Biophysical analysis of the interaction of the serum protein human β2GPI with bacterial lipopolysaccharide

Anna Gries a,∗, Ruth Prassl b, Satoshi Fukuoka c, Manfred Rössle d, Yani Kaconis e, Lena Heinbockel e, Thomas Gutsmann e, Klaus Brandenburg e

a Institute of Physiology, Medical University of Graz, Harrachgasse 21/V, 8010 Graz, Austria
b Institute of Biophysics, Medical University of Graz, Schmiedlstr. 6, 8042 Graz, Austria
c European Molecular Biology Laboratory, c/o DESY, D-22603 Hamburg, Germany
d Institute of Physiology, Medical University of Graz, Harrachgasse 21/V, 8010 Graz, Austria
e Forschungszentrum Borstel, Leibniz-Zentrum für Medizin und Biowissenschaften, Parkallee 10, D-23845 Borstel, Germany

A R T I C L E  I N F O

Article history:
Received 7 February 2014
Revised 23 April 2014
Accepted 23 April 2014

Keywords:
Human glycoprotein β2GPI
Lipopolysaccharide
Cytokine production
Immune modulation
LAL test

A B S T R A C T

There are several human serum proteins for which no clear role is yet known. Among these is the abundant serum protein beta2-glycoprotein-I (β2GPI), which is known to bind to negatively charged phospholipids as well as to bacterial lipopolysaccharides (LPS), and was therefore proposed to play a role in the immune response. To understand the details of these interactions, a biophysical analysis of the binding of β2GPI to LPS and phosphatidylserine (PS) was performed. The data indicate only a moderate tendency of the protein (1) to influence the LPS-induced cytokine production in vitro, (2) to react exothermally with LPS in a non-saturable way, and (3) to change its local microenvironment upon LPS association. Additionally, we found that the protein binds more strongly to phosphatidylserine (PS) than to LPS. Furthermore, β2GPI converts the LPS bilayer aggregates into a stronger multilamellar form, and reduces the fluidity of the hydrocarbon moiety of LPS due to a rigidification of the acyl chains. From these data it can be concluded that β2GPI plays a role as an immune-modulating agent, but there is much less evidence for a role in immune defense against bacterial toxins such as LPS.

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1. Introduction

Serum proteins such as albumin or low- and high-density lipoproteins (LDL and HDL) fulfill a variety of physiological tasks like transport, adhesion, recognition and defenses against invading microorganisms. To this class also belong lipopolysaccharide-binding protein (LBP) and lactoferrin which are considerably up-regulated in acute phase serum i.e. in the case of bacterial infections [1,2]. In contrast the function of the abundant serum protein β2GPI is not completely understood. Various publications presented data on its role as binding molecule to acidic lipids. Actually the analysis of the crystal structure of β2GPI [3,4] gave evidence for domain V being an excellent counterpart for negatively charged phospholipids. These data were extended by Hammel et al. [5] who found a direct interaction of acidic phospholipids such as cardiolipin with the hydrophobic loop adjacent to the positively charged lysine-rich region of domain V of β2GPI.

There is some evidence that β2GPI may also play a role as regulator of the immune response to bacterial toxins such as lipopolysaccharide (LPS, endotoxin). Laplante and co-workers published data giving evidence that β2GPI interacts specifically with LPS and that both LPS and Toll-like receptor 4 (TLR4) are required for β2GPI to activate macrophages [6]. This idea was even more expressed in the investigations of Agar et al. [7] who proposed that β2GPI might be a scavenger of LPS deduced from the inhibition of the LPS-induced expression of tissue factor and interleukin-6 (IL-6) from monocyes.

