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Alginate Trisaccharide Binding Sites on the Surface of β-Lactoglobulin Identified by NMR Spectroscopy: Implications for Molecular Network Formation

Emil G. P. Stender,†,‡ Johnny Birch,†,‡ Christian Kjeldsen,‡ Lau D. Nielsen,§ Jens Ø. Duus,‡ Birthe B. Kragelund,*,§ and Birte Svensson*,‡

Introduction

β-Lactoglobulin (BLG) is an extensively studied lipocalin found in bovine milk,3–6 which is able to bind a large variety of hydrophobic ligands.3–6 The common fold of lipocalins is an 8-stranded antiparallel β-barrel that encloses an internal hydrophobic cavity, the calyx. The BLG fold includes nine β-strands, of which, strands A–H form the central calyx and strand I is involved in BLG dimer interface formation along with an α-helix (Figure 1).7 The tertiary structure of BLG is further stabilized by two disulfide bonds (C66–C160 and C106–C119), which are buried together with a free cysteine (C121) in the hydrophobic core.8 BLG exhibits a variable fi further stabilized by two disulcalyx,3 to which access is controlled by the EF-loop that adopts 8-stranded antiparallel β-sheets, of which, strands A and I is involved in BLG dimer interface formation along with an α-helix (Figure 1).7 The tertiary structure of BLG is further stabilized by two disulfide bonds (C66–C160 and C106–C119), which are buried together with a free cysteine (C121) in the hydrophobic core.8 BLG exhibit a variable structural landscape, comprising a homodimer at neutral pH8,9 and a monomer–dimer equilibrium at pH < 5 and is essentially monomeric at pH < 3.10 Different hydrophobic ligands, e.g., fatty acids and retinoids,5,6 can be accommodated inside the calyx,3 to which access is controlled by the EF-loop that adopts a closed conformation at pH < 7.5 (Figure 1) and an open conformation at a higher pH.10 Notably, BLG also has binding sites on the outer surface. Caprylic acid thus interacts with the loop connecting β-strands C and D (CD-loop, residues W61–C66) at the entrance to the calyx at pH 7.5,6 while at pH 2, vitamin D3 binds at a surface-exposed hydrophobic site (D137–R148) belonging to β-strand I, the α-helix involved at the dimerization interface, and the loop connecting the two (Figure 1).5

Polysaccharides are widely used as food additives such as viscosity agents, pH modulators, water retainers, and antimicrobials11 and constitute important ligands for BLG. Below the pI (4.7–5.2),12–17 BLG interacts with a wide range of polysaccharides, resulting in either insoluble particles12,16,18 or liquid coacervates,15,17 depending on the nature of the polysaccharide and the buffer composition. While particle size, strength of interaction, and formation of insoluble particles or liquid coacervates may be controlled by the nature of the polysaccharide, the factor determining whether BLG adopts a closed or open conformation and is hence able to bind a particular polysaccharide may be a subtle difference in the structure of the polysaccharide. Here, we report that the binding of alginate oligosaccharides (AOSs) prepared by controlled enzyme degradation to the outer surface of BLG is pH-dependent and is facilitated by interactions with both the dimeric and the monomeric BLG isoform A (BLGA). The pH-dependent mechanism identified in this study is likely to have implications for biological function and the binding of other polysaccharides to BLG.

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Supporting Information

ABSTRACT: β-lactoglobulin (BLG) is a promiscuous protein in terms of ligand interactions, having several binding sites reported for hydrophobic biomolecules such as fatty acids, lipids, and vitamins as well as detergents. BLG also interacts with neutral and anionic oligo- and polysaccharides for which the binding sites remain to be identified. The multivalency offered by these carbohydrate ligands is expected to facilitate coacervation, an electrostatically driven liquid–liquid phase separation. Using heteronuclear single quantum coherence NMR spectroscopy and monitoring chemical shift perturbations, we observed specific binding sites of modest affinity for alginate oligosaccharides (AOSs) prepared by alginate lyase degradation. Two different AOS binding sites (site 1 and site 2) centered around K75 and K101 were identified for monomeric BLG isoform A (BLGA) at pH 2.65. In contrast, only site 1 around K75 was observed for dimeric BLGA at pH 4.0. The data suggest a pH-dependent mechanism whereby both the BLGA dimer–monomer equilibrium and electrostatic interactions are exploited. This variability allows for control of coacervation and particle formation of BLGA/alginate mixtures via directed polysaccharide bridging of AOS binding sites and has implication for molecular network formation. The results are valuable for design of polyelectrolyte-based BLG particles and coacervates for carrying nutraceuticals and modulating viscosity in dairy products by use of alginate.
coacervates have been attributed to chemical and physicochemical characteristics of the polysaccharide, through charge neutralization, the residues of BLG involved in the complex formation have so far not been identified. Positively charged patches on the BLG surface have been proposed to mediate interactions with anionic polysaccharides through charge neutralization, leading to coacervation or formation of insoluble particles. However, the large sizes of BLG/polysaccharide complexes have prevented detailed descriptions of their molecular structures. This knowledge void motivated the present analysis of a currently developed model system involving alginate trisaccharides binding to BLG isoform A (BLGA). The major aim was to decipher the details of the molecular interactions.

