease Research Centre, University of Oslo, Oslo, Norway.

Introduction

Celiac disease (CD) is an immune-mediated disorder with autoimmune features caused by immune reactions to dietary gluten proteins from wheat (gliadin and glutenin), barley (hordein), and rye (secalin) (1). CD has strong genetic basis, and the MHC locus with certain HLA-DQA1 and HLA-DQB1 alleles encoding HLA-DQ2 and HLA-DQ8 molecules is the chief genetic risk factor (2). Hallmarks of the disease are gluten-reactive CD4 T cells that recognize deamidated gluten peptides in the context of the HLA-DQ2 or HLA-DQ8 molecules (3), as well as antibodies to the autoantigen transglutaminase 2 (TG2) (4) and to deamidated gluten peptides (5). Both the anti-TG2 antibodies and the anti-deamidated gluten peptide antibodies are diagnostic markers for CD with extraordinary disease specificity and sensitivity (6, 7). TG2 is not only involved in CD by being the target of autoantibodies, but it is also responsible for the deamidation of gluten peptides rendering them immunogenic for CD4 T cells (8, 9). The TG2 enzyme targets Gln residues in polypeptides in a sequence-specific manner (i.e., QXP[F]) and converts them to Glu (deamidation) or crosslinks them to primary amines (transamidation) (10–12). The T cell response to deamidated gluten peptides and the B cell response to TG2 and deamidated gluten peptides are likely mechanistically linked. Both B cells reactive with deamidated gluten and B cells reactive with TG2, the latter via B cell receptor–mediated (BCR-mediated) uptake of TG2-gluten peptide complexes, may serve as antigen presenting cells for gluten-reactive CD4 T cells and thereby engage an amplifying loop for the pathogenic T cell response (3, 13).
results of antibody recognition were used to reinterrogate the HTS sequence data, which revealed selection of a stereotypic antibody response to DGP (23). DGP-reactive PCs had significantly lower numbers of somatic mutations (Figure 1B), as well as lower clonal diversity in comparison with non–DGP-reactive PCs (likely reactive with gut commensals and pathogens) from 10 untreated CD patients. Single cell suspensions of small intestine biopsies were stained and analyzed by flow cytometry. The frequency of DGP-reactive IgA PCs ranged between 0.2% and 2%. DGP-reactive PCs were sorted, and amplicon libraries were prepared, sequenced on Illumina MiSeq platform, and analyzed. The IGHV repertoire of DGP-reactive PCs was relatively narrow, with a preferential usage of the IGHV3-15 and IGHV3-23 gene segments (18). We then solved the structural basis for recognition of the peptide by 2 prototype antibodies encoded by IGHV3-23/IGLV4-69 and IGHV3-15/IGKV4-1 genes (18). Additionally, pull-down and characterization of peptides from complex proteolytic digests of gluten treated with TG2 suggested that the core epitope–recognized -QPEQPF- is only found within or in proximity of many different CD4 T cell epitopes in gluten proteins (21). This suggests that the B cell response toward dietary gluten in CD is connected to the antiglutens T cell response and that the response to DGP plays a central role in the immune reaction to deamidated gluten.

In this study, we have further dissected the CD-specific antibody response to DGP. We first did high-throughput sequencing (HTS) of IGHV genes of PCs from 10 CD patients with active disease. We observed that PCs reactive with the DGP had fewer mutations in the IGHV genes than PCs not reacting with this peptide. These gluten-specific PCs were clonally expanded with a restricted IGHV repertoire and over-usage of the IGHV3-15 and IGHV3-23 gene segments. We then solved the structural basis for recognition of the peptide by 2 prototype antibodies encoded by IGHV3-23/IGLV4-69 (hmAb UCD1002-1E01) and IGHV3-15/IGKV4-1 (hmAb UCD1002-1E03), hereafter termed 1E01 and 1E03, respectively. The structural insight of antibody recognition was used to reinterrogate the HTS sequence data, which together provide insight into the scheme by which CD-specific antibodies are selected in response to an immunodominant and deamidated gluten epitope. Interestingly both antibodies, despite different epitope interaction, recognize signatures of TG2 processing that would allow both B cells to take up and present deamidated gluten peptides to T cells. The study helps to understand, at the molecular level, how the antibody response to post-translationally modified gluten peptides develop in CD.

Results

HTS reveals selection of a stereotypic antibody response to DGP

In order to comprehensively analyze the antibody response to DGP, we undertook HTS of IGHV genes of DGP-reactive and non–DGP-reactive PCs (likely reactive with gut commensals and pathogens) from 10 untreated CD patients. Single cell suspensions of small intestine biopsies were stained and analyzed by flow cytometry. The frequency of DGP-reactive IgA PCs ranged between 0.2% and 2%. DGP-reactive and non–DGP-reactive IgA PCs were sorted, and amplicon libraries were prepared, sequenced on Illumina MiSeq platform, and analyzed. The IGHV repertoire of DGP-reactive PCs was relatively narrow, with a preferential usage the IGHV3-23 and IGHV3-15 gene segments (Figure 1A). There was some variation between individual patients; most patients predominantly used IGHV3-23 (as depicted in Supplemental Figure 1A; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.93961DS1); however, in one subject (CD1195), IGHV3-15 strongly dominated the response (Supplemental Figure 1B). Furthermore, DGP-reactive PCs had significantly lower numbers of somatic mutations (Figure 1B), as well as lower clonal diversity in comparison with non–DGP-reactive PCs, as measured by Hill’s diversity index (Figure 1C; ref. 22). The lower diversity (D) values observed for DGP-reactive PCs indicated that this population consisted of clones that were larger than non–DGP-reactive PCs. The finding of consistent and biased usage of IGHV3-23 and IGHV3-15 across CD patients confirms and extends the results obtained by analysis of 38 single celiac gut PCs (18) and clearly demonstrates that there exists a stereotyped antibody response to DGP in CD.

antigens in rheumatoid arthritis (15, 16). Still, many issues remain unsolved. In rheumatoid arthritis, telling-ly, the question of which enzyme is involved in the citrullination (i.e., bacterial or human peptidylarginine deiminases) and which T cell epitopes are involved in the antibody formation are still elusive (17). The work from CD, including this report on the detailed molecular basis for antibody recognition of posttranslationally modified antigen, illustrates how critical insight can be obtained.

