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Highlights

- Surface plasmon resonance was used to characterize HePS-milk protein interactions
- Binding affinity depends on HePS size, repeat structure, pH and ionic strength
- Heat-treatment of β-lactoglobulin dramatically increased HePS binding capacity

ABSTRACT

Interactions of exopolysaccharides and proteins are of great importance in food science, but complicated to analyze and quantify at the molecular level. A surface plasmon resonance procedure was established to characterize binding of seven structure-determined, branched hetero-exopolysaccharides (HePSs) of 0.14–4.9 MDa from lactic acid bacteria to different milk proteins (β-casein, κ-casein, native and heat-treated β-lactoglobulin) at pH 4.0–5.0. Maximum binding capacity (RU\textsubscript{max}) and apparent affinity (\(K_{A,app}\)) were HePS- and protein-dependent and varied for example 10- and 600-fold, respectively, in the complexation with native β-lactoglobulin at pH 4.0. Highest RU\textsubscript{max} and \(K_{A,app}\) were obtained with heat-treated β-lactoglobulin and β-casein, respectively. Overall, RU\textsubscript{max} and \(K_{A,app}\) decreased 6- and 20-fold, respectively, with increasing pH from 4.0 to 5.0. \(K_{A,app}\) was influenced by ionic strength and temperature, indicating that polar interactions stabilize HePS–protein complexes. HePS size as well as oligosaccharide repeat structure, conferring chain flexibility and hydrogen bonding potential, influence the \(K_{A,app}\).

Keywords:
Hetero-exopolysaccharides (HePSs), β-lactoglobulin, β- and κ-casein, binding parameters, surface plasmon resonance (SPR), dynamic light scattering (DLS)

Abbreviations:
BCN, β-casein; BLG, β-lactoglobulin; DLS, dynamic light scattering; EPS, exopolysaccharide; Gal, galactose; GalNAc, N-acetylgalactosamine; Glc, glucose; GlcNAc, N-acetylglucosamine; hBLG, heat-treated BLG; HePS, hetero-exopolysaccharide; HoPS, homo-exopolysaccharide; KCN, κ-casein; LAB, lactic acid bacteria; nBLG, native BLG; pI, isoelectric point; Pyr, pyruvate; Rha, rhamnose; SEC, size-exclusion chromatography; SNR, signal-to-noise ratio; SPR, surface plasmon resonance.

1. Introduction

Homo- and hetero-exopolysaccharides (HoPSs and HePSs) are secreted by a wide variety of Gram-positive and Gram-negative bacteria, yeasts, molds, and microalgae (Donot, Fontana, Baccou, & Schorr-Galindo, 2012). HoPSs produced from sucrose by extracellular glucan- and fructansucrases (Monsan, Bozonnet, & Albenne, 2001; Zannini, Waters, Coffey, & Arendt, 2015) contain either D-glucopyranose or D-fructofuranose residues connected by a few glycosidic bond types (Laws, Gu, & Marshall, 2001), while HePSs are complex and synthesized intracellularly from nucleotide sugars catalyzed by glycosyltransferases yielding oligosaccharide repeats, which are polymerized and secreted as HePS (Schmid, Sieber, & Rehm, 2015; van Kranenburg, Vos, van Swam, Kleerebezem, & de Vos, 1999). The repeats contain 3–9 monosaccharide residues, typically glucose (Glc), galactose (Gal), rhamnose (Rha), N-acetylglucosamine (GlcNAc), N-acetylgalactosamine (GalNAc), or glucuronic acid, in either pyranose or furanose configuration (Broadbent, McMahon, Welker, Oberg, & Moineau, 2003; De Vuyst, De Vin, Vaningelgem, & Degeest, 2001). Glycerol, pyruvate (Pyr), phosphate, acetate, and
other substituents may occur (De Vuyst et al., 2003; Gruter, Leeflang, Kuiper, Kamerling, & Vliegenthart, 1993, 1992; Nakajima, Hirota, Toba, Itoh, & Adachi, 1992; Van Calsteren, Gagnon, Nishimura, & Makino, 2015; Van Calsteren, Pau-Roblot, Bégin, & Roy, 2002). The molar size of HePSs spans a wide range, and the structural diversity of oligosaccharide repeats is high with regard to monosaccharide composition, anomer configuration, regioselectivity, and substituents.

Exopolysaccharides (EPSs) can protect bacteria against changes in temperature, pH, light intensity, and other environmental stresses (for reviews see (De Vuyst et al., 2001; Donot et al., 2012; Freitas, Alves, & Reis, 2011; Hidalgo-Cantabrana et al., 2014; Patten & Laws, 2015)) and are commonly involved in formation and stabilization of biofilms (Sutherland, 2001). Lactic acid bacteria (LAB) produce HePSs in smaller amount than HoPSs (Torino, Font de Valdez, & Mozzi, 2015), but HePS nevertheless contribute very substantially in improving the physical and rheological properties of fermented milk products (Ayala-Hernández, Hassan, Goff, Mira de Orduña, & Corredig, 2008; De Vuyst & Degeest, 1999; Leroy & De Vuyst, 2004; Mende, Rohm, & Jaros, 2015). Interaction with milk proteins and water-binding capacity of HePSs have been connected with decreased syneresis, increased viscosity, firmness, creaminess, and a shiny surface of fermented dairy products (Folkenberg, Dejmek, Skriver, & Ipsen, 2005, 2006). Additionally, HePS–protein interactions may confer beneficial health effects in humans including antitumor activity (Wang et al., 2014), immunomodulation (Chabot et al., 2001; Hidalgo-Cantabrana et al., 2012), antioxidant properties (Zhang et al., 2013), anti-atherosclerotic activity (Tok & Aslim, 2010), antimutagenicity (Tsuda, Hara, & Miyamoto, 2008), and biosorption of lead (Feng, Chen, Li, Nurgul, & Dong, 2012).

