The J-elongated conformation of $\beta_2$-glycoprotein I predominates in solution: implications for our understanding of antiphospholipid syndrome

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$\beta_2$-Glycoprotein I (G2PI) is an abundant plasma protein displaying phospholipid-binding properties. Because it binds phospholipids, it is a target of antiphospholipid antibodies (aPLs) in antiphospholipid syndrome (APS), a life-threatening autoimmune thrombotic disease. Indeed, aPLs prefer membrane-bound G2PI to that in solution. G2PI exists in two almost equally populated redox states: oxidized, in which all the disulfide bonds are formed, and reduced, in which one or more disulfide bonds are broken. Furthermore, G2PI can adopt multiple conformations (i.e. J-elongated, S-twisted, and O-circular). While strong evidence indicates that the J-form is the structure bound to aPLs, which conformation exists and predominates in solution remains controversial, and so is the conformational pathway leading to the bound state. Here, we report that human recombinant G2PI purified under native conditions is oxidized. Moreover, under physiological pH and salt concentrations, this oxidized form adopts a J-elongated, flexible conformation, not circular or twisted, in which the N-terminal domain I (DI) and the C-terminal domain V (DV) are exposed to the solvent. Consistent with this model, binding kinetics and mutagenesis experiments revealed that in solution the J-form interacts with negatively charged liposomes and with MBB2, a monoclonal anti-DI antibody that recapitulates most of the features of pathogenic aPLs. We conclude that the preferential binding of aPLs to phospholipid-bound G2PI arises from the ability of its preexisting J-form to accumulate on the membrane, thereby offering an ideal environment for aPL binding. We propose that targeting the J-form of G2PI provides a strategy to block pathogenic aPLs in APS.

G2PI is a 50-kDa multidomain glycoprotein that circulates in the plasma at a concentration of ∼0.2 mg/ml (1, 2) (Fig. 1A). It consists of 326 amino acids organized into five domains (DI–V) connected by four short linkers, Lnk1 (residues 61–64), Lnk2 (residues 119–122), Lnk3 (residues 182–185), and Lnk4 (residues 242–244) (3). Domains I–IV are canonical complement control protein (CCP) domains, each containing 4 cysteine residues typically forming 2 disulfide bonds (4, 5). CCP domains are found in regulators of complement activation, such as factor H (FH) (6), complement receptor 1 (CR1), membrane-cofactor protein (MCP), and decay-accelerating factor (DAF) (5). In contrast, DV is aberrant, consisting of one extra pair of cysteines and a 19-residue hydrophobic loop that is responsible for anchoring the protein to negatively charged phospholipids (7, 8) (Fig. 1A). Importantly, G2PI exists in two interconvertible biochemical variants, oxidized (54%) and reduced (46%), depending on the integrity of the disulfide bonds (9, 10). In the oxidized form, 11 disulfide bonds are formed. In the reduced form, the disulfide bonds C288–C326 in DV and C32–C60 in DI are individually or simultaneously broken.

Forgotten for many years since its discovery in 1961 (1), G2PI gained popularity in the fields of hematology and rheumatology in 1990, when it was recognized by two independent studies as the dominant antigen of antiphospholipid antibodies (aPLs) in antiphospholipid syndrome (APS) (11–13), a life-threatening blood-clotting disorder characterized by vascular thrombosis and pregnancy morbidity (10, 14). In this context, G2PI is heavily studied. Indeed, autoantibodies against G2PI (anti-G2PI) are frequently found in young patients with a history of thrombosis (15, 16); they are often associated with lupus anticoagulant, a laboratory test that indicates predisposition for blood clots (17); they induce (18) and potentiate thrombus formation in vivo (14, 18–20) and cause pregnancy complications resulting in fetal loss (21). Thus, a deeper understanding of the structural determinants of antigen-antibody recognition is likely to accelerate the development of new diagnostics and therapeutics for APS patients.

An important feature of all aPLs, and especially highly pathogenic aPLs recognizing the epitope R39–R43 in the N-terminal domain I of G2PI, referred to here as anti-DI antibodies (22–25), is that their detection requires proper immobilization of the antigen onto negatively charged surfaces or lipid
membranes (26–28), raising the question of whether the epitopes recognized by aPLs are cryptic in the circulating form of β2GPI. In support of this viewpoint, structural studies have documented that β2GPI can adopt alternative O-circular (29–31), S-twisted (32), and J-elongated conformations (29–31, 33, 34) featuring different exposures of DI and DV to the solvent (Fig. 1B) and have led to the proposition of a model whereby the O-circular form of β2GPI, which was captured by negative-stain EM (29) and atomic force microscopy (AFM) (31) using β2GPI purified from plasma using mild conditions, is the most abundant (>90%) protein conformation of β2GPI under physiological conditions, which is immunologically inert and incapable of reacting with aPLs (Fig. 1B, left). In contrast, the J-elongated form, originally described in 1999 by X-ray crystallography using β2GPI purified from plasma using the harsh oxidizing agent perchloric acid (33, 34) and, more recently, observed by EM (29) and AFM (31) using “native” β2GPI subjected to high salt (i.e. 1.15 M NaCl) and high pH (11.5) or in complex with the mAb 3B7 (29), bacterial lipopolysaccharide (LPS) labeled with gold nanoparticles (30), and protein H of Streptococcus pyogenes (35), is believed to be the immunogenic conformation of the protein that interacts with aPLs, which appears when β2GPI binds to the membranes (Fig. 1B, right). Since the S-twisted conformation of the protein, inferred by small-angle X-ray scattering (SAXS) (32), was not detected by EM, AFM, or X-ray crystallography, this model also predicts that the S-twisted form represents a transient, unreactive intermediate state that the protein populates while transitioning between the J and O forms (Fig. 1B, central).