We recently isolated plasma cells (PCs) of celiac gut lesions reactive with deamidated gluten peptides, and we generated from such cells a panel of 38 human monoclonal antibodies (hmAbs) (18). Many of the hmAbs were derived from PCs that were selected with a peptide containing the sequence PLQPEQPFF (hereafter termed DGP) that harbos the sequence QPEQPF; a previously identified highly disease-specific and immunodominant B cell epitope in CD (5). In general, the 38 gluten-reactive hmAbs expressed IGHV genes with relatively few somatic mutations — a characteristic shared with TG2 antibodies of CD patients (19, 20). Furthermore, among the 38 hmAbs, there was a restricted usage of VH and VL pairs, with PLQPEQPFF-reactive hmAbs typically using the IGHV3-23/IGLV4-69 and IGHV3-15/IGKV4-1 genes (18). Additionally, pull-down and characterization of peptides from complex proteolytic digests of gluten treated with TG2 suggested that the core epitope–recognized -QPEQPF- is only found within or in proximity of many different CD4 T cell epitopes in gluten proteins (21). This suggests that the B cell response toward dietary gluten in CD is connected to the antiglutens T cell response and that the response to DGP plays a central role in the immune reaction to deamidated gluten.

In this study, we have further dissected the CD-specific antibody response to DGP. We first did high-throughput sequencing (HTS) of IGHV genes of PCs from 10 CD patients with active disease. We observed that PCs reactive with the DGP had fewer mutations in the IGHV genes than PCs not reacting with this peptide. These gluten-specific PCs were clonally expanded with a restricted IGHV repertoire and over-usage of the IGHV3-15 and IGHV3-23 gene segments. We then solved the structural basis for recognition of the peptide by 2 prototype antibodies encoded by IGHV3-23/IGLV4-69 (hmAb UCD1002-1E01) and IGHV3-15/IGKV4-1 (hmAb UCD1002-1E03), hereafter termed 1E01 and 1E03, respectively. The structural insight of antibody recognition was used to reinterrogate the HTS sequence data, which together provide insight into the scheme by which CD-specific antibodies are selected in response to an immunodominant and deamidated gluten epitope. Interestingly both antibodies, despite different epitope interaction, recognize signatures of TG2 processing that would allow both B cells to take up and present deamidated gluten peptides to T cells. The study helps to understand, at the molecular level, how the antibody response to post-translationally modified gluten peptides develop in CD.
Figure 1. *IGHV* repertoire, frequency of somatic mutation, and clonal diversity of DGP-reactive PCs. (A) Bars represent the mean frequencies ± SD of the most abundantly expressed *IGHV* gene segments by DGP-reactive PCs and non-DGP-reactive PCs in 10 CD patients with active disease. (B) Mutation frequency in the *IGHV* genes of non-DGP-reactive and DGP-reactive PCs from 10 CD patients. White dots indicate the median frequency of mutation with 1.5 interquartile range. The differences between DGP-reactive PCs and non-DGP-reactive PCs were assessed by 2-tailed Student’s *t* test (**P < 0.01 and ****P < 0.0001). (C) Diversity and clonal size distribution of DGP-reactive and non-DGP-reactive PCs as measured by Hill’s diversity index. *D* values depict the level of diversity for a given value of q; lower *D* values represent lower diversity. Shaded area represents 95% percentiles.
The prototype IGHV3-23/IGLV4-69 and IGHV3-15/IGKV4-1 hmAbs recognize signatures of TG2 processing in the DGP

We aimed to scrutinize the basis for the predominant usage of IGHV3-23 and IGHV3-15 gene segments in DGP-reactive PCs. Among the 38 hmAbs previously generated from gluten-specific gut PCs, there was a striking heavy-chain variable region/light-chain variable region (VH/VL) pairing preference with IGHV3-23 preferentially pairing with IGLV4-69 and IGHV3-15 with IGKV4-1 (18); hmAbs of these configurations in the panel were all reactive with the DGP. Of note, the IGHV3-23/IGLV4-69 hmAbs also recognize the native form of the epitope, whereas IGHV3-15/IGKV4-1 hmAbs are only reactive with the deamidated peptide (18, 21). The differential reactivity to native/deamidated peptides suggested that the 2 types of antibodies might differ in their fine specificity.

We investigated this by assessing binding of 2 prototypical hmAbs, 1E01 (IGHV3-23/IGLV4-69) and 1E03 (IGHV3-15/IGKV4-1), to Ala-substituted variants of the DGP. Disassociation constants (K) for the 2 hmAbs and the peptide variants were determined by microscale thermophoresis (MST). Impaired binding was observed for Gln3, Pro4, Pro7, and Phe8 for hmAb 1E01 and for Pro4, Glu5, and Gln6 for hmAb 1E03 (Figure 2A). This analysis suggested that the 2 hmAbs make different interactions with the DGP. To obtain insight at atomic level for the antibody-antigen binding, we undertook X-ray crystallography studies. We produced Fab fragments of the 2 antibodies and generated crystal structures of Fab fragments with and without bound peptide antigens — altogether 5 structures: 1E01Fab, 1E01Fab-E, 1E01Fab-Q, 1E03Fab, and 1E03Fab-E.

The crystal structures of 1E01Fab-E and 1E01Fab-Q were highly similar (Figure 2B and Supplemental Figure 2, A and B). The root-mean-square deviation (rmsd) value for superimposing the 2 structures was 0.374, and the spatial orientations of the peptide residues were very similar (Supplemental Figure 2B). The rmsd values
for superimposing 1E01_{Fab} with 1E01_{Fab-Q} and 1E01_{Fab-E} were 0.969 and 0.920, respectively (Supplemental Figure 2, C and D). There were good electron densities for the CDR3 loops of both 1E01_{Fab-Q} and 1E01_{Fab-E} (Supplemental Figure 2, E and F). Further, in both structures, the peptide stretched (N- to C-terminus) in an extended conformation from the CDR-L1, via its CDR-L3 and CDR-H3, to CDR-H2 (Figure 2B and Supplemental Figure 2A). The side chains of the Glu5/Gln5 in both 1E01_{Fab-E} and 1E01_{Fab-Q} faced the solvent and did not interact with the antibody (Supplemental Figure 2, B and G). Residues located in all 3 CDR-H loops, in FR-H2 and FR-H3 as well as in CDR-L1 and CDR-L3, interacted with either main chain or side chain of 8 peptide residues via hydrogen bonds, or hydrophobic or aromatic interactions (Supplemental Figure 2G and Table 1. A summary of the interactions between 1E01_{Fab} and 1E03_{Fab} and the DGP (PLQPEQPFP)

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<th>Pro1</th>
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<th>Gln3</th>
<th>Pro4</th>
<th>Glu5</th>
<th>Gln6</th>
<th>Pro7</th>
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<td>Val38-L†</td>
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<td>Tyr37-L</td>
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<th>Pro4</th>
<th>Glu5</th>
<th>Gln6</th>
<th>Pro7</th>
<th>Phe8</th>
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<td></td>
<td>Main chain-side chain</td>
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<td></td>
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<td>Ile112-H*</td>
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**Peptide main chain; † peptide side chain.**
Table 1). The interface area for 1E01Fab-E and 1E01Fab-Q were 669 Å² and 654 Å², respectively, with about equal contribution by the heavy and light chains in both complexes (Table 2). There were 2 large pockets in the interaction surface of 1E01Fab that accommodated side chains of the peptide, with Gln3 filling pocket A and Pro7 and Phe8 filling pocket B (Figure 2C).