Analysis of HoPS–milk protein interactions by surface plasmon resonance (SPR) has previously showed the binding capacity to decrease with pH increasing from 4.0 to 5.5 and to vary with HoPS linkage type, branching, and molecular size (Babol, Svensson, & Ipsen, 2011; Diemer et al., 2012). In
the present work, the SPR methodology was further developed to describe both HePS–protein binding capacity and apparent affinity. In this study, seven structurally diverse LAB HePSs (HePS-1–HePS-7) were used, all previously described in relation to viscosity regulation in yogurt fabrication (Bouzar, Cerning, & Desmazeaud, 1996; B Degeest, Mozzi, & De Vuyst, 2002; Doleyres, Schaub, & Lacroix, 2005; Kimmel, Roberts, & Ziegler, 1998; Robitaille et al., 2009), immune effect (Bleau et al., 2010; Lebeer, Claes, Verhoeven, Vanderleyden, & De Keersmaecker, 2011), or prevention of antibiotic-associated diarrhea (Vanderhoof et al., 1999). The aim of this study was to test the hypothesis that HePS size and structure determine the ability to bind milk proteins. This procedure can monitor effects of various environmental conditions on polysaccharide–protein interactions in a quantitative manner, disclose functional determinants of HePS–protein complex formation, and screen LAB strains relevant for application in food products as well as the capability to form non-food complexes, e.g. in biofilms.

2. Materials and methods

2.1. HePSs

HePS-1–HePS-7 were purified from culture supernatants of LAB strains, pre-cultured and fermented (20–48 h) in relevant media (Supplementary Table S1; Supplementary methods S1.1), and subjected to structure analyses (Supplementary methods S1.2 and S1.3). HePSs were quantified using the phenol–sulfuric acid method (Dubois, Gilles, Hamilton, Rebers, & Smith, 1956) with monosaccharide mixtures mimicking oligosaccharide repeat compositions as standards.

2.2. HePS molar mass ($M_w$)

Molar mass of HePS-1–HePS-7 was determined by size-exclusion chromatography (SEC) (solvent delivery system (LC-10AD), autosampler (SIL-10A), RI detector (RID-10A); all Shimadzu, Kyoto,
Japan) on an OH-PK SB-805HQ, 300 × 8 mm, pore size 500 Å column (Shoko CO., Ltd, Tokyo, Japan) in 10 mM sodium citrate/citric acid pH 4.0. Dextran standards (4.5 MDa, 1.45 MDa, 560 kDa, 350 kDa (American Polymer Standards Corporation, Mentor, OH, USA), 276.5 kDa, 196.3 kDa, 123 kDa (Pharmacosmos, Holbaek, Denmark)) and pullulan of 22 kDa were used for calibration. Standards (1.0–2.7 mg ml\(^{-1}\)) and HePSs (1.0–2.0 mg ml\(^{-1}\)) dissolved in the above buffer, degassed, kept overnight, and filtered (0.45 µm filters; Frisenette ApS, Knebel, Denmark) were analyzed (100 µl) at a flow rate of 0.5 ml min\(^{-1}\).
<table>
<thead>
<tr>
<th>HePS</th>
<th>LAB strain producing HePS with identical primary structure</th>
<th>Repeat unit (Da)</th>
<th>Molar mass (kDa)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>HePS-1</td>
<td>Lactococcus lactis ssp. cremoris NCIMB 700967</td>
<td>3391 ± 99</td>
<td>810.71</td>
<td>(Gruter et al., 1992)</td>
</tr>
<tr>
<td>HePS-2</td>
<td>Lactobacillus delbrueckii ssp. 372 ± 8</td>
<td>118.99</td>
<td>997.90</td>
<td>(Langersjö, Yang, Huttunen, &amp; Widimál, 2002)</td>
</tr>
<tr>
<td>HePS-3</td>
<td>Lactobacillus rhamnosus ATCC 53103</td>
<td>389 ± 5</td>
<td>1141.04</td>
<td>Streptococcus thermophilus</td>
</tr>
<tr>
<td>HePS-4</td>
<td>Lactobacillus delbrueckii CNRZ 1068</td>
<td>4924 ± 140</td>
<td>689.62</td>
<td>(Steinbüchel, Kamerling, &amp; Vliegenthart, 2001)</td>
</tr>
<tr>
<td>HePS-5</td>
<td>Lactobacillus delbrueckii CNRZ 291</td>
<td>3299 ± 379</td>
<td>810.71</td>
<td>Lactobacillus delbrueckii</td>
</tr>
<tr>
<td>HePS-6</td>
<td>Streptococcus thermophilus RW-9595M</td>
<td>136 ± 3</td>
<td>1141.04</td>
<td>(Van Calsteren et al., 2002)</td>
</tr>
<tr>
<td>HePS-7</td>
<td>Streptococcus thermophilus RD534</td>
<td>385 ± 10</td>
<td>648.57</td>
<td>(Lemoine et al., 1997)</td>
</tr>
</tbody>
</table>
Molar masses of HePS were determined as dextran equivalents by SEC (see 2.2 and Supplementary Fig. S2) and calculated for repeat units from the structures (see Fig. 1; Supplementary methods S1.2 and S1.3; Supplementary Fig. S1 and Supplementary results S2.1).
HePS $M$ values were obtained from the standard curve using a linear regression equation:

