The multimeric glycoprotein von Willebrand factor (VWF) mediates platelet adhesion to collagen at sites of vascular injury. The binding site for collagen types I and III is located in the VWF-A3 domain. Recently, we showed that His1023, located near the edge between the “front” and “bottom” faces of A3, is critical for collagen binding (Romijn, R. A., Bouma, B., Wuyser, W., Gros, P., Kroon, J., Sixma, J. J., and Huizinga, E. G. (2001) J. Biol. Chem. 276, 9985–9991). To map the binding site in detail, we introduced 22 point mutations in the front and bottom faces of A3. The mutants were expressed as multimeric VWF, and binding to collagen type III was evaluated in a solid-state binding assay and by surface plasmon resonance. Mutation of residues Asp979, Ser1020, and His1023 nearly abolished binding, whereas mutation of residues Ile975, Thr977, Val997, and Glu1001 reduced binding affinity about 10-fold. Together, these residues define a flat and rather hydrophobic collagen-binding site located at the front face of the A3 domain. The collagen-binding site of VWF-A3 is distinctly different from that of the homologous integrin αI domain, which has a hydrophobic binding site located at the top face of the domain. Based on the surface characteristics of the collagen-binding site of A3, we propose that it interacts with collagen sequences containing positively charged and hydrophobic residues. Docking of a collagen triple helix on the binding site suggests a range of possible engagements and predicts that at most eight consecutive residues in a collagen triple helix interact with A3.

Under conditions of high shear stress, platelet adhesion to collagen at sites of vascular injury is initiated by the interaction of platelet receptor glycoprotein (Gp)1 Ib-IX-V with collagen-bound von Willebrand factor (VWF) (1). Transient interactions between VWF and GpIb-IX-V mediate platelet rolling, which slows down the platelet and allows other platelet receptors such as integrin αβ3 (2) and GpVI to bind to collagen (2–4). These interactions result in firm adhesion and activation of platelets at the site of vascular injury.

VWF is a multimeric glycoprotein consisting of ~270-kDa monomers that are linked by disulfide bonds (5). The affinity of VWF for collagen depends on multimer size (6). The binding site for fibrillar collagens type I and III is located in the VWF-A3 domain (7), whereas collagen type VI has been shown to bind to the VWF-A1 domain (8, 9). The latter domain also contains the binding site for GpIbα of the GpIb-IX-V complex (10, 11).

VWF A-type domains and homologous integrin I domains adopt a so-called dinucleotide-binding fold, or Rossman fold, composed of a central β-sheet flanked on both sides by amphipathic α-helices (12–15). Binding of the I domains of integrins αβ1, αβ2, and α1β1, and α1β3 to collagen involves a divalent cation (16, 17) located in the metal ion-dependent adhesion site (MIDAS) motif, and amino acid residues at the top face of the domain (18, 19). Binding of the I domain of integrin αβ1 to collagen induces a major displacement of its carboxyl-terminal α-helix that is thought to be critical for integrin signaling (18). The A3 domain of VWF does not contain a functional MIDAS motif, and binding of A3 to collagen is cation-independent (20, 21). The involvement of the top face of A3 in collagen binding has been excluded by mutagenesis studies (13, 22). Recently, we showed that His1023, located close to the edge of the front and bottom face of A3, is critical for binding of VWF to collagen (23).

In this study, we identify the collagen-binding site in the VWF-A3 domain in detail by means of site-directed mutagenesis. We constructed 22 point mutants, expressed these as multimeric VWF, and evaluated their collagen binding characteristics.

