Structural Model of Porcine Factor VIII and Factor Villa Molecules Based on Scanning Transmission Electron Microscope (STEM) Images and STEM Mass Analysis

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Abstract

Porcine plasma factor VIII (fVIII) molecules are heterodimers composed of a 76,000-mol wt light chain (−A2−C1−C2) and a heavy chain ranging in molecular weight from 82,000 (A1−A2) to 166,000 (A1−A2−B). Proteolytic activation of fVIII by thrombin results in fVIIIa heterotrimers lacking B domains (A1, A2, A3−C1−C2). In this study, immunoaffinity purified fVIII was further fractionated by mono S or mono Q chromatography to prepare heterodimers containing a light chain and an A1−A2−B heavy chain (fVIII 166/76) or an A1−A2 heavy chain (fVIII 82/76). Mass analysis of scanning transmission electron microscopic (STEM) images of fVIII 166/76 indicated that heterodimers (mass 237±20 kD) had irregularly globular core structures 10–12 nm across, and frequently displayed a diffuse, occasionally globular to aoid satellite structure extending 5–14 nm from the core, and attached to it by a thin stalk. Factor VIII 82/76 molecules (mass 176±20 kD) had the same core structures as fVIII 166/76 molecules, but lacked the satellite structure. These findings indicate that A1−A2 domains of heavy chains and the light chains of the fVIII procofactor molecule are closely associated and constitute the globular core structure, whereas the B domainal portion of heavy chains comprises the peripheral satellite appendage. Factor VIII core structures commonly displayed a finger-like projection near the origin of the B domainal stalk that was also a consistent feature of the free heavy chains (mass 128–162 kD) found in fVIII 166/76 preparations. Factor VIII light chain monomers (mass, 76±16 kD) were globular to c-shaped particles 6–8 nm across. These chains commonly possessed a v-shaped projection originating from its middle region, that could also be observed at the periphery of fVIII core molecules. Factor VIIIa preparations contained heterotrimers (mass 162±13 kD) that had the same dimensions as fVIII core structures, lacked the B domainal appendage, and sometimes possessed the same core features as fVIII molecules. Molecular species corresponding to heterodimers (mass, 128±13 kD) and unassociated subunit chains (40–100 kD) were also observed in fVIIIa preparations, suggesting that heterotrimers have an appreciable tendency to dissociate, a phenomenon that could explain the decay of fVIIIa activity after thrombin activation of fVIII. (J. Clin. Invest. 1990. 85:1983–1990.) electron microscopy * scanning transmission electron microscopy * factor VIII * factor VIIIa

Introduction

Factor VIII (fVIII)1 is a plasma glycoprotein that is absent or abnormal in hemophilia A. In humans, it is synthesized as a mature polypeptide chain of 2,332 amino acids and contains three types of domains (i.e., A, B, C) in the sequence NH2−A1−A2−B−A3−C1−C2−COOH (1, 2) (Fig. 1). Because of cleavages between the B and A3 domains, perhaps occurring intracellularly (4), fVIII exists in porcine plasma as a divalent cation-linked heterodimer comprised of a heavy chain of variable size and a light chain (−A2−C1−C2) of mol wt 76,000 (5). Heavy chain size heterogeneity is due to proteolytic cleavages within the B domain resulting, with respect to porcine fVIII, in heavy chain peptides of mol wt 166,000 (A1−A2−B), 130,000 (A1−A2−B), or 82,000 (A1−A3) (5, 6). Likewise, human fVIII molecules are heterodimeric, having subunits similar in size to those of porcine fVIII (1, 7–10).

The molecular weight of partially to highly purified human or porcine fVIII, based upon hydrodynamic parameters (7, 10–12) or upon radiation inactivation experiments (13, 14), has been estimated to lie between 140,000 and 285,000. The rather wide range of values obtained may depend to some extent (e.g., reference 7) upon the size heterogeneity of heavy chain components. Summation of the molecular weights of the various constituents of porcine fVIII molecules yields values ranging from 158,000 (A1−A2−A3−C1−C2) to 242,000 (A1−A2−B−A3−C1−C2) (5, 6).