$$\log M = 9.9387 - 0.8712 \cdot V_e \quad \text{(eqn. 1)}$$

where $V_e$ is the elution volume ($r^2=0.992$).

### 2.3. Milk protein stocks

Lyophilized β-lactoglobulin (BLG), β- and κ-caseins (BCN and KCN) (all Sigma-Aldrich, St. Louis, MO, USA) were dissolved (1.0 mg ml$^{-1}$) in 10 mM sodium acetate pH 4.0, centrifuged (20000g, 4°C, 20 min) and the supernatants diluted (50 µg ml$^{-1}$) for sensor chip immobilization (see 2.5.1). BLG (5 mg ml$^{-1}$) in 50 mM sodium phosphate pH 6.8, 30 mM NaCl, was heat-treated (85°C, 15 min) (hBLG) and centrifuged (as above). The supernatant was dialyzed against 10 mM sodium acetate pH 4.0 and diluted (50 µg ml$^{-1}$) for immobilization (see 2.5.1). For dynamic light scattering (DLS) (see 2.4), BLG (5.0 mg ml$^{-1}$) was stirred in deionized water (150 rpm, 13 h, room temperature) and centrifuged (as above). Concentrations were determined spectrophotometrically at 280 nm using molar extinction coefficients ($\epsilon$) 17210, 11460, and 19035 M$^{-1}$ cm$^{-1}$ calculated by ProtParam (Gasteiger et al., 2005) from amino acid sequences for BLG, BCN, and KCN, respectively (GenBank accessions: BLG, 2Q2M_A; BCN, AAA30431; KCN, CAA33034).

### 2.4. Dynamic light scattering

Lyophilized HePS-3 in deionized water (3 mg ml$^{-1}$) was left overnight at 4°C for complete dissolution, filtered (0.45 µm filters; Frisenette ApS), mixed with BLG (see 2.3) to final 0.1 and 1.0 mg ml$^{-1}$, respectively, in 10 mM sodium citrate (pH 3–6) or Tris–HCl (pH 7–8) and equilibrated 30 min prior to DLS analysis. Particle sizes of HePS-3, BLG, and HePS-3+BLG mixtures were analyzed (23°C, scattering angle 90°; BI-200SM; Brookhaven Instruments Corporation, Holtsville, NY, USA)
and the distributions of mean apparent translational diffusion coefficients ($D_T$) determined by fitting the DLS autocorrelation functions (obtained with the Brookhaven system) using nonnegative constrained least squares. Distribution of mean apparent $D_T$ was converted to distribution of hydrodynamic diameter ($d_H$) using the Stokes-Einstein equation:

$$d_H = kT/3\pi\eta D_T \quad \text{(eqn. 2)}$$

where $k$ is the Boltzmann constant ($1.38 \cdot 10^{-23}$ J/K), $T$ the absolute temperature, and $\eta$ the solvent viscosity (0.93 mPa·s; assumed to be that of water at 296 K).

2.5. Surface plasmon resonance

2.5.1. Protein immobilization

Proteins (see 2.3) were covalently coupled to SPR sensor chips (CM5, CM4, C1; GE Healthcare, Uppsala, Sweden) using the automatic immobilization wizard (Biacore T100 Control Software; GE Healthcare) at 10 µl min$^{-1}$ and Amine Coupling Kit (GE Healthcare) for sample flow cell, while the reference flow cell underwent the same treatment without protein. CM4 and CM5 were prepared by 7 min injections of i) 0.05 M N-hydroxysuccinimide, 0.2 M 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide, ii) protein (50 µg ml$^{-1}$ in 10 mM sodium acetate pH 4.0), and iii) 1 M ethanolamine pH 8.5 to block remaining activated carboxyl groups. C1 was washed 2 × 1 min with 0.1 M glycine–NaOH pH 12.0, 0.3% (v/v) Triton X-100 followed by activation, immobilization, and blocking of activated carboxyl groups as above.