Alginate is a linear, acidic polysaccharide composed of α-L-guluronic acid (G) and β-D-mannuronic acid (M) organized as a copolymer with a pattern of 1,4-linked G, M, and M/G blocks. Alginate is widely used as a food additive in dairy products as a modulator of viscosity and consistency. It can also protect whey proteins from gastric digestion, and BLG/alginate particles have been proposed as nutraceutical carriers.

The structure of these compounds has previously been solved by NMR, and the assigned chemical shifts are in good correlation with previous results.

BLGA Has Two Different Binding Sites for AOSs at Low pH. AOS binding to BLGA was investigated under different conditions than those used for earlier NMR chemical shift assignments of BLGA. Therefore, BLGA was prepared with the same primary structure as earlier and the first NMR spectra were recorded under the same conditions as those used in that study. Subsequently, the chemical shifts were followed under conditions used to measure binding via chemical shift perturbations upon addition of AOSs. BLGA has 165 residues including eight prolines and the additional three residues of the N-terminal (see Materials and Methods, Figure S1). Accordingly, excluding the fast exchanging N-terminal and the side chain resonances, a total of 156 resonance peaks are expected. Indeed, the 1H,15N-heteronuclear single quantum coherence (HSQC) spectrum recorded at pH 2.65 showed 156 peaks, of which 145 were unambiguously assigned to BLGA residues (Figure S8). Due to overlap ambiguities and lack of assignment in the reported shift list (L22, E157, E158, I162), nine residues of BLGA were left unassigned. Importantly, the assigned chemical shifts were in good agreement with those previously reported for BLGA (Figure S9, Tables S2, S3), indicative of correctly folded recombinant BLGA WT.

To map where the oligosaccharides bind on the monomer, BLGA was titrated with AOSs at pH 2.65, reaching a molar concentration of 15.5 mM (Figure 3A). Although distinct chemical shift perturbations were seen, the amplitudes of the shifts were generally small. Due to weak binding, saturation was not achieved and fitting to a classical binding model was not possible (Figure S10). Still, chemical shift perturbation analysis by addition of AOSs (Figure 4) identified affected residues particularly in three surface-exposed regions, D11–K14 (N-terminal region), K101–Y102 (FG-loop), and E127–D137 (dimerization α-helix) (Figure 1). Single affected products were separated by high-performance liquid chromatography (HPLC) (see Materials and Methods), resulting in 12 mg of trisaccharides (AOSs) from 100 mg of alginate (Figure S1). Structural analysis by NMR revealed a mixture of two AOSs with G at the reducing end. The dominant species (1) (75%) had G as the central residue, whereas the minor species (2) (25%) had M (Figure 2). As the lyase removes the 4-OH group and the C-5 proton by β-elimination, the C5 epimers β-D-mannuronic acid and α-L-guluronic acid lead to the same nonreducing end product, 4-deoxy-α-L-erythro-hex-4-enopyranosyluronic acid (Figure S1). Structural analysis by NMR revealed a mixture of two main species (1) (75%) had G as the central residue, whereas the minor species (2) (25%) had M (Figure 2). As the lyase removes the 4-OH group and the C-5 proton by β-elimination, the C5 epimers β-D-mannuronic acid and α-L-guluronic acid lead to the same nonreducing end product, 4-deoxy-α-L-erythro-hex-4-enopyranosyluronic acid (Figure 2; Table S1; Figures S2–S7). The structure of these compounds has previously been solved by NMR, and the assigned chemical shifts are in good correlation with previous results.

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**RESULTS**

**Structure Determination of AOSs, an Alginate Model.** Alginate oligosaccharides were prepared by degradation using endoacting alginate lyase, and the di- and trisaccharide end products were separated by high-performance liquid chromatography (HPLC) (see Materials and Methods), resulting in 12 mg of trisaccharides (AOSs) from 100 mg of alginate (Figure S1). Structural analysis by NMR revealed a mixture of two AOSs with G at the reducing end. The dominant species (1) (75%) had G as the central residue, whereas the minor species (2) (25%) had M (Figure 2). As the lyase removes the 4-OH group and the C-5 proton by β-elimination, the C5 epimers β-D-mannuronic acid and α-L-guluronic acid lead to the same nonreducing end product, 4-deoxy-α-L-erythro-hex-4-enopyranosyluronic acid (Figure 2; Table S1; Figures S2–S7). The structure of these compounds has previously been solved by NMR, and the assigned chemical shifts are in good correlation with previous results.