In the structures of 1E03Fab and 1E03Fab-E, electron densities for all CDR loops were intact, with the exception of molecule 1 in the asymmetric unit of 1E03Fab, where 5 residues were missing in the CDR-H3 loop (Supplemental Figure 3, A–D). For 1E03Fab-E, the Fab made an interaction with an extended conformation of the peptide via its FR-H2 and CDR-H3, as well as all 3 CDR-L loops. The orientation of the peptide was distinctly different from that seen in the 1E01Fab-E and 1E01Fab-Q structures (Figure 2B). The interface area was 524 Å², with the biggest contribution from the light chain (Table 2). The rmsd value for superimposing 1E03Fab with 1E03Fab-E was 0.552 (Supplemental Figure 3E), demonstrating that binding of the peptide antigen, as was also observed for 1E01, did not induce changes in structure of the Fab. No real pockets could be defined in the interface area (Figure 2D). Peptide residues Pro4, Glu5, and Gln6 interacted with the antibody (Supplemental Figure 3F and Table 1). In particular, the salt bridge formed between Arg55-H and Glu5 is likely to give an enthalpy change of paramount importance for binding (Figure 2D and Table 1). Importantly, for both 1E01Fab-E/Q and 1E03Fab-E, all residues of the Fab fragments that made contact with the peptides were germline encoded, apart from V38-L of 1E01Fab and R109-L of 1E03Fab.

Results of the Ala-scan affinity resonated well with the insight obtained from the crystal structures for both 1E01 and 1E03. This is also reflected for the B factors of each residue of the peptides when bound to the two Fab fragments (Figure 2A). Higher B factor values were seen for residues at both ends of the 9-mer peptides. For 1E01Fab-E, an increase in B factor values were seen for the Glu5 and Gln6 compared with their neighboring residues, reflecting less interaction between the side chain of the part of the peptide with this hmA. In-depth mutational analysis of positions involved in binding of the DGP

To further understand the basis for antibody-recognition of posttranslational modified gluten antigen in CD and to extend these observations beyond the 2 prototypical hmA, we undertook a detailed analysis of antibody mutations. We interrogated HTS data for positions involved in interactions with the DGP in the X-ray crystal structures (Table 1; position 55 for IGHV3-15 and positions 52, 64, and 66 for IGHV3-23) looking at mutations and affinity-depended selection strength. Data from patients from whom we had at least 30 unique sequences of each gene segment were analyzed (i.e., patients CD1182, CD1195, and CD1287 for

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<th>CDR length:</th>
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<td>Interaction surface area</td>
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<td>Light chain to peptide</td>
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<td>Interface atoms</td>
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### Hydrogen bonds (HB)

| Peptide side chain HB | 3 |
| Peptide main chain HB | 6 |
| Salt bridges | 0 |

Table 2. Interface characteristics of interactions between the DGP (PLQPEQPFP) and Fab fragments

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IGHV3-15 and CD1182, CD1188, CD1203, CD1279, CD1283, and CD1287 for IGHV3-23). Furthermore, we assessed the functional importance of these positions for binding to DGP by testing single amino acid mutants of the prototype hmAbs 1E03 and 1E01.

**Analysis of IGHV3-15.** At position 55 in DGP-reactive PCs of all 3 patients had lower number of mutations than median, with evidence for negative selection in all patients (Figure 3A, Supplemental Figure 4A, and Table 3). Non–DGP-reactive PCs most often mutated to His or Leu at this position. The hmAb 1E03 carries a germline-encoded Arg position, and changing this residue to His abolished binding (Figure 3B). This strong effect is expected, as R55-H in the 1E03Fab-E structure forms a salt bridge with the side chain of the Glu5 peptide residue (Figure 2D and Table 1). The Glu5 side chain also forms a hydrogen bond with Thr113-L (Figure 2D and Table 1). We therefore generated a variant of the 1E03 hmAb carrying Ala at this light chain position and found that this also severely impaired binding (Figure 3B). Our results, thus, demonstrate that the germline-encoded R55-H and T113-L residues are both essential for interactions with the key Glu5 residue of the DGP peptide (Figure 2A).

**Analysis of IGHV3-23.** At position 52, DGP-reactive PCs of all 6 patients analyzed had a lower-than-median number of mutations, but no information on selection strength could be discerned, due to lack of mutations (Figure 4A, Supplemental Figure 4B, and Table 3). Non–DGP-reactive PCs also had few mutations at this position (Figure 4A and Supplemental Figure 4B), which is why we did not test antibody mutants for this position. At position 64, 5 of 6 patients had mutation frequencies higher than median but with no clear sign of positive or negative selection (Figure 4A, Supplemental Figure 4B, and Table 3). We observed a preference for Ser to Gly or Thr substitutions in DGP-reactive PCs at this position, while non–DGP-reactive PCs had a variety of substitutions, with Arg and Asn being the most frequent residues (Figure 4A and Supplemental Figure 4B). The hmAb 1E01 carries a germline-encoded Ser at this position. No changes in binding were observed by mutation of Ser to Gly or Thr (Figure 4B). S64-H is making a hydrogen bond to the main chain oxygen of peptide residue Pro9 (Table 1). Our findings suggest that this hydrogen bond is of little importance for binding. At position 66, DGP-reactive PCs acquired very little

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**Figure 3. Somatic mutations in IGHV3-15 and the effect of positions involved in binding on the affinity of 1E03 hmAb.**