2.5.2. SPR analysis

Standard SPR analysis (25°C; Biacore T100; GE Healthcare) in running buffer of ionic strength similar to milk (10 mM sodium acetate pH 4.0, 70 mM NaCl, 0.005% surfactant P20; GE Healthcare)
(Babol et al., 2011) comprised 180 s association, 60 s dissociation, two consecutive 60 and 30 s regeneration cycles (10 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid pH 9.0, 1 M NaCl, 3 mM ethylenediaminetetraacetic acid, 0.005% surfactant P20) and 300 s equilibration (running buffer). Influence of pH (4.0–5.0) and NaCl (70–200 mM) was assessed. Temperature (25–35°C) effect was analyzed under standard conditions. Binding of HePS-1–HePS-7 was analyzed at seven concentrations (1–200 µg ml⁻¹) using 60 µl min⁻¹ to minimize mass transport effects. The change in refractive index caused by the accumulation of HePS interacting with the immobilized protein on the sensor surface is reported in response units (RU), where the sensorgram is in measured RU plotted versus time. After each HePS series, two samples below the critical HePS concentration (see 3.2.1) were analyzed to evaluate reproducibility. HePS-1 (10 µg ml⁻¹), showing robust and reproducible well-fitted sensorgrams, was used as a control to monitor the protein surface activity before and after each HePS interaction series. Reference cell and buffer blank responses were subtracted from sensorgrams to correct for nonspecific binding to the surface, mechanical drift, and external systemic noise (Myszka, 2000). Data were processed using Scrubber2 (Biologic Software Pty. Ltd., Campbell, Australia) and an in-house script (in MATLAB; The MathWorks, Natick, MA, USA) calculating steady-state response values at equilibrium (RU₂ₑₒₚ) by averaging the signal at 6–4 s prior to dissociation. Steady-state analyses adopted a 1:1 Langmuir interaction model (Myszka, 2000):

\[ RU_{\text{max}} = RU_{\text{eq}} \cdot C / (C + K_{D,\text{app}}) \]  

(eqn. 3)

RU₂ₑₒₚ is the steady-state response at HePS concentration C and \( K_{D,\text{app}} \) the apparent equilibrium dissociation constant. \( RU_{\text{max}} \) is the maximum HePS binding capacity of the protein surface. \( K_{D,\text{app}} \) and \( RU_{\text{max}} \) were calculated by plotting \( RU_{\text{eq}} \) against C and using nonlinear least-squares fitting routines.
(Myszka, 1997; O’Shannessy, Brighamburke, Soneson, Hensley, & Brooks, 1993). $K_{\text{app}}$, calculated as $1/K_{D,\text{app}}$, is the apparent equilibrium association constant.

3. Results and Discussion

3.1. HePS structure and size

HePS-1–HePS-7 oligosaccharide repeat structures (Fig. 1) were confirmed by $^1$H nuclear magnetic resonance spectra (Supplementary Fig. S1) judged to be identical to spectra of HePSs from authentic strains or in the literature (Table 1). Sugar compositions were verified for the rhamnose-containing HePS-2, HePS-3, and HePS-6 (Supplementary results S2.1), and the HePS-7 oligosaccharide repeat was prepared and characterized by mass spectrometry (Supplementary results S2.1). $M_w$ contributes to function (Cerning, Bouillanne, Landon, & Desmazeaud, 1992; De Vuyst & Degeest, 1999; van den Berg et al., 1995) and was determined by SEC (Table 1; Supplementary Fig. S2).
HePS-1

β-D-Galp1 →4)β-D-Galp(1→3)β-D-Galp ↓
→4)β-D-Galp(1→3)β-D-Galp(1→4)α-D-Galp(1→

HePS-2

β-D-Galp1 ↓ 3
β-D-Galp1 ↓ 4 α-L-Rhap ↓ 3
→2)α-D-Galp(1→3)β-D-GlcP(1→3)β-D-Galp(1→4)α-D-Galp(1→

HePS-3

β-D-Galp1 ↓ 6
→3)β-D-Galp(1→3)β-D-Galp(1→4)α-D-GlcP(1→3)α-L-Rhap(1→3)α-D-Galp(1→

HePS-4

α-D-Galp1 ↓ 6
→3)β-D-Galp(1→3)β-D-GlcP(1→3)α-D-GalpNac(1→

HePS-5

β-D-Galp(1→4)β-D-GlcP ↓ 6
→4)β-D-GlcP(1→4)α-D-GlcP(1→4)β-D-Galp(1→

HePS-6

(R)Pyr
/ /
4 6
α-D-Galp1 ↓ 2
→3)α-L-Rhap(1→3)β-D-GlcP(1→3)α-L-Rhap(1→3)α-L-Rhap(1→3)α-L-Rhap(1→2)α-D-GlcP(1→

HePS-7

β-D-Galp1 ↓ 6
→3)α-D-GlcP(1→3)β-D-GlcP(1→3)β-D-Galp(1→

Figure 1 Structures of HePS repeat units. HePS-1 (Lactococcus lactis ssp. cremoris NCIMB 700967), HePS-2 (Lactobacillus delbrueckii ssp. bulgaricus NCIMB 702483), HePS-3 (Lactobacillus rhamnosus GG (ATCC 53103)), HePS-4 (Streptococcus thermophilius CNRZ 1068), HePS-5 (Lactobacillus delbrueckii ssp. bulgaricus CNRZ 1187), HePS-6 (Lactobacillus casei LB31), and HePS-7 (Streptococcus thermophilus RD534).
3.2. SPR analysis of HePS–milk protein interaction

3.2.1. SPR protocol development

SPR analyses were previously used to determine binding between HoPS (0.5 mg ml\(^{-1}\)) and milk proteins immobilized at high density (RU=1130–4200) on a dextran-coated CM4 sensor chip (Babol et al., 2011; Diemer et al., 2012). Surprisingly, HePS-3 (0.5 mg ml\(^{-1}\)) exhibited more pronounced binding to the CM5 and CM4 reference cell compared to the native BLG (nBLG) sample cell, even at lower protein density (RU=395) than used in the HoPS study (Fig. 2A–B). Similar reference cell binding to CM5 and CM4 was observed for all HePSs analyzed in this study (data not shown).