**EXPERIMENTAL PROCEDURES**

**Selection and Construction of VWF Point Mutants—**Selection of amino acid residues for mutagenesis was based on the approximate location of the collagen-binding site as identified in our previous study (23) and the crystal structure of the VWF-A3 domain (12, 13). Selected residues are solvent-exposed in the isolated A3 domain. Residues Glu966, Ser974, Ile975, Thr977, Asp979, Ser981, Asn983, Val984, Val985, Ser986, Val997, Gln966, Glu969, Gln1001, Gln1006, Asp1009, Ser1020, Glu1021, and Met1022 were mutated to alanine. In addition, Pro962, Pro965, and Pro1027 were mutated to histidine to inhibit collagen binding by steric hindrance. Backbone conformations of these proline residues suggested that the histidine side chain would protrude from the protein surface. Point mutations were introduced in the VWF-A3 domain using the QuikChange method (Stratagene, La Jolla, CA) as described previously (23).

**Expression, Purification, and Characterization of VWF—**VWF was...
Collagen-binding Site of VWF-A3

Characterization of VWF-A3 Mutants—The conformation of the A3 domain and the multimeric size of VWF determine its reactivity toward collagen. Multimer distributions of VWF mutants and recombinant wt-VWF were indistinguishable (data not shown). The conformation of the A3 domain was evaluated with conformation-dependent monoclonal antibody RU5. Binding of RU5 to 18 of the 22 point mutants was similar to binding to wt-VWF. As expected, RU5 did not bind to ΔA3-VWF, a deletion mutant lacking the A3 domain (7) (Fig. 1A). Four of the point mutants, P962H, P981A, V984A, and V985A, showed a significantly reduced RU5 binding, ranging from 60% to 80% compared with wt-VWF. However, collagen binding of these mutants was normal (see below). Reduced RU5 binding can be explained by direct interactions of the mutated residues with collagen. Immobile collagen was incubated with expression medium containing 0.1 μg/ml of VWF. Binding of RU5 to 18 of the 22 point mutants was similar to binding to wt-VWF. As expected, RU5 did not bind to ΔA3-VWF, a deletion mutant lacking the A3 domain (7) (Fig. 1A). Four of the point mutants, P962H, P981A, V984A, and V985A, showed a significantly reduced RU5 binding, ranging from 60% to 80% compared with wt-VWF. However, collagen binding of these mutants was normal (see below). Reduced RU5 binding can be explained by direct interactions of the mutated residues with collagen. Immobile collagen was incubated with expression medium containing 0.1 μg/ml of VWF. Binding of ΔA3-VWF to collagen was only 8% compared with wt-VWF. Mutations I975A, T977A, D979A, P981H, V997A, E1001A, and S1020A strongly reduced collagen binding, showing a residual binding of less than 25%.

To further investigate the effect of the seven mutations,
these mutants and mutant H1023A from our previous study (23) were purified by immuno-affinity chromatography, and collagen binding was analyzed by SPR. We also purified and analyzed wt-VWF and mutants P981A and E1021A that bound normally to collagen in the solid-state collagen binding assay. The multimer distribution of the VWF variants after purification was similar (Fig. 2).

Addition of the RU5 Fab fragment to wt-VWF almost completely inhibited collagen binding, confirming that the A3 domain contains the binding site for collagen type III (Fig. 3). Mutants P981A and E1021A, which bound normally to collagen in the solid-state collagen binding assay, also had similar binding affinities and a similar number of binding sites as wt-VWF in the SPR-based collagen binding assay (Fig. 3). Mutants that exhibited strongly reduced collagen binding in the solid-state assay divided into two populations in the SPR collagen binding assay. Mutations I975A, T977A, V997A, and E1001A reduced the affinity of VWF for collagen 5-10-fold, whereas at saturation, the number of binding sites was at least 60% compared with wt-VWF (Table I). Mutants D979A, P981H, S1020A, and H1023A had a residual binding at saturation of less than 20% with wt-VWF (Table I). Mutants I975A, T977A, V997A, and E1001A reduced the affinity of VWF for collagen 5-10-fold, whereas at saturation, the number of binding sites was at least 60% compared with wt-VWF (Table I). Mutants D979A, P981H, S1020A, and H1023A had a residual binding at saturation of less than 20% compared with wt-VWF.