The addition of thrombin to fVIII causes a rapid increase in procoagulant fVIIIa activity, and is accompanied by cleavage between the A2 and B domains, between the A1 and A2 domains, and near the NH2 terminus of the A3 domain (Fig. 1) (1, 3). The resulting fVIIIa molecules form heterotrimers consisting of two heavy chain fragments (A1, A3) plus a light chain component (A3−C1−C2), having a molecular weight of 161,000, as determined by sedimentation equilibrium analysis (15). Although thrombin activation is associated with proteolysis of its constituent peptides (5, 8, 12) and reduction of its size (11, 12, 15, 16), loss of fVIIIa activity after activation (6,

1. Abbreviations used in this paper: fVIII, factor VIII; fVIIIa, factor VIIIa; STEM, scanning transmission electron microscopy.

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17–21) is not prevented by inhibitors of thrombin (11, 21).

This indicates that loss of activity is not due to continued proteolysis by thrombin, a conclusion also supported by the results of Lollar et al. (6), who showed that factor IXa and phospholipid in the presence of calcium ions stabilized the activity of thrombin-activated porcine fVIIIa, although the size of fVIIIa components, as determined by SDS gel electrophoresis, did not differ from that of fVIIIa activated in the presence of thrombin alone.

Immunooaffinity chromatography of partially purified porcine fVIII has provided an effective method for purifying stable, though heterogeneous, fVIII (5). Further processing of immunooaffinity-purified fVIII by HPLC on mono S or mono Q columns has permitted separation of biologically active material into discrete subpopulations of heterodimers, and removal of a high molecular weight constituent devoid of coagulant activity (7, 15, 22). In addition, highly stable and active fVIIIa preparations have been prepared by HPLC of thrombin-activated fVIII on mono S columns (15). One of the main objectives of studying highly purified and stable components of fVIII and fVIIIa has been to facilitate elucidation of their structure in relation to their coagulant function. In this investigation, we have approached that objective using scanning transmission electron microscopy (STEM) coupled with STEM mass analysis to identify and characterize the ultrastructure of fVIII, fVIIIa, and their constituent peptides. A structural model of fVIII based upon these observations is offered.

Methods

Preparation of porcine fVIII and fVIIIa. Immunooaffinity-purified porcine fVIII was prepared as described (5) using a monoclonal antibody to the fVIII light chain (W3-3) that had been bound to Sepharose 4B (Pharmacia, Inc., Piscatway, NJ). This material was further chromatographed on a mono S or mono Q column (Pharmacia, Inc.; reference 15) to remove a high molecular weight constituent lacking fVIII activity, and to isolate a fVIII heterodimer fraction with a predominant heavy chain population of 166 kD (fVIII 166/76), and one with an 82 kD heavy chain (fVIII 82/76) (Fig. 2). Three such preparations were stored at 5°C in 10 mM histidine, 5 mM CaCl2, pH 6.0 buffer, containing 0.5–0.7 M NaCl (mono S) or in 20 mM Tris, 5 mM CaCl2, 0.7 M NaCl, pH 7.4 buffer (mono Q) at concentrations of 46–280 µg/ml. Factor VIIIa was prepared by chromatography on a mono S column as described (15) and stored at 5°C at concentrations of 80 or 173 µg/ml (two preparations). Factor VIII light chains (one preparation) were prepared as described (15) and stored at 5°C in 20 mM Tris, 5 mM CaCl2, 0.45 M NaCl, pH 7.4 buffer at a concentration of 140 µg/ml.