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**Figure 1.** NMR structure of BLGA at pH 2.65 (PDB: 1DV9). β-strands A (green), B (yellow), C (purple), D (orange), E (gray), F (green-gray), G (blue), H (light yellow), and I (pink); dimerization α-helix (white); and the EF-loop (magenta) in the closed conformation. 7

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**Figure 2.** Structures of alginate trisaccharides (AOSs). 1 is the major component (~75%) (4-deoxy-α-L-erythro-hex-4-enopyranosyluronic acid)-(1,4)-(α-L-guluronic acid)-(1,4)-(α-L-gulopyranosyluronic acid); 2 is the minor component (~25%) (4-deoxy-α-L-erythro-hex-4-enopyranosyluronic acid)-(1,4)-(β-D-mannopyranosyluronic acid)-(1,4)-(α-L-gulopyranosyluronic acid).
residues outside these regions were also observed, including S30 (AB-loop), K75 (β-strand D), I84 (FG-loop), and L149 (β-strand I), as well as perturbation of three unassigned peaks (Figure 4, Table 1). Although most of the regions and residues directly contain or have neighboring positively charged residues, mainly lysine, there were several lysines on the BLGA surface that remained unperturbed, e.g., K39, K47, K48, K60, K69, and K70.

Based on these observations, two binding sites, site 1 and site 2, were defined, which, together with the anionic character of AOSs, motivated the production of BLGA single variants, K8A, K75A, and K101A, to investigate the effect on binding by removing a specific positive charge. K8A was chosen based on its location spatially close to K14 and K75 and because it was unassigned in our work due to overlap. K8 was thus hypothesized to be one of the three unassigned spin systems undergoing chemical shift perturbations by addition of AOSs. In the 1H,15N-HSQC spectrum of BLGA K8A, some peak overlap was observed, but it still allowed for K8 assignment (Figure 5A). Notably, after successful assignment, it was clear that K8 was unperturbed at pH 2.65 by addition of AOSs (Figure 4).

The conformational integrity of unlabeled recombinant BLGA WT and mutant proteins was assessed by far-UV circular dichroism (CD). As expected for the β-sheet-rich BLGA,31 a global minimum at 216−218 nm was observed and only very small differences from WT were seen for K8A and K75A (Figure S11). By contrast, the fold of the K101A variant was clearly compromised as deduced from the shifted CD minimum (Figure S11) in agreement with extensive peak collapse in the 1H,15N-HSQC spectrum, indicative of a partially unfolded structure (Figure 5C). AOS binding analysis was therefore not pursued for BLGA K101A. At pH 2.65, AOSs elicited a similar chemical shift perturbation for K8A, K75A, and WT BLGA (Table 1). Thus, not one single residue appeared to be solely responsible for the binding of AOSs to BLGA.

Two AOS binding sites were identified on the surface of monomeric BLGA at pH 2.65, site 1 and site 2 (Figure 4, see also Figure 7). Except for S30 (AB-loop) and L149 (β-strand I) (Table 1, Figure 4), most of the involved residues of BLGA are situated outside the dimerization interface, i.e., the AB-loop (S27−V41) and β-strand I (I147−F150).32 Subsequently, the influence of pH on AOS binding was investigated at pH 4.0 where BLGA is mostly in the dimeric form.10

**pH Tunes Alginate Binding to BLGA and Controls Coacervation.** The effect of BLGA dimerization on the binding of AOSs in a dairy-relevant pH range was investigated by chemical shift perturbation at pH 4.0.33 First, BLGA resonances were assigned at pH 4.0 by titration from pH 2.65 to 3.2 and 4.0 (Figure 3C; Figure S12, Tables S2, S4). Here, at pH 4.0, we obtained unambiguous assignments of 132 residues, compared to 145 residues at pH 2.65. As expected, this increase in pH caused BLGA to dimerize, which seriously affected line widths for several peaks, resulting in assignments of only 132 residues (Table S4, BMRB no: 27668). Mapping these missing assignments onto the BLGA structure clearly showed that residues with broadened peaks located distinctly to the known dimerization interface (Figure 7C). Next, AOSs were added by titration and chemical shift perturbations were analyzed (Figures 6 and 3B). Compared to the BLGA monomer, distinct differences were observed. Notably, Y102 in site 2 was no longer affected by AOS binding, and K101 was less affected at pH 4.0 than at pH 2.65 (Figure 6). By contrast, K75 in site 1 retained a significant chemical shift perturbation at pH 4.0. Moreover, residues belonging to the α-helix contributing to the dimerization interface were unperturbed by
Turbidimetry AOS-induced chemical shift perturbations for both K8A and K75A are similar to those of the free amino acids (Glu, pK\textsubscript{a} 2.65 and 4.07, respectively). Notably, the buried A80 was affected by binding, which may be due to changes in K8, which is located in the flexible, disordered N-terminal region\textsuperscript{,}\textsuperscript{5,6} shielding A80. At pH 4.0, K8 was clearly involved in AOS binding. Taken together, these comparative data highlight that AOS exploits the surface of BLGA in a pH-dependent manner and that the monomer and the dimer of BLGA bind differently to AOSs.