(A) Frequency of amino acid substitution along the IGHV3-15 gene segment of DGP-reactive and non–DGP-reactive PCs from a representative CD patient with more than 30 sequences available for analysis. (B) The effect of single amino acid substitution on binding affinity of hmAbs 1E03 to DGP as assessed by MST (average of 3 measurements ± SD). Variants at R55-H and T113-L were tested. GL, germline; VH, heavy chain; VK, K light chain.
mutations, and only one patient (CD1283) had a mutation number that was higher than median (Figure 4A and Supplemental Figure 4B). There were signs of negative selection in 2 patients and positive selection in 2 patients, as well as a lack of mutations prevented assessment of selection strength in 2 patients. Non–DGP-reactive PCs, by comparison, most often mutated to Phe or Asp at this position (Figure 4A and Supplemental Figure 4B). The hmAb 1E01 carries a germline-encoded Tyr at this position, and mutations to Phe or Asp profoundly impaired binding (Figure 4B). In the 1E01Fab-E structure, Y66-H is involved in binding with a hydrogen bond to the main chain oxygen of peptide residue Pro9, as well as in aromatic and hydrophobic interactions with Phe8. The binding results suggest that this germline-encoded residue is important for the interaction with the DGP.

In summary, from our HTS analysis, we found that some positions involved in binding of the DGP peptide had a higher-than-median number of mutations. Other positions, like position 55 of IGHV3-15, had virtually no mutation and had signs of negative selection. The finding for this position is particularly notable as R55-H makes a salt bridge with the peptide and, thus, has a dominant role in binding of DGP. Overall, our results reveal that germline-encoded residues at interacting positions are favorable for binding to DGP. This resonates well with the general low degree of mutations present among DGP-reactive PCs.

Selection at position 55 of IGHV3-23 for small amino acids that allow for binding to DGP

Strikingly, the degree of mutations at position 55 in IGHV3-23 was more pronounced than what we observed for positions 52, 64, and 66, and we observed that DGP-reactive PCs uniformly across all 5 patients had mutation from Ala (germline) to Gly, with evidence for positive selection in all subjects (Figure 4A and Supplemental Figure 4B). Although this residue was not identified to make interactions with the peptide in the crystal structures, the reason for this mutation pattern could potentially relate to binding of the DGP. We hence undertook a further examination, including analysis of intracodon mutation patterns for this position. In comparison with non–DGP-reactive PCs, DGP-reactive PCs had statistically significant substitution patterns, with fewer nucleotide substitutions, and when acquiring a substitution, this typically was a single nucleotide substitution at position 2 of the codon (i.e., nucleotide 164) (Supplemental Figure 5, A and B). By contrast, non–DGP-reactive PCs had multiple nucleotide substitutions or single substitutions at positions 1 and 2. Moreover, the type of substitution at position 2 was highly selective for DGP-reactive PCs, with 84% of the sequences having a substitution at this position only, compared with 34% of non–DGP-reactive PCs. Typically, the cells had a C164G substitution, encoding an A55G mutation (Supplemental Figure 5A). These findings suggest that G55-H encoded by IGHV3-23 is preferred in interaction with DGP.

We tested single amino acid mutants of the 1E01 hmAb to assess impact of position 55 of IGHV3-23 for binding to DGP. The hmAb 1E01 carries Gly at position 55 that is altered from Ala in germline. In contrast to DGP-reactive PCs, non–DGP-reactive PCs often had Ser or Thr, whereas Ala and Gly were observed in both types of PCs. Substitution from Gly to Ala, Ser, or Thr in hmAb 1E01 led to impaired binding — particularly, the Gly to Thr mutation had a big effect (Figure 5A). In the 1E01Fab-E structure, G55-H lines a binding pocket that accommodates the side chain of peptide residue Phe8 (Figure 5B). Conceivably, A55-H and G55-H could be favorable residues at this position, due to the small or absent side chains. S55-H and T55-H, however, have larger side chains that could cause steric hindrance and, hence, impaired binding (Figure 5, B and C). Phe8 is part of the recognition sequence of TG2 (11, 12), and this observation again demonstrates that the DGP-reactive cells carry BCRs that recognize signatures of the peptides that reflect processing by TG2.

Table 3. Positive and negative selection strengths at positions that are involved in binding to the DGP

<table>
<thead>
<tr>
<th></th>
<th>CD1182</th>
<th>CD1188</th>
<th>CD1195</th>
<th>CD1203</th>
<th>CD1279</th>
<th>CD1283</th>
<th>CD1287</th>
<th>Mean selection (weighted)</th>
</tr>
</thead>
<tbody>
<tr>
<td>IGKV3-15 pos.55</td>
<td>-3.65</td>
<td>NA</td>
<td>-3.02</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>-3.65</td>
<td>-3.13 ± 0.61</td>
</tr>
<tr>
<td>IGKV3-23 pos.52</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>IGKV3-23 pos.55a</td>
<td>0.74</td>
<td>0.66</td>
<td>NA</td>
<td>0.74</td>
<td>NA</td>
<td>0.74</td>
<td>0.48</td>
<td>0.64 ± 0.36</td>
</tr>
<tr>
<td>IGKV3-23 pos.64</td>
<td>-0.07</td>
<td>-0.87</td>
<td>NA</td>
<td>-0.07</td>
<td>NA</td>
<td>-1.41</td>
<td>0.42</td>
<td>-0.39 ± 0.52</td>
</tr>
<tr>
<td>IGKV3-23 pos.66</td>
<td>1.02</td>
<td>-1.79</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>1.29</td>
<td>-3.26</td>
<td>0.05 ± 1.16</td>
</tr>
</tbody>
</table>

aPosition 55 of IGKV3-23 does make interactions with the peptide in the crystal structures but is included, as it demonstrated a uniform pattern of mutation across PCs of all patients. NA, not applicable.
BCR-mediated uptake of DGP facilitates presentation of gluten T cell epitopes

Antigen uptake by specific BCRs can greatly facilitate presentation of T cell epitopes that are physically linked to the B cell epitopes (23). To test whether B cells specific to DGP enhance presentation of gluten T cell epitopes, we designed an antigen presentation assay to T cells using, as antigen presenting cells, A20 B cell lymphoma cells retrovirally transduced with HLA-DQ2.5 encoding genes, as well as genes for 1E03 or a non–DGP-reactive mAb (2F02) in the format of membrane-bound human IgD. IL-2 secretion from transfectant T cells specific for the DQ2.5-glia-ω2 epitope was used as a readout. As antigen, we used a synthetic ω34-mer fragment (ω34mer peptide) that is naturally created by gut enzyme digestion and that contains the DQ2.5-glia-ω2 T cell epitope, as well as 3 copies of the B cell core epitope (QPEQPF) (21). We observed that transfectant B cells expressing 1E03 as BCR rapidly bound the ω34mer peptide on their surface (Figure 6A) and stimulated the T cells more efficiently than B cells with irrelevant BCR (Figure 6B). Hence, cognate T cell and B cell interaction can indeed operate when gluten T cell and B cell epitopes are physically linked.