![Figure 2](image-url)

**Figure 2** Sensorgrams of HePS-3 and native β-lactoglobulin (nBLG) immobilized on different SPR sensor chips. Reference cell: dashed line; sample cell: solid line; difference sensorgram: dotted line. A) HePS-3 (0.5 mg ml\(^{-1}\))
injected on CM5 with 395 RU immobilized nBLG; B) HePS-3 (0.5 mg ml$^{-1}$) injected on a CM4 sensor chip with 190 RU immobilized nBLG. C) 0.4 mg ml$^{-1}$ HePS-3 injected on a C1 chip with 75 RU immobilized nBLG. D) 0.04 mg ml$^{-1}$ HePS-3 injected on a C1 chip with 75 RU immobilized nBLG.

Both the CM5 and the CM4 sensor chips are dextran coated, with the CM4 chip having a shorter dextran layer and a lower carboxyl group density than the CM5 chip. Considering sample and reference cell responses as signal and noise, respectively, the signal-to-noise ratio (SNR) was 0.88 with nBLG CM5 and improved only 5% when using nBLG CM4. Remarkably, adverse HePS–dextran binding was avoided using the matrix-free C1 chip that raised the SNR by 36% at 0.4 mg ml$^{-1}$ and 150% at 0.04 mg ml$^{-1}$ HePS-3 (Fig. 2C-D) compared to CM5. The sensorgram shape clearly indicated rapid saturation of the C1 reference cell by HePS-3 and the binding to nBLG C1 approached steady state at the end of injection (Fig. 2D). Additionally, the density of immobilized protein was kept low minimizing the two common SPR obstacles; mass transport limitations; and crowding (Myszka & Morton, 1998). Slow dissociation, however, as also previously found in SPR analysis—glucan HoPS—milk protein and a lacquer polysaccharide–polylysine binding (Bai & Yoshida, 2013; Diemer et al., 2012) made C1 chip regeneration necessary for all HePSs, and the effectiveness of this was confirmed by superb reproducibility for binding (in duplicate) of 1 and 6 µg ml$^{-1}$ HePS-3 (Fig. 3A). Importantly, at high concentrations of injected HePS, binding to the C1 reference cell was arising causing a decrease in the RU$_{eq}$ upon steady-state binding analysis (Supplementary Fig. S3). The acceptable HePS concentration entailing minuscule binding to the reference cell was HePS specific and referred to as the critical HePS concentration. Below the critical HePS concentrations ranges, the assay reliably featured typical sensorgrams allowing determination of maximum binding capacity (RU$_{max}$) and $K_{D,app}$ (Fig. 3A–B), which represents a major advancement over the previous established SPR procedure (Babol et al., 2011; Diemer et al., 2012). The steady-state binding curve analysis (Fig. 3B) indicated apparent affinity
for protein and HePS – most likely undergoing multiple binding, hence the slow dissociation (Myszka, 2000). Avidity effects are inherent to the polysaccharide structure and have physiological relevance in biofilms (Sheppard & Howell, 2016) as well as importance for industrial associative polysaccharide–milk protein interactions involved in texturizing microstructures of dairy products (Folkenberg et al., 2006).

For statistical evaluation of the SPR procedure, assays with nBLG immobilized at three different levels were carried out (Table 2). Although the immobilization level increased by ~300% from 288 to 866 RU, the apparent affinity was only little affected yielding an average $K_{A,\text{app}}$ of $0.208\pm0.022$ nM.

**Table 2.** Binding of HePS-3 to native $\beta$-lactoglobulin at three different immobilization levels (RU) on a C1 chip.

<table>
<thead>
<tr>
<th>Immobilization level (RU)</th>
<th>$K_{A,\text{app}}$ (nM)</th>
<th>Average $K_{A,\text{app}}$ (nM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>288</td>
<td>0.217</td>
<td></td>
</tr>
<tr>
<td>311</td>
<td>0.230</td>
<td>$0.208\pm0.022$</td>
</tr>
<tr>
<td>866</td>
<td>0.178</td>
<td></td>
</tr>
</tbody>
</table>

Additionally, two negative control polysaccharides, showing only minuscule binding to the reference-cell and no binding to the sample-cell with immobilized BLG, support the protein binding by HePS (Supplementary Fig. S4).
Figure 3 SPR binding curves between HePS-3 and immobilized native β-lactoglobulin (nBLG) on a C1 chip (RU = 288). A) SPR sensorgrams of HePS-3 injected for 180 s followed by 120 s dissociation. HePS was injected with the following concentrations (µg ml⁻¹): 0, blue; 1 in duplicate, red and bordeaux; 3, yellow; 6 in duplicate, purple and dark blue; 12, green; 30, light blue. B) Steady-state analysis determination of RU_{max} and K_{Dapp} using fitting to a 1:1 binding model.