Mapping AOS-induced chemical shift perturbations at pH values 2.65 and 4.0 to the surface of BLGA identified two areas at pH 2.65 (Figure 7), site 1 constituted by D11, E12, K14, and K75 (BLGA N-terminal region and DE-loop) and site 2 constituted by K101, Y102, E126, D128, and D129 (FG-loop and H-dimerization \(\alpha\)-helix loop). Notably, at pH 4.0, residues of site 2 surrounding K101 appeared unaffected by addition of AOSs and the shift for K101 was of much smaller amplitude than at pH 2.65. A change in pH to 4.0 is assumed to affect the protonation state of the acidic side chains. Using PROPKa\textsuperscript{,}\textsuperscript{34} the pH change is suggested to partially deprotonate E126, D128, and D129. The pK\textsubscript{a} values calculated for these residues in the monomeric (PDB 1DV9) BLGA were D128, 2.52; D129, 4.79; and E126, 4.80 and for dimeric (PDB: 1EB) BLGA, D128, 3.69; D129, 4.11; and E126, 4.18. Considering the prediction accuracy of PROPKa, we note that these values are similar to those of the free amino acids (Glu, pK\textsubscript{a} 4.07 and Asp, pK\textsubscript{a} 3.90). Overall, it is reasonable to assume partial deprotonation at pH 4.0. This alters the electrostatic surface of site 2, rendering it binding-incompetent due to charge–charge repulsion between BLGA and the anionic AOSs. Moreover, E126, D128, and D129 are located close to the dimerization interface. At pH 4.0, site 1 surrounding K75 also changes, as K14 no longer contributes, while K8, A80, K83, F82, and D53 appear affected.

Turbidity measurements on BLGA and alginate have been shown to gauge coacervate formation.\textsuperscript{18,20} Since the change in AOS-induced chemical shift perturbations for both K8A and K75A was indistinguishable from WT (Table 1), turbidimetry was performed on BLGA WT, K8A, and K75A at varying alginate concentrations to confirm that the mutations have little effect on the ALG/BLGA interaction. K8A and K75A BLGA gave marginally less protein remaining in solution than WT, albeit BLGA WT at the higher alginate concentrations displayed increased turbidity compared to the two variants (Figure 8). The reason for this may be lower affinity of BLGA variants for alginate, resulting in smaller and/or fewer particles than with WT. The mutant proteins may also have lower solubility due to lower surface charge.

**DISCUSSION**

BLG is known to exploit different binding sites for a wide range of molecules spanning from hydrophobic ligands accommodated inside the calyx\textsuperscript{5,6} to vitamins binding on the surface.\textsuperscript{5} The presently identified AOS binding sites are distinct from previously reported binding areas on the surface of BLGA and the first accommodating carbohydrates.\textsuperscript{35} Notably, a subset of the lysines is consistently involved in AOS binding (K14, K75, K83, K101 at pH 2.65 and K8, K75, K101 at pH 4.0). However, many of the 15 surface-exposed lysines appear unaffected by the presence of AOSs. This highlights that the probed interactions are specific and not a mere electrostatic effect, which is also supported by the mutational removal of a single cationic side chain having little effect on binding, suggesting that other types and not just electrostatic forces are involved in the interaction. Some of the unaffected lysines may engage in local surface electrostatics of their immediate environment, in this way preventing interaction with AOSs. BLG has previously been shown to undergo a conformational change to a more \(\alpha\)-helical structure induced by anionic lipids at pH values 4.6 and 2.6.\textsuperscript{35} However, addition of AOSs did not elicit large changes in the NMR spectra, indicative of helix formation (Figure 3). K\textsubscript{d} values of AOS binding to BLGA were determined previously by isothermal titration calorimetry (ITC) to 1.1 and 0.6 mM at pH 3.0 and 4.0, respectively.\textsuperscript{18} However, none of the present chemical shift perturbations reached saturation to allow for determination of K\textsubscript{d} (Figure S10). Presumably, this is due to the 5-fold higher ionic strength used in the NMR compared to the ITC experiments, weakening the electrostatic interactions between AOSs and BLGA.\textsuperscript{16,18,36}

Binding of aroma compounds onto the surface of BLG at pH 2.0 was described by NMR,\textsuperscript{37} identifying two hydrophobic binding sites, one for \(\gamma\)-decalactone involving K47, L57, K70, and I72 and one for \(\beta\)-ionone involving K60, Y102, L104, and E129.\textsuperscript{37} Both sites are located in the hydrophobic region between \(\beta\)-strand G, the dimerization \(\alpha\)-helix, and \(\beta\)-strand I. The positions of these binding sites are distinctly different from those determined for the AOSs, although Y102 is shared. Notably, BLG is reported to catalyze lipofuscin formation using K60 and K69 as well as K77 and K91 as catalytic
residues.\(^{38}\) Also, none of these residues were found to interact with the AOSs, even though side chain assignments are needed to fully confirm this conclusion.