As ω34mer peptide is an antigen that likely binds directly to surface MHC class II molecules, the difference between the specific and nonspecific B cells in our assay is less than what is usually observed for antigens that are completely antigen processing dependent for T cell recognition (23).

Discussion

In this study, we have dissected the antibody response against the peptide PLQPEQPFP, which harbors an immunodominant epitope of gluten recognized by highly disease-specific antibodies of CD patients.
Figure 5. Steric hindrance at position 55 of IGHV3-23 interferes with binding of DGP. (A) Binding of DGP by hmAb 1E01 mutated at heavy chain position 55 as assessed by MST (average of 3 measurements ± SD.). (B and C) Model for a possible steric hindrance following Gly-to-Thr substitution in the 1E01Fab-E structure. The side chain of Phe8 is accommodated in pocket B and is located in proximity of G55-H. (C) The side chain of Thr is predicted to fill into the pocket that should be accommodated by Phe8, thereby causing steric hindrance. GL, germline; VH, heavy chain.

(5, 18, 24, 25). By Ala scan and X-ray crystallographic studies, we defined the core epitope of the peptide to comprise residues QPEQPF. Two stereotyped classes of CD antibodies recognizing this epitope utilize the gene segments IGHV3-23/IGLV4-69 and IGHV3-15/IGKV4-1 for recognition leading to fundamentally distinct docking modes of the Fab fragments. Strikingly, both antibodies recognize signatures reflecting TG2 processing of the antigen, thereby supporting a critical role of the TG2 enzyme in shaping the anti-gluten antibody response in CD.

Using HTS of IGHV genes, we characterized the repertoire and clonal diversity of IgA DGP-reactive PCs from the gut of CD patients. These PCs had low clonal diversity and a narrow IGHV repertoire with bias in usage of the IGHV3-23 and IGHV3-15 genes, as compared with non–DGP-reactive PCs. This finding confirms and extends finding of 38 hmAbs from single-sorted PCs of CD patients that showed a paring preference of IGHV3-23 with IGLV4-69 and IGHV3-15 with IGKV4-1 (18). Studying 2 prototypic hmAbs 1E01 (IGHV3-23/IGLV4-69) and 1E03 (IGHV3-15/IGKV4-1) with epitope/paratope mutational analysis and X-ray crystallography, we found that the 2 hmAbs, despite interacting with extended peptide conformations with no structure adaptation, employ fundamentally different binding modes. Residues Pro4, Glu5, and Gln6 are essential for recognition of DGP by the 1E03 hmAb, whereas Gin3, Pro4, Pro7, and Phe8 are involved in recognition by the 1E01 hmAb. In both 1E01pα-Fab/E structures, key interactions of the antibody are mediated by peptide side chains (Pro4 and Pro7/Phe8) being buried in pockets in a knob-hole fashion, which is a type of interaction often seen in peptide-protein binding (26).

Importantly, no interaction takes place between 1E01Fab and the side chain of Glu5 in the peptide. In the interface of 1E03pα-Fab-E structure, which was slightly smaller than in the 1E01pα-Fab/E structures, no clear pockets could be defined. The Glu5 peptide residue makes critical contacts with the antibody residues, including a salt bridge with residue Arg55-H. Overall, the results provide an explanation for the differential recognition of deamidated and native variants of the PLQPEQPF peptide by the 1E01 (IGHV3-15/IGKV4-1) and 1E03 (IGHV3-23/IGLV4-69) antibodies.

Only the 1E03 (IGHV3-15/IGKV4-1) hmAb is truly deamidation specific and hence, in a strict sense, qualifies to be termed an anti-deamidated gliadin antibody. Nonetheless, the 1E01 (IGHV3-23/IGLV4-69) antibody recognizes a sequence that intimately associates with deamidation of the gluten peptide. The antibody recognition sequence QPQ/E PF is highly similar to the preferred substrate recognition sequence of TG2, which spans 5 residues from –1 to +3 (targeted Q at position 0) and harbors the core sequence QXP(F) at residues 0 to +3 (11, 12). As we demonstrate, B cells with BCR of this specificity will bind, internalize, and present peptides that are deamidated by TG2 for presentation to CD4+ gluten–reactive T cells and, hence, be in position to receive T cell help. This will be the case for the 1E03 (IGHV3-15/IGKV4-1) type of antibodies, which recognize the introduced glutamate residue itself. Even though the 1E01 (IGHV3-23/IGLV4-69) type of antibodies can also bind native peptides, this would not result in productive T cell help, as most gluten-reactive T cells of CD patients are strictly deamidation dependent (27). Importantly, both types of antibodies recognize residues in a peptide that intimately reflect productive and pathogenic T cell and B cell interaction in CD. Both types of antibodies, thus, serve as proxies for a key step in the pathogenesis of CD; hence, they become excellent surrogate markers for the disease.

The binding of the immunodominant gliadin epitope by both 1E01 (IGHV3-23/IGLV4-69) and 1E03 (IGHV3-15/IGKV4-1) hmAbs was essentially mediated by germline-encoded heavy and light chain residues. This would explain the selection for usage of certain VH and VL gene segments and also the preferential VH/VL paring among PCs reactive with the DGP. Moreover, our findings suggest that the reactivities of IGHV3-23/IGLV4-69 and IGHV3-15/IGKV4-1 stereotyped antibodies are already established with germline configurations, negating a need for naive B cells to undergo substantial somatic hypermutation to differentiate to effector cells. The HTS data demonstrating low levels of mutations further revealed that most sequences were in germline configuration at the positions that involved bonding of DGP. However,
evidence of selection at several of these positions was observed. Most notable was the negative selection at position 55 of IGHV3-15. In the 1E03-Fab-E structure, the germline-encoded Arg served as a lynchpin for the binding of the DGP peptide, explaining why preservation of germline configuration is favorable at this position. Position 55 of IGHV3-23, despite not bonding to the DGP in the crystal structure, was found to be frequently mutated in the HTS data, with evidence for positive selection. DGP-reactive PCs typically mutated from germline Ala to Gly while non–DGP-reactive PCs often also mutated to Ser or Thr. Analysis of the 1E01 crystal structures suggests that this position, which lines a binding pocket harboring the peptide Phe8 side chain, has a requirement for small residues. Ser and Thr with larger side chains could cause steric hindrance and, thus, impaired binding as was also experimentally observed. Taken together, it seems that B cells reactive with the peptide PLQPEQPFP undergo some selection in CD patients. However, at positions that are involved in peptide binding, there is preservation of germline-encoded residues of the antibodies.