From these findings the role of β2GPI still does not seem to be clear. Therefore, we have applied various biophysical techniques to study the interaction of β2GPI with LPS in more detail. This is

Abbreviations: β2GPI, beta2-glycoprotein-I; FRET, fluorescence resonance energy transfer spectroscopy; FTIR, Fourier-transform infrared spectroscopy; HDL, high-density lipoproteins; ITC, isothermal titration calorimetry; LAL, Limulus amebocyte lysate; LBP, lipopolysaccharide-binding protein; LDL, low-density lipoproteins; LPS, lipopolysaccharides; MNC, mononuclear cells; PC, phosphatidylcholine; PS, phosphatidylserine; SAXS, small-angle X-ray scattering

∗ Corresponding author. Tel.: +43 3163804402.
E-mail address: anna.gries@medunigraz.at (A. Gries).
of special importance since LPS belongs to the strongest elicitors of the immune system known in nature. It is responsible, for example, for the induction of many cytokines and chemokines but also for the production of prostaglandins in human mononuclear cells [8]. These induction mechanisms may be beneficial at low amounts of LPS but lead to pathophysiological reactions at higher LPS concentrations such as the induction of sepsis. Although β2GPI binds to LPS we did not get evidence for a significant role of β2GPI as defense protein. We propose rather an activity-modulating role possibly in the sense of an immune modulation which may be important to combat LPS-induced inflammation by enhancing the innate immunity.

2. Materials and methods

2.1. Lipopolysaccharide and β2GPI

Lipopolysaccharides from *Salmonella enterica* Minnesota rough mutants Re and Ra, strains R595 and R60 were extracted from the bacteria grown at 37 °C by the phenol:chloroform:petrol ether (PCP) method, purified and used in the natural salt form [9]. The purity was examined by MALDI-TOF mass spectrometry and LPS samples were only used when the chemical structure in particular of the lipid A part consisted of a diglucosamine to which six acyl chains in amide- and ester-linkage at positions 2 and 2′, and 3 and 3′, respectively, were bound and which were phosphorylated at positions 1 and 4′, according to the known structure of lipid A. LPS Re and Ra were either dissolved in 50 mM Tris–HCl, 25 mM NaCl, 0.1 g/L EDTA, pH 7.4 (Tris–HCl) or in 20 mM Hepes buffer at concentrations of 1 mg/ml. The samples were sonicated for 20 min at 60 °C followed by 30 min cooling to 4 °C and afterwards reheated to 60 °C for 30 min and finally stored at 4 °C for at least 12 h before use.

2.2. Phospholipids

Egg yolk phosphatidylcholine (Egg-PC) or phosphatidylserine (PS) purchased from Avanti Polar Lipids (Alabaster, USA) were dissolved in chloroform/methanol 2:1 (v/v). The organic solvents were evaporated at 40 °C under a stream of nitrogen. The film was dried in high vacuum overnight to remove residual traces of solvent. The dry lipid film (1 mg lipid) was suspended in 1 ml Tris–HCl and hydrated at 50 °C for 1 h interrupted by vigorous mixing every 10 min. Lipid suspensions were extruded through a polycarbonate filter with 100 nm pore size using a LiposoFast pneumatic extruder (Avestin, Inc. Ottawa, ON).

2.3. Isolation of beta2-glycoprotein-I

β2GPI was purified from human plasma by treatment with 1.4% (v/v) perchloric acid followed by affinity chromatography on heparin–sepharose and cation exchange chromatography on Mono S (Pharmacia, Sweden) [10]. The preparation was homogenous as judged by SDS/PAGE (10% resolving and 3.75% stacking gel) yielding a single band.

2.4. Intrinsic Trp-fluorescence

Trp-fluorescence was recorded on a Spex Fluoromax-3 fluorescence spectrometer (Jobin Yvon Horiba, Longjumeau, France) using a 10 mm × 10 mm quartz cuvette. Trp-residues were excited at 292 nm and emission spectra were recorded from 320 to 380 nm with an increment of 1 nm. Band widths of 5 nm were used for excitation and emission with an integration time of 0.1 s, averaging 10 scans. The emission scans were processed for inner filter and instrumental corrections and the background intensities of the samples without protein were subtracted. Titration experiments were performed with continuous stirring by adding aliquots of LPS (0.5 mg/ml suspended in Tris–HCl) into the cuvette containing β2GPI (50 μg/ml, 1.1 μM) dissolved in the same buffer. The ratio of LPS/β2GPI was incrementally increased from 0 to 40 mol/mol by titration. Control experiments were performed in the same way by adding aliquots of Egg-PC unilamellar vesicles.