Although very popular in the field and highly endorsed by the scientific community, there are two important caveats for this commonly accepted model. First, the three structures have been obtained using different protein preparations and experimental techniques. Second, the O-circular form has never been documented in solution. Thus, while consensus exists regarding the fact the J-form is the structure of β2GPI bound to aPLs (29) and perhaps to the membranes (30), unresolved issues remain: 1) which conformation exists and predominates in solution, and 2) what is the conformational pathway leading to the bound state.

Encouraged by our recent results with prothrombin (36–39), the second most common antigen of aPLs in APS, this work was initiated to study the solution structure of β2GPI and provide new insights into the mechanism of autoantibody recognition. Our results, based on X-ray crystallography, single-molecule FRET (smFRET), SAXS, binding kinetics, and mutational studies, were unexpected. In contrast to what was previously thought and widely accepted in the field (2, 29) (Fig. 1B), we found that, in the free form, human recombinant oxidized β2GPI adopts the J-elongated conformation, not O-circular or S-twisted, in which DI is exposed to the solvent and, therefore, is available for autoantibody recognition. Based on this new evidence and previous findings (28, 40, 41), an alternative mechanism to explain how negatively charged phospholipids enhance the affinity toward aPLs without requiring opening of the protein structure or relocation of the glycosylations away from DI is proposed, and its implications for our understanding of APS is discussed here.

Results

Expression, purification, and functional characterization of human recombinant beta-2 glycoprotein I

To get a better grasp of the structural architecture of β2GPI under conditions relevant to physiology, we set out to perform structural and biophysical studies of fully glycosylated human recombinant β2GPI (hrβ2GPI). Two versions of the proteins were successfully expressed and purified under native conditions at high yield and purity. The first version, called LT-β2GPI, contained a long multifunctional cleavable tag at the N terminus, located right before the natural N-terminal sequence (Fig. 2A and Fig. S1). The tag was then cleaved with enterokinase to generate the intact, mature protein (hrβ2GPI). Removal of the tag was confirmed by N-terminal sequencing (Fig. 2B). The second version, called ST-β2GPI, contains a shorter, noncleavable purification tag at the N terminus that, based on our previous work (37), is expected not to affect the conformational properties of the protein (Fig. 2B and Fig. S1). ST-β2GPI was made to eliminate the enterokinase cleavage step, which was very laborious and not as efficient as expected. The presence of the short tag was confirmed by N-terminal
sequencing and accounted for the different electrophoretic mobilities observed between recombinant and plasma purified protein before and after enzymatic removal of the N-glycosylations (Fig. 2B).

To evaluate the functional integrity of the recombinant proteins, LT-β2GPI, hrβ2GPI, and ST-β2GPI were tested in several biochemical assays. β2GPI purified from human plasma using the perchloric acid method (pβ2GPI) was used as a control. Using surface plasmon resonance (SPR), a technique that allows us to measure association (on) and dissociation (off) rate constants in real time, we found that all variants interacted avidly with liposomes containing negatively charged phospholipids, such as phosphatidylserine, yet they failed to interact with phospholipids made entirely of phosphatidylcholine (Fig. 2C and D). Importantly, the values of the affinity constants were similar for all the constructs and consistent with published data (28), and so was the inhibitory effect of physiological concentrations of calcium chloride (data not shown). These results document structural integrity of the hydrophobic loop in DV and also prove that the phospholipid binding activity of β2GPI is not perturbed by the presence or removal of the purification tags.

In addition to properly interacting with phospholipids, the recombinant proteins were also successfully recognized in ELISAs by aPLs isolated from four triple-positive APS patients, which contain anti-DI antibodies (25, 36, 42) (Fig. 2E). In this case, however, LT-β2GPI exhibited significantly lower values of optical density at 450 nm than the other variants and plasma-purified protein, suggesting that the presence of the long tag masks some epitopes or, more likely, changes the preferential orientation of the molecule that is adsorbed onto the plastic surface. Taken together, these studies validate recombinantly made β2GPI as a proxy for plasma-purified β2GPI. They also demonstrate that, under physiological conditions, β2GPI is primed for phospholipid binding.

X-ray crystal structure of human recombinant beta-2 glycoprotein I

In 1999, two identical X-ray crystal structures of β2GPI were simultaneously solved by two independent groups (33, 34). β2GPI used in these studies was purified from human plasma using the perchloric acid method and crystallized at resolutions of 2.87 Å (1C1Z) and 2.7 Å (1QUB). Since then, no other crystal structure has been deposited in the Protein Data Bank (PDB).