As shown in Fig. 4A, mutations that reduce collagen binding are located at the front face of the domain and define a rather flat collagen-binding site. Mutations at the bottom face did not have an effect. Surface properties of the collagen-binding site are shown in Fig. 4, B and C. The upper part of the collagen-binding site contains a small negatively charged patch. In addition, the collagen-binding site contains one large hydrophobic patch and two smaller hydrophobic patches.

Docking of a Collagen Triple Helix on A3—The amino acid sequence of collagen that is recognized by the VWF-A3 domain is not known. Under these circumstances, the use of automated docking procedures, such as FTDock (26) and AutoDock (27), that use scoring functions based on shape complementarity and interaction energies is not meaningful. Therefore, we used the interactive molecular graphics program O (28) to obtain an impression of possible collagen-binding modes compatible with structural and mutagenesis data. Because the amino acids of A3 that are involved in collagen binding define an extended and rather flat surface at the front face of the domain, a bound collagen triple helix must lie nearly parallel to the front face. A model of a triple helix restricted to lie parallel to the front face of A3 was rotated and translated with respect to A3, and two criteria were evaluated to select possible binding modes. Firstly, the collagen triple helix should contact (d < 4 Å) all residues of A3 involved in collagen binding. Secondly, monoclonal antibody RU5 bound to A3, as observed in the crystal structure of the A3-RU5 complex (23), should block (d < 2 Å) binding of collagen by steric hindrance. For the evaluation of distances, we assumed a fairly large radius for the collagen

e triple helix of 9 Å, which corresponds to the approximate distance from the tip of an extended lysine side chain to the center of the triple helix. Using a smaller radius would have reduced the range of possible binding modes, but in the absence of knowledge about the collagen residues actually involved, this did not seem justified.

A range of orientations of the collagen triple helix fulfilled the criteria (Fig. 4D). In these potential binding modes, the collagen triple helix lies at an angle of about 60° to 90° to strand β3 (located at the front face of the domain) and interacts with the A3 domain via six to eight consecutive residues.

**DISCUSSION**

Von Willebrand factor A-type domains are found in many proteins including collagens, complement proteins, and integrins, where they are named I domains. These proteins are involved in several biological functions such as cell-cell interaction and ligand-receptor binding. In integrins, these interactions involve a divergent cation present in the MIDAS motif located at the top of the domain.
VWF contains three A-type domains, of which the A3 domain binds to collagen. The VWF-A3 domain does not contain a functional MIDAS motif, and collagen binding is cation-independent (20, 21). Previously, we excluded the top face of A3 from being involved in collagen binding (22) and showed that His\textsuperscript{1023}, located close to the edge of the front and bottom face, is critical for binding of VWF to collagen (23). Based on these results, we constructed a panel of 22 point mutants in which solvent-exposed residues were mutated to either alanine or histidine. These mutations were introduced in multimeric VWF, and the binding of these VWF mutants to collagen type III was evaluated.

In a solid-state collagen binding assay, 7 of the 22 mutants, namely, I975A, T977A, D979A, P981A, V997A, E1001A, and S1020A, displayed reduced collagen type III binding. None of these mutations are located at the bottom face of A3, excluding its involvement in collagen binding. We further characterized these seven mutants and mutant H1023A, which also displays strongly reduced collagen binding (23), by SPR. In contrast to the solid-state binding assay, SPR analysis measures collagen binding under equilibrium conditions and is not affected by washing steps. The apparent dissociation constant for binding of wt-VWF to collagen type III as determined by SPR was 3.3 nM, which is similar to values previously determined by us and others (22, 29).

Interestingly, mutants that were qualified as "strongly reduced" in the solid-state assay separated in two groups in the SPR analysis. Mutants D979A, S1020A, and H1023A displayed a large reduction in affinity and in the number of binding sites, showing that these residues are critical for collagen type III binding. In contrast, mutants I975A, T977A, V997A, and E1001A were characterized by a 5–10-fold reduced affinity but had a near normal number of binding sites, indicating that these residues contribute to collagen binding but are not essential. Residues essential for collagen binding are located in strand \(\beta_3\) and loop \(\alpha_3\beta_4\) in the lower half of the front face of A3, whereas nonessential residues are located in the upper half of the front face in loops \(\beta_2\beta_3\), \(\alpha_2\alpha_3\), helix \(\alpha_3\), and strand \(\beta_3\) (Fig. 3A).