3.2.2. HePS-3 interaction with native β-lactoglobulin

A pH increase from 4.0 to 4.5 caused K_{A,app} and RU_{max} for HePS-3 and nBLG to decrease 1.3- and 2.9-fold, respectively (Table 3), while no binding was observed at pH 5.0 near the reported nBLG pI of 4.7–5.3 (Mercadante et al., 2012; Sakurai, Konuma, Yagi, & Goto, 2009; Verheul, 1998). Reduced net charge of nBLG or change in ionization state of one or more critical side chains may suppress binding at pH 5.0. At pH 4.0 and 100 mM NaCl, K_{A,app} and RU_{max} decreased 2.3- and 1.4-fold compared to 70 mM NaCl (Table 3); no interaction was detected at 200 mM NaCl.
Table 3. Binding of HePS-3 to native β-lactoglobulin on C1 chip.

<table>
<thead>
<tr>
<th>pH</th>
<th>NaCl (mM)</th>
<th>Temperature (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.0</td>
<td>4.5</td>
<td>5.0</td>
</tr>
<tr>
<td>(K_{A,app} (\mu\text{M}^{-1}))</td>
<td>262.7 ± 10.5</td>
<td>208.6 ± 12.4</td>
</tr>
<tr>
<td>(\text{RU}_{\text{max}})</td>
<td>18.6 ± 0.7</td>
<td>6.5 ± 0.6</td>
</tr>
</tbody>
</table>

n.d. = no detectable signal.

The effects of pH and ionic strength suggest that polar interactions are important for complexation, while the modest increase in \(K_{A,app}\) with increasing temperature (25–35°C) (Table 3) indicated minimal role of hydrophobic interactions.

DLS analysis (Fig. 4) supported the pH dependence of HePS-3 binding. \(d_H\) of nBLG was constant at 6.0±0.4 nm in 70 mM NaCl at pH 3–8, thus close to 7 nm reported for the BLG dimer (Baldini et al., 1999). HePS-3 had \(d_H\) – in the range 100.9±2.7–106.0±3.1 nm at pH 3–6 (Fig. 4, insert). In the absence of salt, BLG and HePS-3 gave \(d_H\) values of 8.2 ± 0.2 nm and 118.3 ± 0.7 nm, respectively, thus no remarkable size difference appeared as a consequence of the presence of NaCl for the two free molecules (data not shown). HePS-3–nBLG mixture had a stable particle mean \(d_H = 37.6±2.9\) nm at pH >5 increasing to a maximum mean \(d_H\) at pH 4 of 184.6±1.3 nm. In absence of NaCl, \(d_H\) of HePS-3–nBLG increased to 448.0±4.8 nm, thus electrostatic shielding weakened HePS-3–nBLG interactions, in accordance with pH and ionic strength effects shown also by SPR. Similarly, DLS data collected at 173° for the HePS-3–BLG mixture in 10 mM sodium citrate pH 4.0 gave \(d_H\) values of 246 ± 11 nm and 50 ± 3 nm in the absence and presence of salt, respectively.
3.2.3. HePS binding capacity

Noise level and activity of the C1 protein surface were assessed as stable during the entire assay (Supplementary results S2.2). Every HePS bound well to nBLG, BCN, KCN, and hBLG (Fig. 5; Supplementary Table S2), with few exceptions. Thus no interaction was observed between nBLG and any of the HePSs at pH 5.0, and for HePS-2 also at pH 4.5 (Fig 5A). No interaction was detected between HePS-6 with any of the four proteins at pH 5.0. Moreover, protein surface saturation was not achieved below the HePS critical concentration level for nBLG–HePS-2 (pH 4.0), BCN–HePS-7 (pH 4.5), KCN–HePS-2 (pH 5.0), hBLG–HePS-5 (4.5), and hBLG–HePS-7 (pH 4.0) (Fig. 5A).