The temperature of the cuvette was maintained at 25 °C and the single measurements were taken after 10 min of equilibration.

2.5. Acrylamide quenching of Trp fluorescence

Aliquots of a 3 M acrylamide stock solution in Tris–HCl were added to a 1.1 μM solution of β2GPI containing 20 mol LPS/mol protein. The acrylamide concentration was varied in a range between 0.02 and 0.45 M. The experimental conditions were as described above. The values obtained were corrected for volume increase and scattering derived from acrylamide titration of LPS.

The fluorescence quenching data were analyzed with a Stern–Volmer plot. In the case where both static and dynamic quenching occur simultaneously in the sample the following modified form of the Stern–Volmer equation holds

\[ I_0 / I = (1 + K_{SV} Q) \exp(-Q) \]

where \( I_0 \) is the fluorescence intensity at zero quencher concentration, \( I \) is the maximum intensity at a given quencher concentration \( Q \), and \( K_{SV} \) is the Stern–Volmer quenching constant. The term \( \exp(V|Q|) \) is used as a phenomenological descriptor of the quenching process where \( V \) represents an active volume of a sphere around the fluorophore to such an extent that any quencher within this volume is able to quench the excited fluorophore at the time of fluorescence excitation.

2.6. Stimulation of human mononuclear cells by LPS

The stimulation of human mononuclear cells (MNC) was performed as described previously [11]. Briefly MNC were isolated from heparinized blood of healthy donors. The cells were resuspended in medium (RPMI 1640) at 5 × 10^6 cells/ml. For stimulation 200 μl MNC (1 × 10^6 cells) were transferred into each well of a 96-well culture plate. LPS Ra and β2GPI mixtures were preincubated for 30 min at 37 °C and added to the cultures at 20 μl per well. The cultures were incubated for 4 h at 37 °C under 5% CO₂. After centrifugation of the culture plates for 10 min at 400g supernatants were collected and stored at −20 °C. TNFα concentrations were quantified by a sandwich ELISA using a monoclonal antibody against TNF (clone 6b from Intex AG, Switzerland).

2.7. X-ray scattering

The aggregate structures of LPS Ra were determined in the absence and presence of β2GPI using small-angle X-ray scattering (SAXS). SAXS measurements were performed at the European Molecular Biology Laboratory (EMBL) outstation at the Hamburg synchrotron radiation facility HASYLAB using the double-focusing monochromator-mirror camera X33. Scattering patterns in the range of the scattering vector 0.01 < s < 0.1 nm⁻¹ (s = 2 sin(θ)/λ, 2θ the scattering angle and λ the wavelength = 0.15 nm) were recorded at 20, 40, 60, and 80 °C with exposure times of 1 min using an image plate detector with online readout (MAR345, MarResearch, Norderstedt, Germany) [12]. The s-axis was calibrated with Ag-Behenate with a periodicity of 5.84 nm. The scattering patterns were evaluated as described previously [13] assigning the spacing ratios of the main scattering maxima to defined three-dimensional...
2.8. Fourier-transform infrared spectroscopy (FTIR)

FTIR was used to determine (a) the gel to liquid crystalline phase transition of the hydrocarbon chains of LPS by evaluating the peak position of the symmetric stretching vibration \( v_s(\text{CH}_2) \) in the wavenumber range \( 2850–2853 \text{ cm}^{-1} \) and (b) the secondary structure of the protein in the range of the amide I vibration in the wavenumber range \( 1700–1600 \text{ cm}^{-1} \). The infrared spectroscopic measurements were performed on an IFS-55 spectrometer (Bruker). LPS and LPS:β2GPI samples dispersed in 20 mM Hepes buffer, pH 7.0 were placed in a CaF\(_2\) cuvette between two 12.5 μm teflon spacers. For the determination of the phase behavior temperature-scans were performed automatically between 10 and 70 °C with a heating-rate of 0.6 °C/min. Every 3 °C, 5 interferograms were accumulated, apodized, Fourier transformed and converted to absorbance spectra. For the determination of the secondary structures an attenuated total reflection unit was used whereupon the protein and protein:LPS mixtures were spread under evaporation of the excess water in a stream of nitrogen and measuring the samples at the temperature of the instrument (30 °C).