To investigate the structural properties of the recombinant proteins, crystallization experiments were performed for all the protein constructs. While it was not possible to crystallize LT-β2GPI, we solved the X-ray crystal structures of hrβ2GPI (Fig. 3A) and ST-β2GPI (Fig. 3B) at 2.6- and 3.0-Å resolution, respectively. We also solved the X-ray crystal pβ2GPI at 2.4 Å as a control (Fig. 3C). Similar to what was previously reported by Bouma et al. (34) and Schwarzenbacher et al. (33),...
diffraction quality crystals were obtained after 2 weeks at 4 °C using ammonium sulfate as a precipitating agent. The crystals belong to the orthorhombic space group C2221 (Table 1). Notably, ST-β2GPI, for which extra electron density was observed at the N terminus (Fig. 3D), crystallized under similar conditions and in the same space group as hrβ2GPI and pβ2GPI, confirming minimal structural perturbation introduced by the artificial tag.

Consistent with previous structural data (33, 34), all three independently solved X-ray crystal structures depicted β2GPI featuring an elongated conformation spanning ~140 Å in length, from the N to the C terminus. The first three domains, DI–DIII, are aligned along the vertical axis of the molecule, whereas DIV and DV bend, forcing the molecule to adopt a characteristic J-shaped elongated form resembling a hockey stick. DI and DV are located >100 Å apart, and both of them are exposed to the solvent. Interestingly, the side chain of residue R43, which is part of the epitope recognized by anti-DI antibodies, is partly exposed to the solvent as it forms a hydrogen bond network with residues R39, G41, and T57 (Fig. 3E).

β2GPI is known to circulate in alternative chemically different species, oxidized and reduced (9, 10, 43). Oxidized β2GPI contains 11 disulfide bonds, whereas reduced β2GPI contains 8 disulfide bonds and four free thiols resulting from the rupture of the disulfide bonds C32-C60 in DI and C288-C326 in DV. In our crystal structures, all 22 cysteine residues were engaged in 11 disulfide bridges (Fig. 3, A–C), indicating that the structure of human recombinant β2GPI purified by immunoaffinity and size exclusion chromatography represents the oxidized form of β2GPI.

Given the similar experimental conditions in which the crystals grew, it was not surprising to observe that, overall, the three structures were superimposable (RMSD of 0.709, 0.516, and 0.630 Å for 6V06 versus 6V08, 6V06 versus 6V09, and 6V08 versus 6V09, respectively) and also very similar to the published ones (RMSD of ≤0.810 Å) (33, 34) (Fig. 4A). There were, however, a few notable differences. First, the phospholipid binding loop in DV (residues 308–319) adopted alternative conformations (RMSD of 3.122 Å), documenting flexibility and exposure to the solvent (Fig. 4A). This is in agreement with spectroscopic data indicating that W316 is exposed to the solvent (44) and that plasmin cleaves β2GPI at position K317-T318 (45). Second, significant extra electron density appeared after molecular replacement, especially in the two datasets at a higher resolution (i.e., 2.4 and 2.6 Å) (Fig. 4B), which we attributed to the N-linked glycosylations (Fig. 4, C and D). Guided by MS analyses (46), we modeled the following sugar sequences: Gal2GlcNAc2Man3GlcNAc2 at N143, Gal2GlcNAc2Man3GlcNAc2 at N164, GlcNAc2 at N174, and Gal3GlcNAc2 at N234. The presence of a putative O-linked glycosylation at T130 could not be confirmed because of weak density, even in the datasets at higher resolution.

smFRET studies of human recombinant beta-2 glycoprotein I

The high salt concentration used in the crystallization buffers may destabilize hydrogen bonds and favor hydrophobic
interactions, forcing the protein to assume a nonnative conformation. To test whether the J-elongated form exists in solution and determine its relative abundance compared with other forms, we applied smFRET to β2GPI (38, 39, 47) (Fig. 5A). By recording the energy that is transferred from an excited fluorophore (donor) to a second fluorophore with spectral overlap (acceptor), smFRET measures distances on a nanometer scale, allowing us to interrogate protein structures and protein

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**Table 1**

Crystallographic data for the structures of human beta-2 glycoprotein I

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value(s) for:</th>
<th>Value(s) for:</th>
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<tbody>
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<td>Buffer/salt</td>
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<tr>
<td>PDB entry</td>
<td>6V06</td>
<td>6V08</td>
<td>6V09</td>
</tr>
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**Data collection**

| Wavelength (Å)            | 1.54          | 1.033         | 1.54          |
| Space group               | C222          | C222₁         | C222          |
| Unit cell dimensions (Å)  | a = 160.8, b = 166.9, c = 114.0 | a = 159.3, b = 173.2, c = 115.2 | a = 160.2, b = 171.2, c = 113.4 |
| Resolution range (Å)      | 40–2.4        | 40–2.6        | 40–3.0        |
| No. of observations       | 513,296       | 363,697       | 158,386       |
| No. of unique observations| 59,497        | 48,503        | 31,543        |
| R_sym (%)                 | 7.9 (50.6)    | 14.3 (64.7)   | 9.5 (46.1)    |
| I/σ(I)                    | 21.1 (2.4)    | 10.0 (1.5)    | 13.7 (2.4)    |