Generally, $R_{\text{Umax}}$ decreased when pH increased from 4.0 to 5.0 (Fig. 5A; Supplementary Table S2), resembling SPR data of α-glucan HoPSs binding milk proteins (Babol et al., 2011; Diemer et al., 2012).
RU<sub>max</sub> decrease from pH 4.0 to 5.0 may stem from reduced protein net charge. Remarkably, average RU<sub>max</sub> at pH 4.0 of all HePSs to BCN (406 RU) was roughly 40% of RU<sub>max</sub> obtained with KCN (406 RU), while nBLG (512 RU) only reached around 15% of average RU<sub>max</sub> with KCN. The same ranking for the proteins were found with α-glucan HoPS where RU<sub>max</sub> at pH 4.0 was about 50% with BCN and 15% with nBLG of the RU<sub>max</sub> value for KCN (Babol et al., 2011). Notably, RU<sub>max</sub> was >10-fold higher with hBLG (1023 RU) than nBLG (512 RU). This, together with binding occurring at pH 5.0 to hBLG but not to nBLG, underscores that heat-treated BLG has increased HePS binding capacity. This may be explained from previous structural analysis establishing that heat treatment of BLG (>80°C) triggers an incipient unfolding of BLG leading to increased exposure of inner hydrophobic amino acid side chains (Moro, Báez, Busti, Ballerini, & Delorenzi, 2011). Finally, to preclude that lack of detected BLG–HePS binding at pH 5.0 was not due to low immobilization level of nBLG (512 RU), a C1 surface with nBLG immobilized at equivalent level (1022 RU) as the hBLG surface (1023 RU) was established and proven unable to bind any of the analyzed HePSs at pH 5.0 (data not shown).
Figure 5 Binding parameters ($RU_{\text{max}}$, $K_{D,\text{app}}$, and $K_{D,\text{app,repeat}}$) for HePS-1–HePS-7 with native β-lactoglobulin (nBLG, magenta), β-casein (BCN, cyan), κ-casein (KCN, white), and heat-treated β-lactoglobulin (hBLG, yellow) as a function of pH in the range 4.0–5.0. A) $RU_{\text{max}}$, B) molar affinities by moles of repeating units, $K_{A,\text{app,repeat}}$ (µM⁻¹), C) molar affinities by moles of HePS, $K_{A,\text{app}}$ (nM⁻¹). $K_{A,\text{app}}$ below 2.5 nM⁻¹ are illustrated in
HePS-2, HePS-6, and HePS-7 gave low (HePS-6 the lowest) \( R_{\text{U,max}} \) values with all four proteins and no binding at pH 5.0. Low \( R_{\text{U,max}} \) values may be explained by the relatively small size of these HePSs (Table 1). HePS-3, however, had similar molar mass as HePS-2 and HePS-7, but gave very high \( R_{\text{U,max}} \), indicating HePS binding capacity and, most probably, binding affinity are influenced by other properties than the molar mass (see 3.2.4).

3.2.4. Affinity of milk proteins for HePSs

The molar \( K_{A,\text{app}} \) for all four investigated protein surfaces varied >3-fold in the pH range 4.0–5.0 for HePS-2, HePS-6, and HePS-7 (except for HePS-7–BCN and HePS-2–KCN), but from negligibly to 3-fold for HePS-1 and HePS-3–HePS-5 (Fig. 5C-D; Supplementary Table S2). By contrast, the \( K_{A,\text{app}} \) for HePS-7–BCN and HePS-2–KCN was only modestly affected (~1.4- and ~2.1-fold) in the pH range 4.0–5.0. The largest affinity reduction was observed for HePS-2–BCN (>20-fold) from pH 4.0 to 5.0 and HePS-6–KCN (>15-fold) from pH 4.0 to 4.5. With HePS-7, \( K_{A,\text{app}} \) for nBLG and KCN decreased >4-fold at pH 4.0 to 5.0 and >9-fold for hBLG at pH 4.5 to 5.0. Notably, \( K_{A,\text{app}} \) for all four milk proteins, except for the HePS-2–nBLG complex, was reduced as pH increased to pH 5.0. By contrast, although changes in \( K_{A,\text{app}} \) for HePS-1 and HePS-3–HePS-5 were modest (<3-fold), their affinity had a tendency to increase with increasing pH in the range 4.0–5.0, except for HeSP-3–nBLG, HePS-4–nBLG and HePS-5–hBLG displaying a small affinity decrease of ~1.1-, ~1.6- and ~1.6-fold, respectively.
Highest apparent affinity of all four protein surfaces was observed with nBLG and BCN binding HePS-1, HePS-4, HePS-5, and to some degree with HePS-3 (Fig. 5C-D; Supplementary Table S2). A large molar mass of HePS-1, HePS-4, and HePS-5 (3.3–4.9 MDa; Table 1) compared to HePS-2, HePS-3, HePS-6, and HePS-7 (0.14–0.39 MDa; Table 1) and high $K_{A,\text{app}}$ probably reinforce multiple binding and avidity effects (Fig. 5C-D; Supplementary Table S2). Still, HePS-2, HePS-3, and HePS-7, despite similar molar mass of around 0.3 MDa, bound milk proteins with different affinity and pH dependency, indicating that other properties of HePSs than molar mass contribute to the protein affinity.

Molar affinity ($K_{A,\text{app},\text{repeat}}$) based on repeat unit size (648.57–1141.04 Da; Table 1) ignores the large differences in HePS molar size (Fig. 5B; Supplementary Table S2). The relative $K_{A,\text{app},\text{repeat}}$ for milk proteins was generally highest for HePS-3 and HePS-4 and lowest for HePS-6 and HePS-7 (Fig. 5B; Supplementary Table S2). HePS-3 and HePS-4 differ from the other HePSs by containing backbone GlcNAc and GalNAc residues, respectively (Fig. 1), which may impose higher binding energy than regular sugar residues, as further supported by comparing $K_{A,\text{app},\text{repeat}}$ for HePS-4 and HePS-7, having similar repeat structures both containing 3:1 1→3:1→6 (branch) linkages combined with compositions of 1:2:1 Glc:Gal:GalNAc and 2:2 Glc:Gal, respectively (Fig. 1). Highest apparent affinity for the four protein surfaces was seen with HePS-4 and the lowest with HePS-7, suggesting GalNAc has a positive effect on the interaction. However, avidity effects due to large differences in molar mass probably also influence the difference in $K_{A,\text{app},\text{repeat}}$ between HePS-4 and HePS-7. Notably, HePS-3 and HePS-7 had similar molar mass but differed significantly in $K_{A,\text{app},\text{repeat}}$, being higher for HePS-3 with all proteins (Fig. 5B; Supplementary Table S2). Thus, different repeat unit structures were demonstrated to result in different $K_{A,\text{app},\text{repeat}}$. $K_{A,\text{app},\text{repeat}}$ of HePS-6 was generally low, except for KCN and hBLG at pH 4.0 (Fig. 5B; Supplementary Table S2). The rhamnose-rich HePS-6 repeat (4:2:1:1 Rha:Glc:Gal:pyruvate)
may have low affinity, as this 6-deoxyhexose can form one hydrogen bond less than regular hexoses. A potential negative charge from the pyruvate in HePS-6 (Fig. 1) may also affect binding.