2.9. Isothermal titration calorimetry (ITC)

Microcalorimetric measurements of the binding of β2GPI to LPS Ra were performed on an MCS isothermal titration calorimeter (Microcal Inc.) at 37 °C as described previously [13]. LPS Ra (0.05 mM) was dispensed into the microcalorimetric cell (volume 1.3 ml) and the protein solution (1 mM) was filled into the syringe compartment (volume 100 μl). After temperature equilibration β2GPI was titrated every 5 min in 3 μl portions into the LPS-containing cell under constant stirring and the heat of reaction was plotted versus time.

2.10. Fluorescence resonance energy transfer spectroscopy (FRET)

The ability of β2GPI to bind to and intercalate into phospholipid liposomes or into LPS R60 aggregates was investigated by FRET as described earlier [13]. Briefly, phospholipid liposomes from phosphatidylserine (PS), phosphatidylcholine (PC) or LPS R60 were doubly labeled with the fluorescent phospholipid dyes N-(7-nitrobenz-2-oxa-1,3-diazol-4-yl)-phosphatidyl ethanolamine (NBD-PE) and N-(lissamine rhodamine B sulfonyl)-phosphatidylethanolamine (Rh-PE) (Molecular Probes). Intercalation of unlabeled molecules into the doubly labeled liposomes leads to probe dilution and thereby to a lower FRET efficiency. Thereby the emission intensity of the donor \( I_D \) increases and that of the acceptor \( I_A \) decreases (for clarity, only the quotient of the donor and acceptor emission intensity is shown here).

In all experiments 100 μl of 100 μM β2GPI were added to doubly labeled PC, PS or LPS R60 (900 μl of 10 μM) at 50 s after equilibration. NBD-PE was excited at 470 nm and the donor and acceptor fluorescence intensities were monitored at 531 and 593 nm respectively and the fluorescence signal \( I_D/I_A \) was recorded for further 250 s.

2.11. Determination of endotoxin activity by the chromogenic Limulus test

To test a possible contamination of β2GPI with endotoxins the LPS content was determined by a quantitative kinetic assay based on the reactivity of Gram-negative endotoxin with Limulus amebocyte lysate (LAL) at 37 °C, using test kits of LAL Coomatic Chromo-LAL K (Chromogenix, Haemochrom). The standard endotoxin used in this test was from Escherichia coli (O55:B5) and 10–15 endotoxin units (EU)/ml correspond to 1 ng/ml. In this assay saturation occurs at 50 EU/ml and the resolution limit is 0.05 EU/ml (maximum value for ultrapure water, Aqua B. Braun).

3. Results

3.1. Purity of β2GPI

Despite the extraction and purification procedure leading to a single band in SDS-PAGE as described above contamination by LPS cannot be excluded. Therefore endotoxin was determined in the Limulus amebocyte lysate (LAL) assay and the results are shown in Table 1. Clearly the protein is not free of LPS. From the data of Table 1 an endotoxin concentration of ~2 EU/μg can be estimated which corresponds to approximately 150–250 pg LPS per μg protein.

3.2. Stimulation of mononuclear cells

Serum proteins such as β2GPI are important LPS binding proteins. Therefore the ability of β2GPI to influence the inflammation induction by LPS was investigated by stimulating mononuclear cells (MNC) with LPS at different concentrations (100 and 10 ng/ml) in the absence and presence of varying concentrations of β2GPI [LPS]:[β2GPI] 1:1, 1:10, 1:100, and 1:1000 weight:weight). The addition of the protein to LPS at a ratio of 1:1 and 1:10 respectively influenced the LPS-induced cytokine (TNFα) secretion only marginally (data not shown). At ratios of 1:100 and 1:1000 however a clear suppression of the cytokine production was observed.