In conclusion, in this study, we have combined HTS and X-ray crystallography to decipher the antibody response toward the immunodominant deamidated gliadin B cell epitope PLQPEQPFP in CD. We demonstrate that 2 stereotyped forms of antibodies bind to the same epitope fundamentally differently; one targets the recognition sequence of TG2, while the other targets the Glu residue formed by TG2-mediated deamidation. In essence, these 2 types of recognition lead to the same results: uptake of the deamidated peptide by BCR of gluten-reactive B cells, which in turn can receive T cell help from gluten-reactive T cells and thereby undergo clonal expansion. Interestingly, the fine specificity of the 2 stereotyped antibodies recognizing the immunodominant B cell epitope reflects key events in the pathogenesis of CD. On a general level, the isolation of antigen-specific PCs/B cells followed by detailed structural analysis of epitopes illustrates a strategy by how key aspects of disease that involve posttranslational modification and generation B cell and of T cell epitopes can be identified.

**Methods**

**Subjects and cells.** Ten patients with untreated, active CD were enrolled in this study. All were diagnosed according to the guidelines of the British Society of Gastroenterology (28). Single cell suspensions were obtained from small intestinal biopsies of the duodenum obtained by endoscopy as previously described and cryopreserved until use (18, 20).

**Isolation of DGP-reactive PCs and library preparation of IGHV HTS.** PCs reactive with the immunodominant gliadin epitope PLQPEQPFP were stained using antigen tetramers as previously described (18). In short, biotinylated-deamidated gliadin peptide (biotin-GSGSGS-PLQPEQPFP, GL Biochem) were incubated with SA-APC (PhycoLink) at a 4:1 molar ratio for 1 hour in PBS supplemented with 3% FCS on ice to create PLQPEQPFP-tetramers. Similarly, TG2 was tetramerized with SA-PE (Invitrogen) and used to exclude TG2-specific PCs from the analysis. Single cell suspensions were incubated with the 2 tetramers for 45 minutes, and the following PC markers were added for the last 20 minutes to stain for specific cellular lineage markers: IgA-FITC (SouthernBiotech, catalog 2050-02), Brilliant Violet 570 CD3 and CD14 (BioLegend, clones UCHT1 and M5E2, respectively), CD19 Pacific Blue (BioLegend, clone HIB19) and CD27-PE-Cy7 (eBioscience, clone LG.7F9). IgA intestinal PCs were defined as live, large, and CD14/CD3-negative, with high expression of CD27. PCs were sorted for being either negative or positive for the PLQPEQPFP-tetramers into 25 μl catch buffer (10 mM DTT and RNAsin 2 U/μl [Promega] in PBS) and kept in –70°C until cDNA synthesis. For cDNA synthesis, 13.5 μl of sorted PC extracts were added to the mix containing indexed IGJH reverse primer (1.4 μM), 0.45% v/v NP-40, and 0.5 μl RNasin 40 U/μl (Promega). DEPC-treated water was added to the final volume of 25 μl, incubated at 65°C for 5 minutes, and put on ice. Next, 10 μl of 5× reverse transcription
(RT) buffer, 3.5 μl of 100 mM DTT, 4.3 μl of 10 mM dNTP-Mix (Promega), 0.7 μl of 40 U/μl RNasin (Promega), 0.85 μl superscript III (Invitrogen), and 4.5 μl DEPC-treated water were added. cDNA was synthesized at 42°C for 10 minutes, 25°C for 10 minutes, 50°C for 60 minutes, and 94°C for 5 minutes and stored at –20°C. Second-strand cDNA was synthesized using AmpliTaq Gold polymerase (Applied Biosystems) with forward IGHV1-6 framework region (FR) 2 (29), which were indexed and included 6 random nucleotides (further used as unique molecular identifiers [UMIs]) and a part of Illumina adapt-ers. Second-strand synthesis was carried out at 95°C for 7.5 minutes, 52°C for 2 minutes, and 72°C for 10 minutes. Double-stranded cDNA was purified using AMPure XP (Beckman Coulter) at a 1:1 ratio according to the manufacturer’s instructions. Next, the second part of the Illumina adapter was connected using R1 and R2 primers. PCR was performed using Qiagen Multiplex PCR at 95°C for 15 minutes, ×25 (95°C for 30 seconds, 60°C for 45 seconds, 72°C for 90 seconds), and 72°C for 10 minutes. Primers were previously described (20). Final IGHV amplicon libraries were purified using AMPure XP (Beckman Coulter) and further run and extracted from agarose gel. Paired-end sequencing of 250 × 2 or 300 × 2 bp was performed using Illumina MiSeq at the Norwegian Sequencing Centre in Oslo, Norway (http://www.sequencing.uio.no). Raw sequencing data was processed using pRESTO (30) as previously described (20, 31). Unique sequences to which at least 2 copies were present were further aligned using ImMunoGeneTics (IMGT) database (32), and their IGHV and IVHJ segments and CDR3 were determined. The number of unique sequences is given in Supplemental Table 1.

**Diversity of B cell clone measurements.** Sequences were clustered into clones originating from common ancestors using Change-O (33) as described previously (20). Diversity scores for clones from DGP-reactive and non–DGP-reactive PCs were calculated using the Hill diversity index, which measures diversity in a population (qD) as a smooth function of a single parameter (q) (22, 34).
\( D = \left( \sum_{i=1}^{S} p_i^q \right)^{1/(1-q)} \)

where \( S \) is the number of clones, \( p_i \) is the frequency of the \( i \)th clone, and the parameter \( q \) determines how clonal frequencies are weighted (34).

Using the Hill diversity index, we determined whether the repertoire diversity indices differed between these subsets. To eliminate sampling effects, repertoire subsets were subsampled (1,000 times) according to the number of sequences in the smallest sample (i.e., the DGP-reactive PCs), and Hill’s curve was calculated in 1,000 equally spaced \( q \) values between 0 and 15. At a given value of \( q \) (x-axis in Figure 1D), lower values of \( D \) (y-axis) indicate lower diversity (i.e., more focused repertoire).