Mutation of Pro\textsubscript{981} to alanine did not affect collagen binding, whereas mutation to histidine markedly decreased VWF binding to collagen. Apparently, Pro\textsubscript{981} is not required for binding to collagen type III, but the introduction of a bulky histidine side chain at the lower half of the front face of A3 interferes with collagen binding via sterical hindrance. This observation further supports our conclusion that the collagen-binding site is located at the front face of A3.

Docking of a collagen triple helix on the front face of the A3 domain suggested a range of possible engagements (Fig. 4D) and predicts that at most eight consecutive residues in a collagen molecule interact with A3. Based on the surface characteristics of the collagen-binding site (Fig. 4, B and C), we propose that collagen sequences recognized by A3 contain positively charged and hydrophobic residues.

Despite their similarity in fold and ligand, VWF-A3 and collagen-binding integrin I domains have distinctly different binding sites. In A3, a rather hydrophobic and flat binding site to collagen. Apparently, Pro\textsubscript{981} is not required for binding to collagen type III, but the introduction of a bulky histidine side chain at the lower half of the front face of A3 interferes with collagen binding via sterical hindrance. This observation further supports our conclusion that the collagen-binding site is located at the front face of A3.


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binding that makes them particularly suited for signal transduction and modulation of ligand binding affinity (18). In contrast, the VWF-A3 domain appears to function as an independent structural unit, and there is no evidence for modulation of its collagen binding affinity, nor does binding of A3 to collagen affect the affinity of the VWF-A1 domain for platelet receptor GpIbα. Thus, the collagen-binding site of A3 merely performs an adhesive function, whereas binding sites of I domains are more sophisticated and also play a regulatory role (31).

Recently, we determined the crystal structure of the VWF-A1 domain in complex with an amino-terminal fragment of platelet GP Ibα. Like A3, the A1 domain does not contain a MIDAS motif. GpIbα binds to two distinct sites on A1. The larger of the two binding sites is located at the front face of A1 and consists of residues from strand β3, helix α3, and loop α3β4. Interestingly, the same structural elements contribute residues to the collagen-binding site of A3, suggesting that ligand-binding sites in A-type domains that lack a MIDAS motive may all be located in a similar position at the front face of the domain.

Genetic screening has identified four mutations, S968T, Q971H, I978T, and Q999R, that affect binding of VWF to collagen (32, 33). These mutations are located at or just below the front surface of the domain, and their effect on collagen binding is consistent with our mutagenesis data. Surprisingly, these mutations caused no (32) or only moderate (33) bleeding phenotypes. Wu et al. (34) recently showed that collagen binding by the A3 domain is relevant because an antibody blocking the A3 domain is relevant because an antibody blocking the A3 domain is relevant because an antibody blocking the VWF-A3-collagen interaction prevented the formation of platelet-rich thrombi and prolonged the skin bleeding time at high doses. Further investigations are required to reconcile these conflicting observations and to establish the physiological importance of VWF-A3 in platelet adhesion.

After submission of this manuscript, Nishida et al. (35) reported mapping of the collagen-binding site of the VWF-A3 domain using a novel NMR technique. Their findings with A3 are consistent with our data. In summary, the binding site for collagen type III is located at the front face of the VWF-A3 domain. Residues in the lower half of the collagen-binding site are essential for collagen binding, whereas residues in the upper half contribute to binding but are not essential. We suggest that a collagen triple helix that interacts with A3 contains hydrophobic and positively charged residues. Further understanding of the VWF-collagen interaction requires the identification of specific collagen sequences involved in VWF binding.

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