**Refinement**

| Resolution (Å)            | 40–2.4        | 40–2.6        | 40–3.0        |
| R cryst, R free           | 0.201, 0.236  | 0.200, 0.232  | 0.223, 0.246  |
| No. of protein atoms      | 2540          | 2510          | 2517          |
| No. of solvent molecules  | 431           | 377           | 15            |
| RMSD* bond lengths (Å)    | 0.013         | 0.010         | 0.011         |
| RMSD angles (°)           | 2.0           | 2.0           | 1.7           |
| RMSD ΔB (Å²) (mm/ms/ssb)  | 5.12/5.28/6.93| 4.54/5.13/5.85| 3.97/3.58/4.38|
| Protein (Å²)              | 63.8          | 68.5          | 71.0          |
| Solvent (Å²)              | 64.4          | 62.8          | 49.6          |

**Ramachandran plot:**

| Most favored (%)          | 98.9          | 100.0         | 99.6          |
| Generously allowed (%)    | 1.1           | 0.0           | 0.4           |
| Disallowed (%)            | 0.0           | 0.0           | 0.0           |

| a RMSD from ideal bond lengths and angles and RMSD in B-factors of bonded atoms. |
| b mm, main chain–main chain; ms, main chain–side chain; ss, side chain–side chain. |

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**Figure 4. Location and structural role of the N-linked glycosylations.** A, superposition of five X-ray crystal structures of β2GPI highlights structural similarities yet diversity of the phospholipid binding loop in DV (residues 308–319). B, extra electron density detected in the structure of pβ2GPI solved at 2.4-Å resolution attributed to the N-linked glycosylations. The domains of β2GPI are color coded as shown in Fig. 1. The presence of a putative O-linked glycosylation at T130 could not be confirmed because of weak density. Side (C) and top (D) views of the N-glycosylations surrounding DIII are shown. The electron density 2F₀-Fc map is countered at 0.8σ.
conformational changes one molecule at a time. This overcomes averaging effects seen in traditional bulk experiments.

Guided by our structures, we generated four FRET pairs in the ST background, S13C/S112C, S13C/S312C, S112C/S312C, and S190C/S312C (Fig. 5, B and C), by replacing the natural serine residues with isosteric cysteines and then reacting those newly engineered cysteines with Alexa Fluor 555 (AF555)-maleimide as the FRET donor and Alexa Fluor 647 (AF647)-maleimide as the FRET acceptor. Residue 13 is located in DI, residue 112 is located in DII, residue 190 is located in DIV, and residue 312 is located in DVI. Labeling occurred only at the engineered sites, as no fluorescence was observed for WT β2GPI (Fig. 5C).

This result is consistent with our structural data and provides additional support to our conclusion that human recombinant β2GPI secreted in the media and purified for our structural studies does not contain free thiols. Given the Förster radius, R₀ = 50 Å, of the 555/647 FRET couple, the crystal structure predicts no FRET for the mutants S112C/S312C and S13C/S312C. In contrast, high FRET and low but measurable FRET values are expected for the FRET pair S13C/S112C and S190C/S312C, respectively. This is because residues 13 and 112 are located ~24 Å apart while the Ca-Ca distance between residues 190 and 312 is ~46 Å. Remarkably, the experimental results were fully consistent with our structure-based calculations, validating our previous structural work and proving that the elongated conformation predominates (>90%) under physiological pH and salt concentrations (Fig. 5E).

Specifically, probes located at positions C13, C312, C112, and C312 reported a negligible FRET signal, whereas probes attached to the S13C/S112C and S190C/S312C mutants reported high (E_FRET = 0.92) and low (E_FRET = 0.26) FRET values, respectively. Interestingly, the construct 190/312 displayed a FRET distribution wider than the theoretical distribution predicted by shot noise (48), documenting the existence of multiple conformations at equilibrium mediated by the flexibility of Lnk4.

We also tested the effect of sodium chloride (0.025 versus 0.145 versus 1 m, Fig. 5F) and pH (4.0 versus 7.4 versus 10.0, Fig. 5G). In contrast to previous findings in which low pH and low concentrations of sodium chloride favored circularization of the protein (29), no significant FRET differences were observed, suggesting that variation of [NaCl] and pH produces conformational changes, such as further elongation of the protein structure or rotations of the domains, that could not be detected by our FRET pairs.

**SAXS studies of human recombinant beta-2 glycoprotein I**

To rule out potential artifacts arising from the replacement of natural amino acids with cysteine and incorporation of fluorescent dyes, we collected SAXS data for ST-β2GPI and pβ2GPI under physiological pH and salt concentrations (Fig. 6). SAXS is a biophysical method that is particularly useful to assess the overall shape of biological macromolecules in solution (49), i.e. linear versus globular, and, similar to smFRET, therefore is ideal to detect large conformational changes in β2GPI. The radius of gyration (R_g) for ST-β2GPI and β2GPI purified from plasma was 44.19 ± 2.22 and 44.39 ± 2.10, respectively (Fig. 6A). These values and the maximum linear dimension (D_max) obtained by computing the pair distance distribution function, P(r), shown in Fig. 6B (i.e. ~160 Å), agree very well with the values calculated using the J-elongated structure captured by X-ray crystallography (i.e. R_g = 45.22 and D_max ~140 Å, Fig. 3A) and explain the lack of FRET for the FRET couples 13/312 and 112/312.
How anti-DI antibodies recognize β2GPI