**Fab fragments and antibody mutants.** The hmAbs 1E01 (IGHV3-23/IGLV4-69) and 1E03 and (IGHV3-15/IGKV4-1) were studied (18). Fab fragments of these 2 hmAbs were generated by introducing a stop codon in the plasmid after residue 232 (C288PKSCD232). PCR was performed by using forward primer 5’-TTTCTAGTAGCAACTGCAAC-3’ and reverse primer 5’-GTGGACAAGAAAGTTGAGC-CCAAATCTTGGTACGCTGAAGCTTG-3’, and the products were inserted into the expression vector IgG1 between the AgeI and HindIII restriction sites. The light chains of the 2 antibodies have been previously cloned (18). Plasmids encoding the heavy and light chains were cotransfected into HEK 293-F (ThermoFisher Scientific, R79007) cells by using 293fectin (Invitrogen) or polyethyleneimine (Polysciences Inc.). Transfected HEK 293-F cells were cultured at 37°C with shaking for 6 days. Anti-gliadin autoantibodies were purified from culture supernatants. The 1E01 Fab fragment was affinity purified by using CaptureSelect LC-lambda (Hu) Affinity Matrix (Life Technologies), whereas the 1E03 Fab fragment was affinity purified using a protein L column (GE Healthcare). Purity of the proteins was confirmed by SDS-PAGE.

Site-directed mutagenesis in 1E01 and 1E03 hmAbs was induced using Quickchange Multisite-Directed Mutagenesis Kit (Agilent) following the manufacturer’s instructions. Mutations were verified by sequencing of the expression vectors (GATC Biotech). Primers that were used to induce mutations are listed in Supplemental Table 2. Expression vectors containing the original or the mutated genes were cotransfected into HEK 293-F cells by using polyethyleneimine (Polysciences Inc.). Transfected HEK 293-F cells were cultured at 37°C with shaking for 6 days. Anti-gliadin autoantibodies were purified from culture supernatants. The 1E01Fab fragment was affinity purified by using CaptureSelect LC-lambda (Hu) Affinity Matrix (Life Technologies), whereas the 1E03Fab fragment was affinity purified using a protein L column (GE Healthcare).

**Crystallization and data collection.** The purified Fab fragments were concentrated to 12 mg/ml in 20 mM Tris, pH 7.2, 150 mM NaCl, 1 mM EDTA. Complexes between the Fab fragments and peptide were formed by incubating them at 1:8 molar ratio overnight at 4°C. Initial large-scale screenings were performed by using the sitting drop vapor diffusion method using Cartesian Dispensing system (Genomic Solutions, Madison, Wisconsin, USA). Promising hits were chosen for crystallization optimization. Crystals of 1E01Fab alone (1E01Fab), 1E01Fab in complex with deamidated peptide (1E01Fab-E), 1E01Fab in complex with native peptide (1E01Fab-Q), 1E03Fab alone (1E03Fab), and 1E03Fab-E in complex with deamidated peptide (1E03Fab-E) (Supplemental Table 3) were generated and resolved: 1E01Fab-E, alone (1E01Fab), 1E01Fab-E in complex with deamidated peptide (1E01Fab-E), 1E01Fab-E in complex with native peptide (1E01Fab-Q), 1E03Fab-E in complex with deamidated peptide (1E03Fab-E) (Supplemental Table 3). Crystals of 1E03Fab-E in complex with native peptide could not be generated, as 1E03Fab does not bind the native peptide. The cryo conditions for freezing the crystals were 25% v/v glycerol in the mother liquor, except for 1E03Fab-E, which required 20% v/v glycerol for cryo solvents. Each single crystal was transferred to the cryo solution and flash-frozen in liquid nitrogen for data collection. Data were processed and scaled by XDS (37) for all crystals, except that of 1E03Fab-E, which was processed by XDS (37), POINTLESS (38), and AIMLESS packages (39).

**Structure determination and refinement.** All the 5 structures were determined by molecular replacement using CCP4 Phaser (40), and the models used for searching are listed in Supplemental Table 3. Manual buildings of all the 5 structures were carried out by Coot (41), and refinements were implemented by Refmac (42) and PHENIX (43). Validation was done by the validation servers in Protein Data Bank (PDB) and MolProbity (44). The residues were numbered according to the IMGT numbering system (32). Data processing and refinement statistics are summarized in Table 4. Ramachandran statistics indicated that, for all structures, 95.5%–97.8% of the residues had a favored geometry, with 0.4%–0.7% of the residues being outliers. Interactions between the peptide and Fab fragments were analyzed by PIC (45). The determination of interaction surface area, was analyzed by PISA (http://www.ebi.ac.uk/pdbe/prot_int/pistart.html) (46). Analysis of pockets on the protein monomers of bound structures was done using the CASTp server (47), and the pockets in the interface area were selected. Each asymmetric unit had several molecules (Supplemental Table 3) and the interaction analysis was made on the following chains: 1E01Fab (PDB 5IHZ), chains A, B; 1E01Fab-E (PDB 5IFJ), chains D, E, F; 1E01Fab-Q (PDB 5IG7), chains D, E, F; 1E03Fab (PDB 5IHY), chains A, B.
non–DGP-reactive PCs and were compared using a χ² test for goodness of fit. For each patient, intracodon substitution patterns were summarized for DGP-reactive and non–DGP-reactive PCs were analyzed by assigning each amino acid position along the gene segments to low/high mutation rates relative to the median mutation rate for each IGHV gene of each sample. Affinity-dependent selection strength among DGP-reactive PCs was quantified for selected residue in the structures interrogating positions involved in direct bonding with the peptide. Sequences of DGP-reactive and non–DGP-reactive PCs were analyzed by assigning each amino acid position along the gene segments to low/high mutation rates relative to the median mutation rate for each IGHV gene of each sample. Affinity-dependent selection strength among DGP-reactive PCs was quantified for selected residue in the IGHV gene segments using a tailored version of BASELINe (49, 50), adapted to single codons. A local test that compares frequency of observed nonsynonymous mutations with their expected frequency was applied with the SSF targeting model to account for somatic hyper mutation hot/cold spots (31). To avoid biases that may have emerged due to unknown order of mutation acquisition, only sequences with no more than 1 mutation per codon were used. Intracodon mutation pattern was analyzed for position 55 of IGHV3-23. This analysis is based on the fact that, for each codon, 8 distinct combinations of substitutions were possible. For each patient, intracodon substitution patterns were summarized for DGP-reactive and non–DGP-reactive PCs and were compared using a χ² test for goodness of fit. P values were adjusted using Benjamini-Hochberg correction.