STEM. High-resolution STEM was performed at the Brookhaven National Institutes of Health Biotechnology Resource Facility. The preparation of ultrathin carbon films and the method of specimen preparation and application for freeze-drying were as described (23, 24). Briefly, protein solutions for analysis were diluted to a final concentration of 5–10 µg/ml with 0.15 M NaCl, 1 mM CaCl2, 10 mM Hepes buffer, pH 7.0, and injected into a droplet of the same buffer on a grid coated with an ultrathin (~3 nm) carbon film. After a 1-min attachment time, the specimen on the film was washed 8–10 times with 50 mM ammonium acetate, pH 7.0 solution to remove nonvolatile salts, freeze-dried slowly overnight, transferred to the STEM micro-

Figure 1. Schematic model of the domainal structure of single-chain fVIII and the structures of the three commonly identified heterodimeric fVIII molecules that are isolated from porcine plasma. Cleavages of single-chain fVIII that result in the heavy and light chain populations found in plasma are indicated, as well as thrombin (T) cleavages of the heavy and light chains that accompany its activation to fVIIIa (1, 3). Heavy chains and light chains are linked noncovalently by a divalent cation (Me2+)–dependent mechanism. Large heavy chains containing B domainal structures are designated A3–A2–B (mol wt 166,000) or A4–B2 (mol wt 130,000); small heavy chains, A4–A3 (mol wt 82,000); fVIII light chains, A4–C1–C2 (mol wt 76,000); fVIIIa light chains, A4–C1–C2 (mol wt 69,000).

Figure 2. Composite SDS Laemmli PAGE of the nonreduced silver-stained fVIII, fVIIIa, and fVIII light chain preparations used in this study. The apparent molecular weights (x10−2) of the components are indicated. Lane 1, fVIII 166/76; lane 2, fVIII 82/76; lane 3, fVIII light chains; lane 4, fVIIIa.
scope stage (–140°C) under vacuum, and imaged using a 40-kV probe focused at 0.25 nm. For some experiments, specimen wash solutions also contained 1 mM CaCl₂.

Image resolution under the specimen sampling conditions (1 nm/ pixel) and the electron beam dose we used (1,000 e⁻/nm²) was ~ 2 nm (24). Particle mass measurements were performed off-line using a “circle of integration” program (23–25) with a variable integration radius depending upon the shape and area covered by any given object. Tobacco mosaic virus particles that had been added to the specimen were used as an internal mass calibration standard (26, 27). The accuracy of STEM mass measurements ranges from ± ~ 15 kDa at an integration radius (r) of 10 nm to ± ~ 25 kDa at an r of 20 nm (24). The mean mass values reported for the various populations of molecules found within a specimen were computed over a range of values that encompassed the species of interest.

STEM images were reproduced from a televised tape file onto Polaroid type 665 film and the negative was then acquired through a video camera (DAGE-MTI, Michigan City, IN) into an image processing system (model 8502, Tracor Northern, Middleton, WI). The images were then filtered to reduce background noise using a nonlinear grey level contrast processing program, and occasionally were smoothed using a median smoothing function to reduce the prominence of pixel lines. The filtered images were recovered by photographing the processed images on the computer TV screen. Assessments of the presence and form of subdomainal molecular features were made from filtered images.

Results

Factor VIII. Mass analysis of images of a FVIII 166/76 preparation revealed particles in a range of 70–450 kDa (Fig. 3). The

![Figure 3](image-url)

**Figure 3.** STEM images selected from FVIII 166/76 preparations, plus a frequency histogram of the size distribution (number of particles vs. mass) of particles in one such preparation (r, 18 nm). The data in this histogram were obtained from a different FVIII 166/76 preparation than that presented in Fig. 4 and in the text. The pattern of particle distribution is essentially the same for both, but the mean value for the mass in the range 210–290 kDa is insignificantly higher, 255±22 kDa, than that found for the preparation reported in Fig. 4, 237±20 kDa. The value in the 130–210 kDa range, 179±20 kDa, is virtually the same as in the preparation reported in Fig. 4, 177±19 kDa. Panels a–e represent molecules selected from the size range shown by the dot filled bars, except in panel a in which only molecule I (58 kDa) fulfills that requirement. Molecule 2 (162 kDa) represents a free heavy chain, whereas molecule J represents a small FVIII heterodimer (193 kDa). Panels f–h show molecules in the size range > 350 kDa. Panels i–n were selected from the size range shown by the striped bars, and correspond to free heavy chains (k–n) and small factor VIII heterodimers (i and j). Arrows indicate fingerlike structures; split lines, v-shaped structures. Satellite structures are indicated by the brackets labeled B. Bar (panel a), 20 nm.