Table 1

| Dilution | LAL result (diluted) [EU/ml] | LAL result (undiluted) [EU/ml] | App. LPS-content [EU/μg] |
|----------|-----------------------------|-------------------------------|--------------------------|
| 800      | 1                           | 1863.3                        | 2.3                      |
| 80       | 10                          | 152.99                        | 1.9                      |
| 8        | 100                         | 14.641                        | 1.8                      |

![Fig. 1](image.png)

Fig. 1. Secretion of tumor-necrosis-factorα from human mononuclear cells induced by lipopolysaccharide LPS Ra, and LPS:β2GPI mixtures. The error bar results from the determination of TNFαs in an ELISA in duplicate.
change of the cytokine secretion was observed (Fig. 1). TNFα concentrations in the range 2000–1750 pg/ml for pure LPS are significantly reduced at [LPS]:[β2GPI] 1:100. Interestingly there is no further decrease at the highest concentration. In contrary, at [LPS]:[β2GPI] 1:1000 a highly significant increase of TNFα secretion can be observed. The bar at the right hand side in Fig. 1 shows that the protein itself seems to be an activator of MNC. This activity however could be explained at least partially by the presence of LPS in as much as 10 μg/ml β2GPI were contaminated by 2 ng/ml LPS.

3.3. Interaction of β2GPI with LPS-aggregates assessed by Intrinsic Trp-fluorescence

Interaction of β2GPI with LPS was monitored by measuring alterations in the fluorescence spectra of tryptophan (Trp), which occur due to changes in the microenvironment of Trp-residues. As shown in our previous paper on the binding behavior of β2GPI to negatively charged cardiolipin vesicles [5], the Trp-emission spectrum of β2GPI is rather broad with a maximum (λmax) at about 350 nm. This feature arises by reason of different microenvironments of the five Trp-residues in β2GPI. Four Trp-residues located in domain 1–4 are embedded in the interior hardly accessible to water whereas Trp316 located in domain V is surface exposed [3]. Despite its apolar nature Trp316 is positioned in a hydrophobic loop which is surface exposed and was found to be essential for the lipid-binding capacity of β2GPI, as a single point mutation of this amino acid completely abrogates lipid binding [16,17]. Thus one can assume that upon lipid association Trp316 becomes located in a more hydrophobic lipid environment seen as a blue shift in the fluorescence emission spectra. Indeed a significant blue shift
from 350 nm to 336 nm was observed when cardiolipin vesicles were added to a solution of β₂GPI. The magnitude of the blue shift was dependent on the molar ratio of lipid-to-protein being constant above a molar ratio of 20 mol lipid per mol protein [5].

Here we have performed an analogous experiment applying increasing LPS concentrations between 0 and 45 μM corresponding to molar ratios up to 40 mol LPS/mol β₂GPI. Depending on the concentration of LPS a blue shift from 350 nm to 339 nm was observed which in contrast to cardiolipin increases linearly up to 5 μM LPS and remains unchanged above a molar ratio of about 10 (Fig. 2A and B). As β₂GPI lacks the ability to bind to zwitterionic choline headgroups [5,18], the fluorescence emission maximum observed for egg-PC at 350 nm remained unchanged (Fig. 2B).

Fig. 3 shows the Stern–Vollmer plots for the quenching of the intrinsic Trp fluorescence of β₂GPI by acrylamide. We observed a non-linear biphasic characteristic of the quenching curves with an upward curvature indicative for simultaneous static and dynamic quenching. The effective quenching constants were determined from the curvature indicative for simultaneous static and dynamic quenching with an upward biphasic characteristic of the quenching curves with an upward curvature indicative for simultaneous static and dynamic quenching. The effective quenching constants were determined from the curvature indicative for simultaneous static and dynamic quenching. The effective quenching constants were determined from the curvature indicative for simultaneous static and dynamic quenching.