Figure 6. Elongated conformation of β2GPI revealed by SAXS. Scattering profiles (A) and pair distribution functions (B) for pβ2GPI (black) and ST-β2GPI (magenta) collected at 2 mg/ml under physiological conditions (20 mM Tris, pH 7.4, 145 mM NaCl). The calculated values of the radius of gyration, Rg, are very similar for pβ2GPI and ST-β2GPI. The blue curve in panel B, which is significantly different from the experimental scattering profiles, represents the theoretical pair distribution function for a hypothetical circular conformation. C, ab initio envelope calculated from scattering profiles for ST-β2GPI (magenta). Note that the linear arrangement of domains in the structure 6V09 is consistent with the elongated SAXS envelope.

(Fig. 5E), confirming that β2GPI is extended in solution, not circular. Whether the S- or J-form predominates in solution cannot be concluded from these studies due to the low resolution of SAXS. However, it is worth noting that the molecular envelope calculated ab initio from the SAXS data of ST-β2GPI returned a linear and not a twisted envelope, as documented before (32), in which the structure of ST-β2GPI fits nicely (Fig. 6C) without having to rotate or bend the domains.

Autoantibody binding studies

In addition to demonstrating that oxidized β2GPI adopts an elongated conformation in solution, our structural analysis predicts that this form may be primed for autoantibody binding, especially anti-DI antibodies. To test this hypothesis, we took advantage of the reactivity of MBB2 (21), a newly developed recombinant mAb raised against DI that, upon complement fixation, recapitulates, in vivo, most of the clinical characteristics assigned to pathogenic aPLs. The binding of β2GPI to MBB2 was monitored using SPR (Fig. 7).

To retain the native conformation of β2GPI in solution, we immobilized β2GPI to the chip’s surface and injected β2GPI in the fluid phase. Binding between MBB2 and β2GPI should occur only if DI is exposed to the solvent. This experimental setup is different from previously reported interaction data between MBB2 and β2GPI in which β2GPI was covalently immobilized on a dextran-based chip and the antibody was used in the fluid phase to mimic binding of MBB2 to β2GPI bound to negatively charged phospholipids (21). The results of the experiments shown in Fig. 7A demonstrate that MBB2 interacts with β2GPI in solution with a modest but measurable affinity, characterized by a dissociation constant, Kd, of 2.2 ± 0.2 μM (Fig. 7B). The interaction was characterized by very fast on and off rates, suggesting that electrostatic interactions dominate the binding interface. This was demonstrated by systematic experiments in which we varied the concentration of sodium chloride in the running buffer from 300 to 15 mM. As expected, the affinity constant (Kd) of MBB2 for β2GPI increased ~700-fold at low salt concentrations, from 4.4 × 10⁴ M⁻¹ at 300 mM NaCl to 3.2 × 10⁷ M⁻¹ at 15 mM NaCl (Fig. 7C).

Interestingly, the Kd value determined for MBB2 at 145 mM NaCl is similar to the Kd value obtained by Dienava-Verdoodt et al. for patient-derived monoclonal antibodies targeting DI (50). Furthermore, the affinity of MBB2 for soluble β2GPI is 200-fold weaker than the affinity previously determined for MBB2 toward immobilized β2GPI (i.e. Kd of 11 nM) (21), as expected for pathogenic aPLs. To test whether MBB2 binds to the epitope R39-R43 in DI that is targeted by pathogenic aPLs in APS patients (22), positively charged residues R39, R43, and K44 were mutated to the neutral amino acid alanine. Remarkably, a significant ~30-fold reduction of the affinity under physiological conditions was detected for the triple mutant (Fig. 7, D and E). This result is consistent with the electrostatic nature of the interaction and demonstrates that these three positively charged amino acids are part of the binding epitope.

The ability of MBB2 to interact with residues R39-K44 in DI provides the opportunity to further and independently test the model in Fig. 1B envisioning the epitope R39-R43 to be cryptic, either buried by DV in the O-circular form (29) or shielded by the N-linked glycosylations in the S-twisted form (32). To this goal, we measured the affinity of three new constructs, isolated DI (residues 1–60), β2GPI deletion Lnk2 (β2GPI Δ120–122), and recombinantly deglycosylated β2GPI (degβ2GPI) (ID305/ N143Q/N164Q/N174Q/N234Q) toward immobilized MBB2, which were designed to force the exposure of DI to the solvent and, in principle, should have higher affinity for MBB2. Our binding data shown in Fig. 7F indicate that all three constructs interact with immobilized MBB2 with micromolar affinity comparable to that of full-length β2GPI WT. This conclusively rules out a significant contribution of the neighboring domains and the glycosylations in shielding the R39-R43 epitope when the protein is free in solution and further validates the conclusions derived from our structural studies.