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same mass distribution and particle shapes were observed whether or not 1 mM CaCl₂ was included in the specimen wash buffer (data not shown). Most objects within the range 130–290 kD were irregularly globular particles varying from 8 to 14 nm across. At an integration radius of 9–11 nm, the circle of integration centered about these objects, a single population of molecules between 140 and 220 kD (mean 178±17 kD; r₁ 11 nm) was suggested by the relatively Gaussian shape of this portion of the frequency histogram (data not shown). However, as the integration radius was increased, two distinct size populations of molecules were revealed (Fig. 4). The larger population had a mass of 237±20 kD at r₁ 19 nm (range 210–290 kD), and corresponded to heterodimeric fVIII 166/76 molecules (Table I). The mass of the second population of molecules did not increase significantly above r₁ 11 nm and had a mean value of 177±19 kD (range 120–200 kD) at r₁ 19 nm.

Examination of the larger fVIII 166/76 heterodimer population revealed irregular globular core structures 8–12 nm across, which often possessed a smaller, globular to ovoid, relatively less dense, satellite structure extending 5–14 nm from the denser core and connected to it by a thin stalk 1–2 nm wide (Fig. 3, B, panels a, b, c and e). This structure was sometimes diffusely spread on the surface and did not always terminate in a well-defined globular configuration (e.g., Fig. 3, panel f). Many molecules corresponding in mass to a heterodimeric fVIII 166/76 structure (e.g., Fig. 3, panel d) did not display a discrete satellite structure, suggesting that in these cases this region of the molecule was superimposed upon the fVIII core. The dimensions of the core structure of such particles tended to be somewhat larger (up to 14 nm) and the shape more irregular than that of molecules with recognizable peripheral satellite structures.

Recognizable subdomainal features appearing in profile at the periphery of core structures were a fingerlike projection (17 of 56 structures; 30%) ~3–5 nm long, arising near the origin of the stalk of the satellite structure (e.g., Fig. 3, arrow, panels b, e, and f), and a thin v-shaped projection (15 of 57; 26%), each limb of which protruded 2–3 nm from the core (e.g., Fig. 3, panels b and c, split lines). This latter structure was situated opposite to the origin of the stalk of the satellite structure.

Particles with masses of 300–450 kD tended to have “core” structures that were larger and more irregular than heterodimeric fVIII 166/76 structures (Fig. 3, panels g and h), consistent with them representing aggregates of two fVIII 166/76 molecules or perhaps two or more heavy chains, or some combination of these. Some heterodimeric fVIII 166/76 molecules with a high mass were situated so near to other smaller structures that both were included in the mass measurement (e.g., Fig. 3, panel f). Particles with masses <110 kD represented 5–7% of the objects observed, and were not investigated further in these studies.
Evaluation of particles with masses between 120 and 200 kD revealed that there were two discernible molecular forms within this size range. The first had mass values of 180–200 kD and corresponded in shape and size to a fVIII heterodimeric core structure with a truncated heavy chain (e.g., Fig. 3, panel a—molecule 3; panels i and j). The second form had masses between 128 and 162 kD and corresponded in size to large free heavy chains (i.e., A1–A2–B/ A1–A2–B). These particles were irregularly globular and smaller (~8 nm) than the heavy chain/light chain core structures (Fig. 3, panel a—molecule 2; panels k–n). One consistent aspect of these structures (seven of eight; 88%) was the presence of a bent fingerlike projection (e.g., Fig. 3, arrow, panels a, k, and n) corresponding in appearance to these seen at the periphery of fVIII core structures. Another feature was the presence of thin irregularly curled strands 1–2 nm wide, resembling the stalk of the satellite structure (e.g., Fig. 3, panel a—molecule 2, panel k).