Charged patches on BLG were previously suggested to engage in interaction with anionic polysaccharides,\(^{13,15,16,18,20}\) and binding regions for anionic polysaccharides were hypothesized to include residues A1–K14, V41–K60, T76–K83, and A132–R148.\(^{15,16}\) Notably, at pH 4.0, the AOS-perturbed residues in BLGA are K8, G9, D11, D53, K75, A80, F82, K83, and D85, whereas no residues from the stretch V41–K60 underwent noteworthy chemical shift perturbation, except the surface-exposed D53 and the buried V41 and L57 (Figure 6). At pH 2.65, two AOS binding areas are identified on BLGA, of which site 1 remains as the sole, albeit rearranged, binding site at pH 4.0. However, the AOSs do bind near residues from the segments A1–K14 and K75–D85 at both pH 4.0 and 2.65, suggesting these regions to be the main sites. In addition, at pH 2.65, significant chemical shift perturbation occurred in the region A132–R148 as well as around K101–Y102 in site 2, but these disappeared at pH 4.0 (Figure 6). The observation of two binding sites at pH 2.65 is in agreement with previous ITC stoichiometry of two BLGA AOS binding sites at pH 3.0 and one at pH 4.0.\(^{18}\)

The charged state of AOS is very important for a valid extrapolation of the AOS binding sites to alginate/BLGA interaction and subsequent coacervation. The pKa values of M and G monosaccharides are 3.38 and 3.65, respectively.\(^{39}\) Thus, there is a possibility that AOS could be neutral at pH 2.65. The individual pKa values of the acid groups in the sugar units and the charged state of the molecule were predicted using Chemicalize. At pH 2.65, it is predicted that 64.21% of the population is charged (25.2% \(-2\), 39.01% \(-1\)), and at pH 4, 100% of the molecules are charged (77.55% \(-3\), 21.11% \(-2\), 1.34% \(-1\)). Thus, the AOS should be charged under all conditions used in this study and the results thus enable extrapolation to alginate binding. Site 1 is also affected at both pH 2.65 and 4, indicative of ionic contribution in the interaction at both pHs. The presence of the double bond at the nonreducing end of the trisaccharides is a consequence of the enzymatic degradation as no alginate hydrolase has been discovered to date.\(^{40}\) It will cause a change in the sugar ring that goes from being in a chair conformation to a boat conformation. This combined with short trisaccharides will have significant border effects that are not found with the polysaccharide. The AOS, however, is the only alginate oligosaccharide published to date that does not form insoluble particles with BLGA, while the interaction with BLGA remains.\(^{18}\) This is why AOS was chosen as a model system for NMR studies. Enzymatic degradation affords a greater control over the polymerization as opposed to partial acid hydrolysis and was chosen for this reason.

AOSs are a mixture of two anionic trisaccharides (Figure 2), which are expected to interact with positively charged residues such as lysines as found for other anionic oligosaccharide protein complexes.\(^{41}\) In BLGA also, hydrophobic residues undergo chemical shift perturbations including Y102 at pH 2.65 and L10, A80, and F82 at pH 4.0. This is also supported by the little difference observed by mutating K8 and K75 in the interaction sites (Table 1 and Figure 8). This suggests that other types of interactions are involved in AOS binding to BLGA, thus going beyond what is classically seen for oppositely charged flexible polymers.\(^{42}\) This could be a consequence of BLGA having a well-defined tertiary structure.

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\(\text{[x]}\) indicates peaks lost at pH 4.0.
at pH values 2.65 and 4.0 \cite{10,43} and therefore charged or hydrophobic residues cannot move freely relative to the binding area upon interaction with AOSs.

The mutational analysis of alginate/BLGA binding/coacervation did not reveal big differences between the mutant proteins and WT (Figure 8). However, the observed differences in the alginate concentration required to reach a saturation could indicate a change in binding stoichiometry; something that will be needed to be addressed in later studies.

Overall, these results identify binding sites on BLGA for anionic oligosaccharides and hence also polysaccharides, which are dependent on the pH and thus on the oligomeric state of BLGA. Through binding of longer polysaccharides, this forms the basis for coacervate formation. Thus, the pH dependence is one way to regulate the behavior of the coacervates, opening up for the design of polyelectrolyte carbohydrates to achieve desirable coacervate behavior.

\section*{CONCLUSIONS}

Two surface-exposed AOS binding sites on monomeric BLGA centered around K75 (site 1) and K101 (site 2) were identified. The sites are distinctly different from previously reported binding sites on BLG for other compounds. Notably, in the dimeric BLGA, only site 1 is functional at pH 4.0. Although the AOSs/BLGA interaction is electrostatically driven, the binding sites are specific and not structurally dynamic, suggesting that other types of important interactions occur, e.g., stacking interactions between monosaccharide rings and aromatic side chains. A further scrutiny of these interactions by mutations can help elucidate their importance including their influence on macromolecular particle formation. Overall, the distinct BLGA binding sites and their pH dependence are fundamental characteristics that can help guide design of polyelectrolyte carbohydrates to achieve desirable coacervate behavior.