Discussion

In recent years, our structural understanding of multidomain clotting and complement factors, such as prothrombin (37, 39), plasminogen (51), FXII (52), ADAMTS-13 (53), factor H (6), and properdin (54), has evolved, and so have the technologies capable of capturing protein conformational changes in solution. Owing to its flexibility, the solution structure of β2GPI has remained controversial, and so is the mechanism through which pathogenic anti-DI antibodies recognize β2GPI in APS
patients, thereby promoting thrombosis. To fill this gap in knowledge, this study was initiated to provide a rigorous structural and functional assessment of plasma-purified and human recombinant B2GPI under conditions relevant to physiology, using traditional (X-ray crystallography, SAXS, and mutagenesis studies) and state-of-the-art (smFRET) technologies. To our knowledge, this is the first application of smFRET to a CCP-containing protein.

A first major conclusion emerging from this study is that human B2GPI expressed in mammalian cells and purified under native conditions adopts an elongated, flexible conformation in which DV and DI are exposed to the solvent. Importantly, this conformation has 11 disulfide bonds (Fig. 3) and does not react with thiol-specific maleimide dyes (Fig. 5B), thereby representing the oxidized form of B2GPI, which accounts for 54% of the protein in human plasma (9). In the free form, under physiological pH and salt concentrations, we propose that B2GPI is primed for phospholipid binding and autoantibody recognition. As such, these findings contrast with current structural and mechanistic models envisioning B2GPI primarily (~90%) adopting an O-circular conformation in human plasma and formation of the J-elongated form only after binding to the membranes (Fig. 1B).

A second major conclusion of this study is that recombinantly made B2GPI protein is structurally and functionally identical to B2GPI purified from plasma using the perchloric acid method. Hence, in contrast to what was previously thought (29), the elongated conformation of B2GPI is not an artifact caused by the harsh purification methods or crystallization conditions but a genuine conformation of the protein in solution. Whether the treatment of B2GPI with perchloric acid introduces additional chemical modifications to specific amino acids (e.g., oxidation and deamination) remains a possibility that cannot be ruled out by our current studies.

In addition to settling previous controversies in the field, the recognition that the elongated form of oxidized B2GPI...
How anti-DI antibodies recognize β2GPI

preexists and, according to our smFRET experiments (Fig. 5E) and data from Ioannou et al. (9), predominates in human plasma bears important implications in our understanding of the APS pathology. It also provides new ideas for the development of APS-focused diagnostics and therapeutics.

Regarding the mechanism of anti-DI antibody recognition, our structural and binding data indicate that the opening of the protein structure and relocation of DI away from the glycosylation sites are neither necessary nor sufficient to explain how β2GPI becomes a better antigen for anti-DI antibodies upon binding to the membranes. They instead strongly suggest that, in agreement with previous oligomerization models (39, 49, 48, 49), binding of the preexisting elongated conformation of β2GPI to the membranes gives rise to an ideal surface in which β2GPI has a sufficiently high density and adopts a favorable orientation that promotes bivalent binding. Indeed, recent theoretical calculations (55) and binding studies (56) elegantly demonstrated that the energetic gain associated with limiting antigen diffusivity is quite substantial, and such a gain is expected to be even higher for low-affinity antibodies, such as aPLs. In this context, rotation or bending of the CCP domains relative to the plane of the membrane, such as those documented here, by the FRET couple 190/312 may be important for proper packing of β2GPI onto the lipids. A contribution of local conformational changes, such as exposure of R43 upon binding to the lipids, is also possible, yet, considering the intrinsic low affinity of aPLs for their targets, the modest effect caused by mutations in isolated DI (51), and the key role of bivalence documented before (49, 52), the energetic contribution of this process is expected to be minor.

One of the strongest arguments in favor of the O-circular/J-elongated conformational model is that immunocomplexes are rarely found in patients’ plasma (57, 58). If β2GPI were primed for autoantibody binding, then immunocomplexes should be seen more often in APS patients. An alternative view for this phenomenon, which is consistent with the preexistence of the J-elongated form in solution, comes from our SPR binding kinetics and analysis of the literature. Figure 7A shows that, under physiological pH and salt concentrations, the interaction of MB2B and β2GPI is characterized by a low affinity (Kₜ'' of 2.2 μM) and fast on/off rates, indicating that, in solution, the immunocomplexes form rapidly but also dissociate very rapidly. According to previously published data (20), the plasma level of anti-β2GPI antibodies in thrombotic APS patients represents 0.5% or less of the total IgG pool, which corresponds to molar concentrations of ~0.1 μM. Given the micromolar affinity of MB2B for its target, this translates into <90% complex formation at equilibrium. Hence, the immunocomplexes are extremely difficult to detect and even more challenging to purify because of their thermodynamic instability and low concentration of autoantibodies with respect to their targets. In this context, a possible role of the negatively charged surfaces, as suggested by our SPR binding experiments performed at low concentrations of NaCl (Fig. S3), could be to stimulate the binding of anti-DI antibodies to β2GPI by slowing down the dissociation rate, resulting in complex stabilization. Such a mechanism is fully consistent with recent observations (35) and previous data (49) and might be conserved among other aPLs.

An important and relatively unknown aspect of the pathogenic APS is how aPLs potentiate thrombus formation (10, 14, 61–64). The discovery that the J-elongated form dominates in solution provides new insights into this mechanism. Since binding between aPLs and oxidized β2GPI can occur in solution, our studies support the hypothesis that stimulation of circulating and endothelial cells requires clustering of the antigen onto the membranes, which leads to aPL-dependent complement activation (18) and/or aPL-induced receptor dimerization (63–65).