HePS-2 had medium affinity for the four protein surfaces, except KCN at pH 5.0 and nBLG at pH 4.0, where saturation was not achieved, indicative of low affinity (Fig. 5B; Supplementary Table S2). Furthermore, no interaction was detected with nBLG at pH 4.5–5.0. The HePS-2 repeat unit of seven sugar residues (5:1:1 Gal:Glc:Rha) includes three branches assumed to cause substantial chain stiffness that may lead to lower binding compared to less branched HePSs (Fig. 1). HePS-1 and HePS-5 had very similar molar mass, linkage type, monosaccharide configuration, and number of branches and also gave similar $K_{A,app,repeat}$ with all four proteins (Fig. 5B; Supplementary Table S2).

As seen with regard to binding capacity, affinity of hBLG and nBLG differed remarkably with all HePSs (Fig. 5A-B; Supplementary Table S2). HePS-1 and HePS-3 showed 5–7-fold higher $K_{A,app}$ for nBLG than for hBLG, while HePS-4 and HePS-5 showed 2–5-fold higher $K_{A,app}$ for nBLG (Fig. 5C-D; Supplementary Table S2). By contrast, HePS-6 and HePS-7 had 7–9-fold higher $K_{A,app}$ for hBLG as compared to nBLG. The difference in binding properties probably results from the conformational change of BLG induced by heat treatment (Moro et al., 2011). Furthermore, increase of $K_{A,app,repeat}$ was observed with increasing pH as before for all four protein surfaces for HePS-1 and HePS-3–HePS-5, except for HePS-3–nBLG and HePS-4–nBLG and HePS-5–hBLG, while binding capacity of the proteins decreased in the same pH range (Fig. 5A-B; Supplementary Table S2). Thus, a simple correlation was not observed between binding capacity and apparent affinity within the pH range 4.0–5.0, which emphasized the importance of ability to determine both $R_{U,\text{max}}$ and $K_{A,app}$ of HePS–protein interactions by this SPR procedure.
Overall, the outcome of the present study presumably reflects that structural determinants of repeat units elicit variation in HePS functionality. Usually, α-linkages are more flexible than β-linkages (Laws et al., 2001), furanose confers higher flexibility than pyranose residues (Seo et al., 2008), while branches reduce flexibility (Tuinier et al., 2001). The diversity in HePS linkage type, monosaccharide ring configuration, and degree of branching thus invokes considerable span of HePS flexibility. This in turn has direct implication for manufacture of fermented dairy products where the milk is severely heat treated (i.e. at 90–95°C for several minutes) prior to fermentation in order to ensure better texture as a consequence of disulfide interactions between BLG and KCN on the surface of the casein micelles as well as in the serum phase (Donato & Guyomarc’h, 2009). Subsequent production of HePS by the starter culture used during fermentation will then further influence texture through potential interaction (aggregative or segregative) with the milk proteins (van de Velde, de Hoog, Oosterveld, & Tromp, 2015). These authors have also noted that molecular weight, linkage type, and charge of HePS are the main factors determining their effect on structure. From the present results, it would appear that a viable strategy to control texture in fermented dairy products could be to characterize starter cultures in terms of the parameters inducing flexibility and the charge density of the HePS produced in order to maximize binding affinity. This, however, has to be complemented with textural and microstructural characterization on actual dairy products and verified with a larger set of samples.

4. Conclusions

The established SPR protocol provides a tool for screening and validating HePS–milk protein interactions by steady-state analysis of the maximum binding capacity $RU_{max}$ and the apparent association constants $K_{A,app}$ and $K_{A,app,repeat}$. The procedure is sensitive, uses a small amount of HePS for
full binding analysis, and is applicable in evaluating both food and non-food protein–polysaccharide interactions. To our knowledge, this is the first study where comparative HePS–milk protein binding affinities have been reported. The effects of pH, ionic strength, and temperature changes suggest HePS–protein complex formation to be driven mainly by polar interactions. Heat treatment (85°C/15 min) of BLG dramatically increased its HePS binding capacity but reduced the affinity. Even though differences in binding properties were not assigned to specific chemical properties of HePSs, the data support that both molar mass and oligosaccharide repeat structure are important for complexation as measured by maximum binding capacity $R_{U_{\text{max}}}$ and apparent association constants $K_{A,\text{app}}$ and $K_{A,\text{app,repeat}}$.

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