Mass analysis of a fVIII 82/76 preparation showed particles in a range of 35–400 kD (Fig. 5). The same mass distribution and particle shapes were observed whether or not 1 mM CaCl2 was included in the specimen wash buffer (data not shown). Objects smaller than 120 kD probably reflect the presence of free heavy chains or light chains. Particles with masses above 220 kD were rare. Molecules in the 140–220-kD range had the same globular shape noted in fVIII 166/76 core structures, and commonly (13 of 30; 43%) possessed a fingerlike (Fig. 5, arrows, panels a–c) and/or v-shaped (9 of 30; 30%) projection (Fig. 5, split line, panel b). However, fVIII 82/76 molecules uniformly lacked the satellite structure seen in fVIII 166/76 molecules, and because of this, had a smaller mass (176±20 kD) that could be encompassed at an integration radius of 10 nm (Fig. 4) and that corresponded to the estimated size of heterodimers containing an –A1–C1–C2 light chain and an A1–A2 heavy chain.

**Factor VIIIa.** Mass analysis of fVIIIa preparations revealed a polydisperse distribution of particles in a range of 40–290 kD (Fig. 6), and indicated that several populations of molecules were present. These corresponded to that expected for dissociated chains (40–100 kD), fVIIIa heterodimers (110–140 kD), fVIIIa heterotrimers (140–190 kD), and particles >190 kD. The same mass distribution and particle shapes were observed whether or not 1 mM CaCl2 was included in the specimen wash buffer (data not shown).

Molecules in the fVIIIa heterotrimer mass range appeared very similar in shape and size to the core structures observed in the fVIII 166/76 (Fig. 3) and fVIII 82/76 (Fig. 5) preparations, and none possessed a satellite appendage. Consequently, their mass was entirely encompassed at an r1 of 10 nm (162±13 kD; Fig. 4). Owing to irregular serrations in the contour of fVIIIa heterotrimers, plus the absence of a satellite structure to help
orient the location of peripheral core features, identification of a fingerlike (13 of 37; 35%; Fig. 6, arrows, panels a, c, and h), and particularly, a v-shaped projection (8 of 37; 22%; Fig. 6, split lines, panels a, e, and g) was sometimes ambiguous. A small proportion of fVIIIa heterotrimers (4 of 37; 11%) possessed both v-shaped and fingerlike appendages in the same general relationship as was found in fVIII molecules (Fig. 6, panel a).

Molecules in the heterodimer mass range (128±13 kD) tended to be globular or sometimes oblong (Fig. 6, panel p) in shape and somewhat smaller than heterotrimers (8–10 nm). Commonly (13 of 16; 80%) they possessed a fingerlike structure (Fig. 6, panels m, p, and q), or less commonly (3 of 16; 19%) a v-shaped projection (Fig. 6, panels n and r). Particles < 100 kD varied in shape from globular (Fig. 6, panels t, w, and x) to c-shaped (Fig. 6, panels s and v) or oblong (Fig. 6, panel u), and commonly displayed either the v-shaped or the fingerlike feature characteristic of assembled core structures and fVIII subunit chains (Fig. 6, arrow, panel u; split line, panel s). The few particles > 190 kD (panels i–l) were globular in shape and resembled heterotrimetric structures. Their identity is uncertain but they probably represent fVIIIa heterotrimers in association with a fVIIIa subunit (Fig. 6, panels i, k, and l), or a molecule situated in the vicinity of an uncomplexed small peptide whose mass had been included within the radius of integration (Fig. 6, panel j).

Factor VIII light chains. Mass analysis of a fVIII light chain preparation indicated that in addition to a monomeric population of chains with masses between 45 and 105 kD (mean, 76±16 kD), there were dimeric and higher-order light chain aggregates, some of which were outside the range shown in the histogram (Fig. 7). Monomeric chains tended to be c-shaped (Fig. 7, panels a–c) or irregularly globular (Fig. 7, panels d–f) structures 6–8 nm across. Commonly, a v-shaped projection (7 of 30; 23%) was evident (Fig. 7, panels b, d, and e) and appeared to originate from the middle region of the chain.