Regarding the development of new diagnostics and therapeutics, if the main role of the membranes is to increase the local concentration of the elongated form, we speculate that immobilization of human recombinant β2GPI produced in this work at the desired density and with a defined orientation should provide a novel, efficient, robust, and cost-effective method to detect anti-β2GPI antibodies. On the other hand, blocking the binding of the elongated form of β2GPI to cell receptors and phospholipids should limit the pathogenic effects of anti-DI antibodies. This approach could complement current strategies aimed at competing with anti-DI antibodies in solution (13, 57, 58, 61), as it would theoretically block other potentially pathogenic aPLs, in addition to those targeting DI. Consistent with this premise, antibodies against DV found in APS patients do not induce thrombosis but are protective instead (66), and a novel dimeric molecule, A1-A1, protects mice from aPL-induced thrombosis by interfering with ApoER2 and phospholipid binding (63).

In conclusion, our study provides clear-cut evidence that the monomeric oxidized form of β2GPI adopts a J-elongated conformation under physiological pH and salt concentrations, not O-circular or S-twisted. Whether the reduced form of β2GPI adopts alternative structures in solution remains a possibility. However, answering this question is challenging and will require rigorous structural, biophysical, and mutagenesis studies.

Experimental procedures

Protein production and purification

β2GPI WT and mutants were expressed in BHK (LT-β2GPI) and HEK293 (ST-β2GPI) mammalian cells and purified to homogeneity by immunopurification and heparin and size exclusion chromatography after swapping the signal peptide of β2GPI with the one of the coagulation factor X to boost expression and adding a furin specific recognition motif RRKR for quantitative posttranslational processing. The purity and chemical identity of each fragment were verified by SDS-PAGE and N-terminal sequencing. Domain I (1–60) was chemically synthesized and refolded as done before (67). Plasma-derived β2GPI (pβ2GPI) was purified using the perchloric acid method, as described previously (67). MB2B was produced as described before (21). Liposomes composed of phosphatidylcholine (PC) or phosphatidylcholine and phosphatidylserine (PS) in a 4:1 molar ratio (PC:PS) were prepared by extrusion using 100-nm polycarbonate membranes (Avanti Polar Lipids, Alabaster, Alabama), kept a 4 °C, and used within 7 days. ELISAs were performed as described before (25, 36, 67). Protein concentrations
were determined by reading at 280 nm with molar extinction coefficients adjusted according to the amino acid sequence. All other chemicals were purchased from Sigma-Aldrich.

**SPR experiments**

Binding affinities for liposomes were measured as done before (36), using an L1 sensor chip in which liposomes were immobilized at 1600 response units (RU). Titrations were performed by injecting increasing concentrations (0–2 μM) of β2GPI and its variants in running buffer (20 mM Tris, pH 7.4, 150 mM NaCl, 0.1%, w/w, BSA) at a flow rate of 25 μl/min at 25 °C. Binding affinities for MBB2 were measured using a CM5 sensor chip in which MBB2 was immobilized at 6000 RU using NHS/EDC chemistry. Titrations were performed by injecting increasing concentrations (0–20 μM) of β2GPI and its variants in running buffer (20 mM Tris, pH 7.4, 25–300 mM NaCl, 0.01%, w/w, Tween 20) at a flow rate of 25 μl/min at 25 °C. All experiments were carried out on a BIAcore-S200 instrument (GE-Healthcare). The dissociation constants \(K_d\) were obtained as a fitting parameter by plotting the value of the RU at the steady state for each concentration using the BIAevaluation software and Origin Pro 2015.

**X-ray studies**

Crystalization of human recombinant (hrβ2GPI and ST-β2GPI) and pb2GPI was achieved at 4 °C by the vapor diffusion method, mixing 3 μl of protein (10 mg/ml) with equal volumes of reservoir solution. Optimization of crystal growth was achieved by the hanging drop vapor diffusion method, mixing 3 μl of protein (10 mg/ml) with equal volumes of reservoir solution at 4 °C. After 7–10 days at 4 °C, crystals were frozen with 25% glycerol from original mother liquid. X-ray diffraction data were collected at 100 K with a home source (Rigaku 1.2-kW MX007 generator with VHF optics) Rigaku Raxis IV++ detector for pb2GPI and ST-β2GPI and with detector Pilatus of beamline ID23 at the Advanced Photon Source, Argonne, IL, for hrβ2GPI. Data sets were indexed, integrated, and scaled with the HKL2000 software package (68). All structures were solved by molecular replacement using PHASER from the CCP4 suite (46) and the structure of β2GPI (PDB entry 1C1Z) as the starting model. Refinement and electron density greration were performed with REFMAC5 from the CCP4 package. 5% of the reflections were randomly selected as a test set for cross-validation for four structures. Model building and analysis of the structures were carried out using COOT (69). Ramachandran plots were calculated using PROCHECK. Statistics for data collection and refinement are summarized in Table 1.