3.4. Gel to liquid crystalline phase transition of the acyl chains of LPS

The gel (β) to liquid crystalline (α) phase transition of LPS ranges from 30 to 35 °C for enterobacterial LPS. Binding to serum proteins may play a role in modulating immune responses to LPS. In Fig. 4, in FTIR experiments as sensitive indicator of the phases the peak position of the symmetric stretching vibration at 2850–2853 cm⁻¹ is plotted for pure LPS as well as in the presence of β₂GPI at a ratio of [LPS]:[β₂GPI] 1:1 weight%. For LPS alone the wavenumber values changed from 2850.5 to 2852.5–2853.0 cm⁻¹ indicating a midpoint of the transition at 32.4 °C which considerably increased to a value of 36.3 °C in the presence of [LPS]:[β₂GPI] 1:1 weight%. At the same time, at all temperatures the wavenumber values are lower for the sample with protein. Both effects shift in Tₘ to higher values as well as decrease in wavenumbers are characteristic for a rigidification of the entire LPS aggregate.

3.5. Secondary structure of β₂GPI

The secondary structures of proteins such as α-helical and β-sheets and their changes due to interaction with LPS may be of relevance for an understanding of the mode of action. FTIR allows an analysis of the secondary structures by evaluating the amide I vibration in the spectral range of 1700–1600 cm⁻¹. The main secondary structural elements α-helix, β-sheets, and random coils have maxima at 1650–1657, 1625–1640, and 1645–1660 cm⁻¹, respectively. A particular helical substructure, 3₁₀ helices, has bands around 1637–1643 cm⁻¹.

In Fig. 5, data are presented for pure β₂GPI and in the presence of LPS ([LPS]:[β₂GPI] 10:1). The pure protein exhibits a main vibrational maximum at 1640.7 cm⁻¹, which is shifted to a value of 1648.4 cm⁻¹ in the presence of LPS. The position of the former band may indicate a mixture of a β-sheet and a 3₁₀ helical structure. In the presence of LPS the significant shift to higher wavenumbers is indicative for a transformation, at least partially, into a random coil organization. Since the assignment to the different secondary structures may be difficult due to the superposition of the single band components we have also applied resolution-enhancing techniques such as Fourier self-deconvolution (FSD) and second derivatives of the spectra, but these analyzes did not provide any extra support for an increase in random structure.

3.6. Three-dimensional aggregate structure of LPS

The supramolecular aggregate structure of LPS has been shown to represent an important determinant for its ability to be biologically active i.e. by inducing the cytokine production in human MNC. Changes of the aggregate structure due to protein binding may strongly influence the capability of LPS to induce cytokines.

The determination of the aggregate structure can be performed by SAXS which shows characteristic scattering patterns depending on the kind of aggregates. For pure LPS the scattering patterns exhibit broad intensity distributions between s values of 0.15–0.35 nm⁻¹ indicating the existence of LPS bilayer systems (form factor), which do not show a particular aggregation behavior (no structure factor), in accordance to numerous data published earlier [19].

In the presence of the protein ([LPS]:[β₂GPI] 1:4) the scattering patterns in the range of 20–80 °C (Fig. 6) exhibit sharp reflections

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**Fig. 3.** Quenching of the intrinsic Trp fluorescence of β₂GPI by acrylamide. The protein concentration was set to 1.1 μM. The acrylamide concentration was increased to 0.45 M. The Stern–Volmer plot is shown for β₂GPI in the absence (▲) and in the presence of 20 mol LPS/mol protein (●).
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Fig. 4. Gel to liquid crystalline phase transition of the acyl chains of LPS Ra in the absence and presence of β2GPI at [LPS]:[β2GPI] 10:1 weight%. Shown is the peak position of the symmetric stretching vibration of the methylene groups νs(CH2) versus temperature. In the gel phase, the position lies around 2850.5, in the liquid crystalline phase at 2852.5–2853 cm−1.