Such “precursor” fVIII light chains resembled c-shaped particles corresponding in shape and mass to dissociated fVIIIa light chains (e.g., Fig. 6, panel s, 74 kD).

Discussion

Porcine plasma fVIII heterodimers are composed of a heterogeneous population of heavy chains, ranging in molecular weight from 166,000 to 82,000 plus a light chain of mol wt 76,000 (≈A₁–C₁–C₂) (5, 6) (Fig. 1). These forms of fVIII, including the porcine 82/76 species and those of human origin, require proteolytic activation to participate as a cofactor for factor IXa during the activation of factor X (7). The B domain does not appear to have a coagulant function since plasma-derived fVIII heterodimers lacking this domain have similar coagulant potential as B domain-containing heterodimers (7) and since deletion of most of the B domain from recombinant human fVIII by site-directed mutagenesis does not abolish coagulant potential (28, 29). Activation of porcine fVIII to fVIIIa by thrombin is accompanied by cleavages between the A₁ and A₂ domains, between the A₂ and B domains, and within the light chain near the amino terminus to produce a 69,000-mol wt derivative (A₁–C₁–C₂) (5, 6). The resulting heterotrimeric fVIIIa molecule consists of the A₁, A₂, and A₂–C₁–C₂ domains (15). These present studies coupling mor-
phological and mass analysis of high resolution STEM images, have provided the first detailed views of the ultrastructure of these molecules, and have enabled us to propose a model of the structure of fVIII and its activation product, fVIIla (Fig. 8).

Mass analyses of fVIII molecules confirmed the molecular weight value determined by physical measurements or that predicted by summation of its constituent subunits (Table 1). They indicate that components of fVIII 166/76 molecules extend well beyond the vicinity of its core structure (Fig. 4). These observations indicate that fVIII 166/76 molecules consist of a compact globular core of 10–12 nm that accounts for ~75% of its total mass, plus a peripheral satellite structure connected to it by a thin stalk, that accounts for the remainder. Factor VIII 82/76 molecules consist of a core structure that is indistinguishable from that of fVIII 166/76, but they lack a satellite structure, and thus their mass is fully accounted for by the core structure itself (Fig. 4). These findings clearly indicate that the A₁-A₂ heavy chain and the light chain components of the fVIII procofactor molecule are closely associated with one another to form the globular core structure, whereas the B domainal portion of the heavy chain is represented by the peripheral satellite appendage.

Factor VIIIa molecules consisted of several populations corresponding in mass to heterotrimers, heterodimers, and dissociated subunits. Molecules in the heterotrimer range were very similar in shape and size to the fVIII core structures, and none possessed a satellite appendage. Thus the globular structure comprising fVIIIa heterotrimers is derived directly from the heavy chain and light chain components of the fVIII core structure.

There were two subdomainal structures observed in fVIII core molecules: a fingerlike projection arising in the vicinity of the B domainal stalk and a v-shaped appendage situated opposite to the origin of the stalk. Visualization of these structures in any given molecule depends upon their proper extension in profile from the core and/or upon appropriate orientation of molecules on the grid surface, and could account for the relatively small proportion of molecules (26–30% in fVIII 166/76) displaying such features. V-shaped structures, in particular, are thinner and less dense than fingerlike structures, and may be more readily subject to beam damage, or more difficult to appreciate above the background noise from the carbon film itself, or both. Additionally, even when properly positioned, these features may become obscured or their identification made more ambiguous when the B domain is superimposed upon the core structure, such as sometimes occurs in fVIII 166/76 molecules (e.g., Fig. 3, panel d).

As assessed by STEM mass analysis, there were free large heavy chains in fVIII 166/76 preparations. These chains were smaller and more irregular than fVIII core structures, and possessed irregularly curled strands resembling the stalk of the B domainal satellite structure. A fingerlike projection was a prominent and consistent feature of these chains and closely resembled the fingerlike structure seen in fVIII core molecules, indicating that they represent the same region of the fVIII molecule.