\section*{MATERIALS AND METHODS}

\subsection*{Preparation of Alginate Oligosaccharides (AOSs).}

AOSs were prepared essentially as described.\cite{18} Briefly, 10 mg mL$^{-1}$ alginate ($M_n = 40$ kDa, $M/M_0$ ratio = 0.6, and polydispersity = 2.6; kind gift of Finn Madsen, DuPont Nutrition Biosciences, Brabrand, Denmark) was incubated with endoacting alginate lyase from \textit{Sphingomonas} sp. (Megazymes, Ireland) at 42 °C for 6.5 h, followed by enzyme inactivation (90 °C, 10 min), centrifugation (20 000 g, 10 min), and desalting of the supernatant into water (Hiprep Desalt 26/10; GE Healthcare). Oligosaccharide products were monitored spectrophotometrically at 235 nm.\cite{18,44} The fractions containing AOSs were added acetonitrile to 50% (v/v) and purified by HPLC (TSKgel Amide-80 column 5 μm particle size; 4.6 by 250 mm with 4.6 × 10 mm guard column (Tosoh, Japan); Ultimate 3000 HPLC (Dionex, CA) equipped with RI-101 refractive index detector (Showa Denko, Japan)) eluted by 70% (v/v) acetonitrile in water as mobile phase at 70 °C at a flow rate of 1 mL min$^{-1}$. AOSs were quantified using the phenol sulfuric acid method.\cite{18,44} Purity was assessed by thin layer chromatography (TLC; silica gel 60 F$_{254}$; Merck) 2 × 3
μL spotted effluent, developed twice in 50% butanol/25% acetic acid/25% MilliQ water and visualized by taring (300 °C) in 10% sulfuric acid, 80% ethanol, 8% H2O, and 2% orcinol (Supporting Information, Figure S1).

**Figure 7.** Mapping of AOS-induced chemical shift perturbations (blue) onto the surface of monomeric BLGA WT (PDB: 1DV9) at (A) pH 2.65 and (B) pH 4.0. (C) Residues assigned to peaks lost during the pH titration (pH 2.65 to 3.2 to 4.0) are in red and also mapped to the BLGA dimer (PDB: 1BEB).

pKₐ values of the acid groups in AOSs were predicted by submitting the structure to chemicalize.com (ChemAxon, 06/02-2019).

**BLGA Cloning, Mutagenesis, and Pichia pastoris Transformation.** The vector pPICZαA harboring the gene encoding BLGA (UniProt: P02754) was optimized for P. pastoris to produce a recombinant BLGA variant with the N-terminal extension EAE (here numbered −1 to −3) and three point mutations, changing L1I2 to A1,Y2 and the V105F (Figure S1). These changes were done to enable transfer of assignments reported previously for structure determination of BLGA by NMR. The full primary structure of this BLGA variant is given in the Supporting Information. The DNA sequence encoding the BLGA V105F (referred to as BLGA) was cloned in-frame with the *Saccharomyces cerevisiae* α-mating factor to give pPICZαA−BLGA (purchased from GeneArt; ThermoFisher). An XhoI restriction site was introduced after the α-factor gene to avoid additional amino acids to remain after cleaving off the α-factor. Single BLGA mutants K8A, K75A, and K101A were made by site-directed mutagenesis (Quick Change lightning mutagenesis kit; Agilent) using the primers (purchased from Eurofins Genomic; Germany) K8A 5′cgtccacccagacccgtagttgctgccga3′; K8A_anti 5′tggatatcctgttgcgactac3′; K75A 5′gtgcctgcaacctgatcttctg3′; K75A_anti 5′cagggatcttggttgcttctgctt3′; K101A 5′ggtggttggacaccgactac3′, and K101A_anti 5′tggtgtgtgcttgtgtgtg3′.