Data presented in Fig. 7 indicate an exclusively exothermic process, which however does not show a sigmoidal saturation profile as normally observed in this type of measurement. A sigmoidal response is not necessarily to be expected in the case of specific interactions, in particular for binding proteins such as lactoferrin. It was not possible over the range of [protein]:[LPS] ratios used here to evaluate binding constants. Furthermore, the observed small enthalpy changes in the range −8 to −2 kJ/mol indicate only a weak binding of the protein to LPS.

3.8. Intercalation of β2GPI into phospholipid or LPS

FRET was applied to study a possible intercalation of β2GPI into phospholipid liposomes made from PC or PS or into LPS aggregates. As shown in Fig. 8A–C there is only a very weak interaction of β2GPI with PC (Fig. 8A) leading to an insignificant change of the FRET signal of 0.02. For the interaction of β2GPI with the negatively charged phospholipid PS there is a much higher change of the FRET signal of nearly 0.2 (Fig. 8B). Despite the highly negative head group charge of LPS the interaction with β2GPI leads to an increase of the FRET of only 0.05 (Fig. 8C).

4. Discussion

We have performed a biophysical analysis of the abundant serum protein β2GPI with bacterial lipopolysaccharides and in some cases also with phospholipids to understand the still insufficiently defined role of β2GPI. We have found that β2GPI provokes an increase in the phase transition temperature of LPS (Fig. 4). Additionally a change of the aggregate structure of LPS with a moderate tendency to aggregate into a multilamellar phase was observed (Fig. 6). At the same time the interaction indicates a clearly exothermic process (Fig. 7) which, however, shows no saturation characteristics. This is in accordance with the rather weak interaction in the FRET experiment (Fig. 8C) being lower than that with the negatively charged PS (Fig. 8A), despite the fact that the surface charge density of PS is lower than that of LPS. Concomitantly to these physical data the biological experiment exhibits a moderate decrease of the LPS-induced stimulation of human mononuclear cells (Fig. 1) in a concentration range ([LPS]:[β2GPI] 1:1–1:100) in which the endotoxin contaminant within the β2GPI does not yet play a role.

The experimental data of the secondary structures of the protein (Fig. 5) are indicative of a slight transformation of the β-sheet and a 3_10 helical structure of the protein into – as one possible interpretation – a random coil conformation. Literature data of secondary structures of β2GPI have been published by Paolorossi and Montich [20] and Borchman et al. [21]. The former group found – among others – a temperature-dependent distribution of β-sheets (bands at 1631–1636 cm−1) and α-helical structures with the former structures dominant at 25 °C and the latter at 50 °C. In the presence of negatively charged phosphatidylglycerols (POPG, DMPG, DPPG), the main components at 1631–1633 cm−1 remained corresponding to β-sheets but may be influenced by the addition of NaCl leading partially to more α-helical structures. The data of Borchman et al. [21] show significant variations already for different preparations of the protein with 27 and 39% α-helix and β-sheets for one and 36 and 53% respectively for the other. Also the turns vary considerably while no random coil structures were observed. In the presence of phosphatidylcholine the structural units corresponding to turns nearly disappeared whereas the addition of the negatively charged cardiolipin led to a significant increase of random coil structures. This phenomenon was also observed in this work after addition of LPS. Summarizing the different data and interpretations in the literature and observed in our experiments

3.7. Thermodynamics of binding of β2GPI to lipopolysaccharide

Isothermal titration calorimetry (ITC) is an important technique to study the thermodynamics of molecule interactions. In this way characteristics of binding (endotherm, exotherm), the binding stoichiometry and saturation can be analyzed.

with one main reflection d1 and the second order at d1/2, at 80 °C also with the third order reflection at d1/3. This behavior can be interpreted as occurrence of a multilamellar LPS aggregate due to protein binding with a lamellar periodicity of 9.12–8.77 nm generated i.e. by distances between neighboring bilayer planes. However from the sharpness of the single reflections it can be deduced that the multilamellarization is only weakly expressed.
do not give a satisfactory picture for an understanding of the complex behavior of β2GPI alone and in the presence of different lipids. Therefore further detailed analyzes of the interaction of the protein with the most important lipid systems deserve closer attention.