Factor VIII light chains (A₁-C₁-C₂), which encompass the carboxy-terminal region of the molecule, tend to have a c-shape which seems to be preserved in free light chains (A₁-C₁-C₂) found in fVIIla preparations (e.g., Fig. 6, panel d). Owing to the small differences in size between fVIII and fVIIla light chains, we cannot distinguish them from one another by this type of analysis. A v-shaped projection is a feature of both fVIII as well as fVIIla light chains, and can also be identified at the periphery of assembled core structures in fVIII 166/76, positioned opposite to the stalk of the satellite appendage.

These observations permit us to suggest the orientation of light chains in fVIII core molecules relative to its heavy chains (Fig. 8), based upon (a) the relative positions of v-shaped and fingerlike projections in core structures and (b) the assumption that uncomplexed light chains retain the same general shape as they do when they are complexed with heavy chains. The same two features were detected in some fVIIla molecules in the same relative positions. This observation is consistent with the notion that the basic arrangements of light and heavy chains are preserved in the conversion of fVIII to fVIIla; this issue will require further investigation.

As assessed by mass analysis, free heavy chains and free light chains were evidently present not only in images of fVIII 166/76 preparations but also in fVIII 82/76 preparations. We have also detected them in images of immunoaffinity purified fVIII preparations before mono S or mono Q chromatography (data not shown). They occur as well in fVIII preparations in which the specimen wash buffer contained 1 mM CaCl₂, thus eliminating the possibility that the absence of calcium ions, per se, from buffer solutions during specimen preparation might have contributed to their generation from heterodimeric forms during processing. It is possible that dissociation of heavy and light chains occurs after specimen dilution for application to the EM grids, and that their appearance represents a sort of dilutional "artifact." This would be consistent with the observation that multiple boundaries have not been observed during analytical velocity sedimentation of fVIII 166/76 preparations (30). It is interesting to note in this context that we have made several attempts to prepare fVIII heavy chains for EM analysis by releasing them from W3-3 affinity columns with EDTA-containing buffer solutions. These attempts at imaging monomeric chains have been uniformly unsuccessful, in that at most, we have observed aggregates of protein (data not shown). It may be that heavy chains have a tendency to self-associate in the absence of light chains.

Based upon ultracentrifugal analyses of fVIIla (15), it seems likely that the fVIIla preparations we studied exist in stock solution mainly as heterotrimers. Sedimentation equilibrium experiments at loading concentrations of 60–160 µg/ml showed a concentration dependent increase in apparent molecular weight from 148,000 to 161,000 (15). Sedimentation velocity analyses carried out at a concentration of 50 µg/ml revealed a single boundary sedimenting at 7.2S, thus excluding the possibility that there were large amounts of free light chains or heavy chain fragments in the preparation. STEM mass analysis of fVIIla preparations indicated that heterotrimers were the predominant form, but heterodimers and dissociated subunits were also present in substantial amounts. Heterotrimeric fVIIla structures resembled fVIII core structures, both in shape and submolecular detail. Heterodimers tended to be smaller than heterotrimers but many retained the heavy chain or light chain landmark features of heterotrimeric or fVIII core structures from which they had originated. Considering the characteristics of the fVIIla stock solutions as summarized above, it seems that the molecular heterogeneity we observed in our STEM images of fVIIla is the result, to some extent, of subunit dissociation occurring during specimen dilution and washing that accompany processing for subsequent STEM imaging.

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The model we have proposed for fVIII and fVIIa molecules should provide a useful tool for investigating and refining our understanding of the functional features of their domains, the nature of their complexation with other molecules, and for comparing their structure with that of other related proteins such as factor V (31, 32). Given the high degree of sequence and functional homology between fVIII and fV molecules, it is no surprise that their structures resemble one another. However, there are some noteworthy differences. The satellite structures (B domains) of fV molecules tend to extend further from the core structure than those of fVIII (up to 35 nm vs. up to 14 nm). More importantly, in contrast to fVIIa, there is evidence for only a small degree of dissociation of factor Va molecules. Perhaps the tendency for fVIIa molecules to dissociate explains the instability of fVIIIa activity compared with that of fV.

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