Figure 8. Turbidity (600 nm) of BLGA WT and variants mixed with alginate at pH 4.0 and absorbance (280 nm) of supernatants after centrifugation (see Materials and Methods): BLGA WT (circle), K75A (square), and K8A (triangle).
*catcagaaacaaacctagctggtttagctgggtgccaaaccc3′. Wild-type (WT) and mutant plasmids were transformed into *Escherichia coli* DH5α by heat shock and selected for zeocin resistance (Novagen, U.K.), and the mutations were confirmed by sequencing (GATC Biotech, Germany). pPICZαA–BLGA WT and mutant plasmids were linearized using *PmeI* and transformed into *P. pastoris* X-33 by electroporation (EasySelect Pichia Expression Kit; Invitrogen). Transformants were selected on zeocin containing yeast extract peptone dextrose medium (BMM: 0.1 M K2HPO4/KH2PO4 (KPi) pH 6.0, 0.34% (w/v) yeast nitrogen base without amino acids and ammonium chloride, 4 × 10−5% (w/v) biotin, 1% ammonium sulfate, 0.5% (v/v) glycerol), growing for 5 days at 22 °C evaluated on aliquots removed with 24 h intervals and analyzed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). The transformants secreting the highest amount of protein were propagated in 25 mL of buffered glycerol-complex medium (BMGY: 1% (w/v) yeast extract, 2% (w/v) peptone, 0.1 M KPi pH 6.0, 0.34% (w/v) yeast nitrogen base, 4 × 10−5% (w/v) biotin, 1% (v/v) glycerol) (30 °C, 150 rpm) until OD600 = 2–6, inoculated into 1 L BMGY medium in baffled 3 L shake flasks, and grown until OD600 = 2–6. Cells were harvested by centrifugation (1500 g, 5 min, 22 °C), resuspended in 1 L BMGY medium, incubated 110 h (22 °C, 150 rpm), and methanol being added to 0.5% (v/v) every 24 h. Cells were pelleted (10 000 g, 30 min, 4 °C), and the supernatant was filtered (0.45 μm) and concentrated by cross-flow filtration (SARTOFLOW Slice 200 Benchtop System, 5 kDa Hydrostart ultrafiltration cassette at 4 °C, Sartorius, Germany). Protein concentration was determined spectrophotometrically using a predicted molar extinction coefficient of 18 700 M−1 cm−1 (Protparam).45 Conformational integrity of BLGA variants was confirmed by circular dichroism spectroscopy. For NMR analysis 15N-labeled BLGA was produced essentially as described above in 2 L of BMGY (0.1 M KPi pH 6.0, 0.34% (w/v) yeast nitrogen base without amino acids and ammonium chloride, 4 × 10−5% (w/v) biotin, 1% 15N-ammonium sulfate (Cambridge Isotope Laboratories Inc., Andover), 0.5% (v/v) glycerol). Purification of Recombinant BLGA WT and Mutants. All steps were performed at 4 °C. Concentrated culture supernatant was buffer-exchanged to 50 mM KPi pH 6.0, 150 mM NaCl (Hiprep Desalt column 26/10; GE Healthcare) at a flow rate of 2 mL min−1 and concentrated (3 kDa cutoff, Amicon Ultra 15 centrifugal filter; Merck). Recombinant BLGA WT and mutant proteins were purified by size exclusion chromatography (HiLoad Superdex 75 26/60; GE Healthcare) at a flow rate of 1.4 mL min−1 in 50 mM KPi pH 6.0 and 150 mM NaCl and dialyzed (MilliQ water, 3 kDa cutoff Spectra/ Por membrane; Spectrumlabs) by 3 × 100-fold dilution, each for 4 h, evaluated by SDS-PAGE to be >95% pure (data not shown), lyophilized, and stored at −20 °C until use. For NMR experiments, proteins were dissolved in 55 mM KPi pH 2.65, 3.2, or 4.0 and dialyzed against the buffer (as above). Yields of purified BLGA WT, K8A, K75A, and K101A were 40.4, 20.6, 35.2, and 6.3 mg, respectively, per liter of culture containing 15N-ammonium sulfate.

**Production of Isopeptide-Labeled Recombinant BLGA.** Eight clones from each *P. pastoris* transformation were restreaked on YPD agar zeocin plates and selected for expression level in 50 mL of buffered minimal methanol medium (BMM: 0.1 M K2HPO4/KH2PO4 (KPi) pH 6.0, 0.34% (w/v) yeast nitrogen base without amino acids and ammonium chloride, 4 × 10−5% (w/v) biotin, 1% ammonium sulfate, 0.5% (v/v) glycerol), growing for 5 days at 22 °C evaluated on aliquots removed with 24 h intervals and analyzed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). The transformants secreting the highest amount of protein were propagated in 25 mL of buffered glycerol-complex medium (BMGY: 1% (w/v) yeast extract, 2% (w/v) peptone, 0.1 M KPi pH 6.0, 1.34% (w/v) yeast nitrogen base, 4 × 10−5% (w/v) biotin, 1% (v/v) glycerol) (30 °C, 150 rpm) until OD600 = 2–6, inoculated into 1 L BMGY medium in baffled 3 L shake flasks, and grown until OD600 = 2–6. Cells were harvested by centrifugation (1500 g, 5 min, 22 °C), resuspended in 1 L of BMM medium, incubated 110 h (22 °C, 150 rpm), and methanol being added to 0.5% (v/v) every 24 h. Cells were pelleted (10 000 g, 30 min, 4 °C), and the supernatant was filtered (0.45 μm) and concentrated by cross-flow filtration (SARTOFLOW Slice 200 Benchtop System, 5 kDa Hydrostart ultrafiltration cassette at 4 °C; Sartorius, Germany). Protein concentration was determined spectrophotometrically using a predicted molar extinction coefficient of 18 700 M−1 cm−1 (Protparam).45 Conformational integrity of BLGA variants was confirmed by circular dichroism spectroscopy. For NMR analysis 15N-labeled BLGA was produced essentially as described above in 2 L of BMGY (0.1 M KPi pH 6.0, 0.34% (w/v) yeast nitrogen base without amino acids and ammonium chloride, 4 × 10−5% (w/v) biotin, 1% 15N-ammonium sulfate (Cambridge Isotope Laboratories Inc., Andover), 0.5% (v/v) glycerol).