Agar and co-workers published data from which they deduced that β2GPI belongs to the class of defensive proteins by scavenging LPS [7]. These data result among others from the inhibition of the LPS-induced production of tissue factor and interferon-6 in monocytes. In contrast to this hypothesis our data do not favor an important role of β2GPI as defense structure. Biophysical data of other defense structures such as polymyxin B [22], high-density lipoprotein [23], lactoferrin [24] and various antimicrobial peptides [25] indicate a much higher change of LPS membrane fluidity leading in most cases to complete fluidization, considerably higher multilamellarization of the LPS aggregates and a much higher exothermic reaction in the ITC experiments with starting values of 80 kJ/mol and with saturation characteristics. Furthermore the decrease of the LPS-induced immunostimulatory activity elicited by the protein in the range [LPS]:[β2GPI] 1:1–1:100 weight% shown in Fig. 1 is much lower than for the defense proteins and peptides described above [18–21].

The weak and non-saturable binding of β2GPI to LPS – resembling a catalysis-like process, but not necessarily being catalytic – could be a signal of an immune modulating function of the protein. This view is supported by FRET data (Fig. 8B) and is in accordance with the strong interaction of β2GPI with phosphatidylserine (PS) as reported earlier [18]. In this work a direct interaction of the positively charged lysine-rich region of β2GPI with the PS head group was suggested. As also presented in our work PS is an important phospholipid compound in the context of apoptotic processes. Thereby PS is exported from the inner monolayer of immune cells to its outer side thereby being directly accessible to the action of β2GPI. In this sense we would agree with the statement of Agar et al. [7] that β2GPI could play a role as component of innate immunity. Regarding their data of cytokine inhibition however, which is only moderate even at rather high excess of β2GPI with respect to LPS we would not support the view that β2GPI has a significant role for LPS neutralization and clearance. The interpretation of an immune modulating property of β2GPI seems also to be in accordance with data of Ninivaggi et al. [26] who found that β2GPI incubated with phospholipids inhibited the generation of thrombin.

Laplante et al. [6] also investigated the interaction of β2GPI with LPS. They reported that β2GPI interacts specifically with LPS and that this interaction is responsible for apparent TLR4 activation.
Fig. 8. Förster resonance energy transfer spectroscopy (FRET) of mixtures from doubly labeled 0.01 mM phosphatidycholine (A), phosphatidylserine (B) and LPS Ra (C) with biGPI (10 μM) added after 50 s. The FRET signal $I_d/I_a$ is a sensitive measure of incorporation of the protein into the lipids.
by β2-GPI. We do not agree that (i) the binding of β2-GPI to LPS is specific (see above) and (ii) there is a significant interaction of β2-GPI with TLR4. Laplante et al. were unable to prove the latter interaction because they found no immune stimulatory action of β2-GPI in cases where the contaminating LPS was removed by purification or by polymyxin B. Thus, their statement that ‘both LPS and β2-GPI are required for the direct interaction of β2-GPI to bind to and activate macrophages’ remains not fully supported.

Further experiments seem to be necessary to elucidate the interaction of β2-GPI with LPS and other negatively charged lipids from different membranes. Particularly atomic force microscopy should be applied to give a view on the immune cell’s membrane to explain the direct interaction of β2-GPI and β2-GPI:LPS mixtures with the membrane and its components, in particular with phosphatidylserine.

Acknowledgements

We kindly acknowledge the help of Nina Hahlbrock and Christine Hamann for performing the IR spectroscopic and cytokine as well as the FRET experiments. The technical assistance of Josef Kellner is also gratefully acknowledged.

We are indebted to the German Ministry (Ministerium für Bildung und Forschung) BMBF, project 01GU0824 and the Else Kröner-Fresenius-Stiftung, project 2011_A140 for financial help.

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