**Antibody affinity measurement using MST.** Binding affinity between hmAbs (1E01, 1E03, or their mutants) and the peptide PLQPEQPFPF or Ala-substituted variants of the same peptide were determined using MST. Analyses were performed on a Monolith NT.LabelFree Instrument using Monolith NT.LabelFree capillaries (Nano Temper Technologies). For MST measurements, hmAbs were used at a constant concentration of 0.25 µM. Gliadin peptides were diluted 16 times in a 1:1 serial dilution from 0.5 mM to a final concentration of 15.25 nM and incubated at room temperature with the antibodies for at least 5 minutes before measurement. All measurements were performed in triplicate, and data analysis was performed with the Monolith software (Nano Temper Technologies).

**Data alignment and genotyping.** Sequences were aligned to the IMGT database using IMGT/HighV-QUEST (32). Allele frequency was calculated for each gene segment. To be included in an individual’s reference, an allele had to appear in at least 20% of nonmutated sequences assigned to the same gene or at least 20 times, regardless of the frequency. Realignment was accomplished using IgBLAST (48) with the personalized germline reference.

**Analysis of mutations, selection, and intracodon nucleotide substitution patterns.** Analysis of mutations in HTS data of IGHV3-23 and IGHV3-15 was done using data from patients from whom we had at least 30 unique sequences of each gene segment with more that 5 clones that were not dominated by a single clone (i.e., patients CD1182, CD1188, CD1203, CD1279, CD1283, and CD1287 for IGHV3-23 and CD1182, CD1195, and CD1287 for IGHV3-15). Mutational analysis was done in the context of the X-ray crystal structures interrogating positions involved in direct bonding with the peptide. Sequences of DGP-reactive and non–DGP-reactive PCs were analyzed by assigning each amino acid position along the gene segments to low/high mutation rates relative to the median mutation rate for each IGHV gene of each sample. Affinity-dependent selection strength among DGP-reactive PCs was quantified for selected residue in the IGHV gene segments using a tailored version of BASELINe (49, 50), adapted to single codons. A local test that compares frequency of observed nonsynonymous mutations with their expected frequency was applied with the SSF targeting model to account for somatic hyper mutation hot/cold spots (31). To avoid biases that may have emerged due to unknown order of mutation acquisition, only sequences with no more than 1 mutation per codon were used. Intracodon mutation pattern was analyzed for position 55 of IGHV3-23. This analysis is based on the fact that, for each codon, 8 distinct combinations of substitutions are possible. For each patient, intracodon substitution patterns were summarized for DGP-reactive and non–DGP-reactive PCs and were compared using a χ² test for goodness of fit. P values were adjusted using Benjamini-Hochberg correction.

**T cell and B cell collaboration assay.** Murine A20 B lymphoma cells were engineered by retroviral transduction to express HLA-DQ2.5 and 1002-1E03 BCR in an IgD format as previously described (19). To assess capacity of these transfectant B cells to bind deamidated gluten peptide, cells were stained with tetramers for 30 minutes on ice and analyzed on a FACSCalibur flow cytometer (BD Biosciences). Gluten peptide tetramers were generated by incubating the biotinylated 34-mer α gliadin peptide (QPQQPFQPQOEPOEPFEQPFPWPQPFPQ, GL Biochem) with SA-APC (PhycoLink) at a 4:1 molar ratio for 2 hours in PBS supplemented with 2% FCS on ice. Murine BW519-β2 cells devoid of endogenous TCR were engineered to express human CD4 and T cell receptor from a DQ2.5-glia-α2-reactive CD4 T cell clone (TCC737.30) (51) as previously described (52). Transfected B cells (1 × 10⁶ cells/ml) expressing HLA-DQ2.5 alone, HLA-DQ2.5 with DGP-negative (693-2F02) BCR, or HLA-DQ2.5 with DGP-reactive BCR (1002-1E03) were incubated in RPMI supplemented with 10% FCS with indicated concentrations of α34-mer peptide for 10 minutes on ice and washed with RPMI to remove unbound peptide. Transfected B cells (50,000) were incubated for 4 hours in 96-well U-bottom plates at 37°C before transfectant T cells (25,000) were added. After 18 hours, culture supernatants were collected and assayed for murine IL-2 production by ELISA as previously described (52). IL-2 concentrations were calculated from a standard curve generated using recombinant murine IL-2 (BioLegend).

**Accession codes.** HTS data have been deposited at the National Center for Biotechnology Information Sequence Read Archive (SRA) (accession code SRP098633, bioproject PRJNA368623). Protein structure data have been deposited in the worldwide PDB (wwPDB) under accession codes 5IHZ, 5IFJ, 5IG7, 5IK3 and 51JK.
Statistics. A two-tailed t test was applied to compare the number of acquired mutations in non-DGP-reactive and DGP-reactive PCs. P < 0.05 was considered significant. χ² test for goodness of fit followed by Benjamini-Hochberg correction for the adjustment of the P values were used to compared the calculated intracodon mutations in non-DGP-reactive and DGP-reactive PCs. Hill’s diversity index (22) was used to determine the repertoire diversity as reflected by the clonal size distribution. The repertoire was subsampled 1,000 times according to the number of sequences in the smallest subsets (as described above). Statistically significant differences between samples can be visualized when curves describing the diversity of 2 PC subsets are separated and fall outside the 95% that resulted from random sampling.

Study approval. All subjects had signed informed consents, and the study was approved by the Regional Ethics Committee of South-Eastern Norway (2010/2720).

Author contributions
ΟS, OS, and LMS conceived the project. OS processed gut biopsies, sorted PCs, prepared the amplicon library for MiSeq sequencing, performed binding affinity studies, and produced mutants of hmAbs. XC produced and purified proteins for crystallography, generated crystals, solved protein structures by X-ray crystallography, and performed binding affinity studies. MG processed and analyzed sequence data. MFDP performed T cell and B cell collaboration assays. YZ was involved in crystallography work. ΟS performed PC sorting. KEAL performed the endoscopies and provided small-intestinal biopsies. LMS and OS wrote the manuscript with contributions from other authors. GY and LMS supervised the study.

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Address correspondence to: Ludvig M. Sollid, Department of Immunology, Oslo University Hospital–Rikshospitalet, NO-0372 Oslo, Norway. Phone: 47.230.73.811; Email: l.m.sollid@medisin.uio.no.