**Circular Dichroism Spectroscopy.** Lyophilized BLGA WT and mutant proteins were dissolved in 10 mM NaPi pH 7.0, centrifuged (20 000 g, 20 min, 4 °C), and dialyzed (3 kDa cutoff, Spectra/Por membrane; Spectrumlabs) by 3 × 100-fold dilution, each against the buffer for >4 h, at 4 °C. Far-UV CD spectra were recorded at 250−190 nm in a 1 mm quartz cuvette (50 nm min−1, 1 nm bandwidth, 2 s response time, 25 °C; Jasco J810 Spectropolarimeter, Peltier controlled). Ten scans were averaged, and a buffer background recorded using identical parameters was subtracted. The molar ellipticity was calculated by

\[
[\theta] = 100 \times \frac{\theta}{m} \times d
\]

where *m* is the molar protein concentration, *θ* is the ellipticity at wavelength *λ*, and *d* is the path length in centimeter.48

**NMR of BLGA WT and Mutant Proteins.** 15N-BLGA WT and mutant proteins were diluted to 50 mM KPi H2O/D2O (9/1 v/v) and 0.1 mM 2,2-dimethyl-2-silapentane-5-sulfonic acid (DSS) and centrifuged (12 000 g, 20 min, 4 °C) prior to analysis by NMR spectroscopy in 5 mm Shigemi microtubes. The buffer was chosen to match the conditions used by Uhrinova et al.5 and to ensure consistent pH values outside the main buffering range of the buffer. Upon addition of ligand, pH was corrected in all samples prior to the measurement and measured immediately after, to secure consistency. BLGA WT is monomeric48 and stable for many days at pH 2.65,5 while a significant amount of dimer is present at pH values 3.2 and 4.0.10 The H1, 15N-heteronuclear single quantum coherence (HSQC) spectra were recorded on an 800 MHz Varian INOVA spectrometer equipped with a 5 mm triple resonance room temperature probe with a Z-field gradient at 37 °C46 using a Varian/Agilent BioPack sequence or on a 600 MHz Bruker AVANCE system equipped with a cryoprobe. Recorded free induction decays were processed with nmrPipe.47 Proton chemical shifts were referenced to internal DSS at 0.00 ppm and indirectly using gyromagnetic ratios for 14N and 13C chemical shifts.

**NMR Assignment and Chemical Shift Perturbation Analysis.** Assignments of N and H20 nuclei of BLGA at pH 2.65 were done using the assignment list of BLGA obtained from triple resonance spectra recorded at exactly the same conditions as those for template.7 To transfer the assignments to pH 4.0, 1H,15N-HSQC NMR spectra of 400 μM BLGA WT were recorded at pH values of 2.65, 3.2, and 4.0, and the signals were followed during the titration to allow assignments at pH 4.0. These assignments are deposited in the Biological Magnetic Resonance Data Bank (BMRB, http://www.bmrb.wisc.edu/) entry 27668. For mapping of binding sites, a series of 1H, 15N-HSQC spectra were recorded for 100 μM BLGA WT or mutant protein and 15.5 mM AOSs at pH 2.65 or pH 4.0 prior to mixing in five steps with a BLGA sample not containing AOSs to give seven spectra in total recorded per protein sample at 0, 2.6, 5.2, 7.8, 10.4, 12.9, and 15.5 mM AOSs. Chemical shifts were in fast exchange and could be followed during the titration. Chemical shift perturbations were calculated to map the binding site on the surface of the protein by

\[
\Delta \delta = \sqrt{(\Delta \delta H)^2 + (\Delta \delta N/S)^2}
\]

where \(\Delta \delta H = \Delta \delta H_{\text{free}} - \Delta \delta H_{\text{obs}}\) and \(\Delta \delta N = \Delta \delta N_{\text{free}} - \Delta \delta N_{\text{obs}}\). Changes in the proton and nitrogen chemical shifts (in
Significant chemical shift perturbation was defined as the average chemical shift perturbation plus one standard deviation.

**Turbidimetry.** The formation and assessment of solubility of BLGA/alginate particles were done as previously described. Briefly, alginate, BLGA WT, or mutant proteins were dissolved in 50 mM NaPi pH 4.0. BLGA (1 mg mL⁻¹ or 54 μM) was mixed with alginate (0–0.2 mg mL⁻¹ or 0–5 μM using Mₙ = 40 kDa), incubated for 10 min at room temperature, and the turbidity was measured at 600 nm. Subsequently, the mixtures were centrifuged (20 000g, 20 min, 20 °C) and remaining soluble protein was determined spectrophotometrically at 280 nm.

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**REFERENCES**