**Protein labeling for single-molecule detection**

Selective labeling of the unpaired Cys residues with AF555–C2-maleimide as the donor and AF647–C2-maleimide as the acceptor was achieved as described recently for prothrombin, with minor modifications (37, 38). Briefly, β2GPI (8–10 μM) was used in the presence of 350 mM NaCl, 20 mM Tris, pH 7.4, at room temperature for 1 h in the dark and in the presence of DTT at a molar ratio \([−\text{SH}]:[\text{DTT}]\) of 1:1.6. Constructs were then labeled with an equal molar mixture of the thiol-reactive dyes, using AF555–C2-maleimide as the donor and AF647–C2-maleimide as the acceptor (Thermo Fisher Scientific, MA). The labeling reaction was carried out for 3 h at room temperature in the dark. The monomeric protein free of unreacted dyes was purified on an analytical Superdex 200 column (GE Healthcare, PA), and the efficiency of derivatization was assessed by UV-Vis measurements.

**Single-molecule FRET measurements**

FRET measurements of freely diffusing single molecules were performed with a MicroTime 200 confocal microscope (PicoQuant, Berlin, Germany) as detailed elsewhere (37, 38). Briefly, as shown in Fig. 5A, experiments were carried out with pulsed interleaved excitation, which reports the status of both donor and acceptor fluorophores by sorting molecules on the basis of relative donor:acceptor stoichiometry and apparent FRET efficiency. The donor and acceptor dyes were excited with a ps-pulsed diode laser at 532 and 638 nm, respectively. To achieve pulsed interleaved excitation (70), the 532-nm laser was electronically delayed 25 ns relative to the 638-nm laser (48, 71). A dual-band dichroic mirror reflecting 532 nm and 638 nm guided the light to a high-numerical-aperture apochromatic objective (60×, numeric aperture of 1.2, water immersion, Olympus) that focused the light to a confocal volume of 1.0 fl for excitation at 532 nm and detection at 575 nm. Fluorescence from excited molecules was collected with the same objective and focused onto a 50-μm-diameter pinhole. The donor and acceptor emissions were separated via a dichroic long-pass filter with a dividing edge at 620 nm. Suited bandpass filters were inserted to eliminate the respective excitation wavelength and minimize spectral cross-talk. The fluorescence was detected with two avalanche photodiodes using time-correlated single-photon counting with the TimeHarp 200 board. Data were stored in the time-tagged, time-resolved mode. Measurements were performed 25 μm deep in the solution with a total acquisition time of 1 h and repeated fresh up to four times on each protein sample (50–100 pm). Signals from single molecules were observed as bursts of fluorescence. Bursts with more than 40 counts were searched with the APBS algorithm, while the integration time was set to 0.5 ms (72). Appropriate correction for direct excitation of the acceptor at the donor excitation wavelength, leakage of the donor in the acceptor channel, and the instrumental \(γ\) factor were calculated using a mixture of dsDNA models with known FRET efficiency (E) and stoichiometry (S) labeled with dyes AF555 and AF647 (73). Only molecules with a stoichiometry in the range of 0.25–0.75 were considered in the final analysis, and their distribution was fit to Gaussian curves using Origin 2015 (OriginLab Corporation, Northampton, MA). Data recording was performed using SymphoTime software 6.4 (PicoQuant, Berlin). Data analysis was carried out with PAM (74).

**Small-angle X-ray scattering measurements**

SAXS data were collected at beamline 12-ID-B of the Advanced Photon Source at Argonne National Laboratory.
How anti-DI antibodies recognize β2GPI

(Argonne, IL) on ST-β2GPI and pβ2GPI at concentrations of 0.5, 1, 2, and 5 mg/ml. The $R_g$ was determined using the Guinier approximation in the low q region ($qR_g < 1.3$), and its linearity served as an initial assessment of data and sample quality. Maximum particle dimension, $D_{\text{max}}$ and distance distribution function, $P(r)$, were calculated using GNOM. The low-resolution envelopes were produced using both GASBOR (75) (q up to 0.8 Å$^{-1}$) and DAMMIN (76) (q up to 0.3 Å$^{-1}$) by directly fitting the reciprocal space-scattering profile. Twenty models were generated for every calculation and then aligned and averaged using DAMAVER (76). The results of GASBOR and DAMMIN were very similar, but only GASBOR results are reported here. Structural figures were prepared using Pymol.

Data availability

Atomic coordinates and structure factors have been deposited in the Protein Data Bank (accession codes 6V06 for pβ2GPI at 2.4 Å, 6V08 for hrβ2GPI at 2.6 Å, and 6V09 for ST-β2GPI at 3.0 Å).


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Abbreviations—The abbreviations used are: β2GPI, β2, Glycoprotein I; aPLs, antiphospholipid antibodies; APS, antiphospholipid syndrome; AFM, atomic force microscopy; SAXS, small-angle X-ray scattering; smFRET, single-molecule FRET; SPR, surface plasmon resonance; RMSD, root mean square deviations; AF555, Alexa Fluor 555; $R_g$ radius of gyration; $D_{\text{max}}$ maximum linear dimension; RU, response units; CCP, complement control protein; PC, phosphatidylcholine; PS, phosphatidylserine.

References


How anti-DI antibodies recognize β2GPI


