Definition of an N-Terminal Actin-binding Domain and a C-Terminal Ca\textsuperscript{2+} Regulatory Domain in Human Brevin

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Abstract. Brevin is a Ca\textsuperscript{2+}-modulated actin-associated protein that will sever F-actin and cap barbed filament ends. Limited proteolysis with chymotrypsin or subtilisin cleaves the molecule approximately in half. Cleavage is ~10-fold more rapid in Ca\textsuperscript{2+} than in EGTA. The two fragments are readily separated from each other and from undigested brevin by high pressure liquid chromatography on a DEAE resin. A 40,000-mol-wt fragment from the N-terminal is not retained by DEAE, while a 45,000-mol-wt C-terminal fragment binds more tightly than brevin. The N-terminal fragment retains ~10% of the nucleation activity, caps barbed ends, and retains 50% of the total severing activity defined by dilution induced depolymerization of pyrenyl actin, but, in contrast to brevin, none of these functions are affected by Ca\textsuperscript{2+}. Fluorescent actin binding studies and gel-filtration demonstrate that the 40,000-mol-wt fragment binds two actin monomers. The 45,000-mol-wt C-terminal fragment has no severing, nucleating, or capping activity. Cross-reaction with two monoclonal antibodies against two specific Ca\textsuperscript{2+}-induced conformations of human platelet gelsolin suggest that both Ca\textsuperscript{2+} binding sites are located on the carboxyl half of the brevin molecule. One epitope, defined as the rapidly exchanging Ca\textsuperscript{2+} binding site in the gelsolin–actin complex, is lost when a 20,000-mol-wt fragment is cleaved from the carboxyl terminal. The second epitope, related to the poorly exchanging Ca\textsuperscript{2+} binding site in the complex, is nearer the middle of the brevin molecule.
Materials and Methods

Protein Purification

Brevin was immunopurified from pooled human sera following the procedures outlined in Hwo and Bryan (9). Monoclonal antibodies were isolated from serum-free harvest fluid. The purification and characterization of 4F8 IgA and 8G5 IgG are given in Hwo and Bryan (9). Actin was prepared according to Spudich and Watt (16) and labeled with pyrene iodoacetamide as described by Bryan and Coluccio (2). Protein determinations were done by the method of Bradford (1). The brevin concentration was estimated using an extinction coefficient of 150,000/M per cm determined for bovine brevin (10).

Fluorescence Measurements

Dilution induced depolymerization experiments using pyrene F-actin were done as described by Bryan and Coluccio (2). Actin monomer binding was measured using NBD(nitrobenzoxadiazole)-actin as described by Bryan and Kurtz (3). Pyrene actin excitation and emission wavelengths were 365 and 407 nm, respectively; NBD-actin excitation and emission wavelengths were 475 and 530 nm, respectively.

SDS Gel Electrophoresis and Immunoblotting

SDS gel electrophoresis was done as outlined by Laemmli (13). Immunoblots were performed as described by Towbin et al. (19) modified as described by Hwo and Bryan (9).

Sequence of Peptides

A model 470A gas phase sequencer (Applied Biosystems, Inc., Foster City, CA) was used for all analyses. PTH amino acids were identified by high pressure liquid chromatography (HPLC) using a Beckman model 344 chromatograph equipped with two model 112 pumps and two model 160 detectors (Beckman Instruments, Inc., Palo Alto, CA). Samples were automatically injected using a model M7108 Waters Intelligent Sample Processor (WISP, Waters Instruments Rochester, MN). PTH amino acids were analyzed using a 5-μm pore size IBM cyano column (4.5 mm × 25 cm) (IBM Instruments, Inc., Danbury, CT) with either a DuPont ETH guard column or a pellicular CN guard column (DuPont Co., Wilmington, DE). The samples were analyzed at 35°C using a gradient system that consisted of buffer A (0.02 M sodium acetate, 5% tetrahydrofuran, pH 4.3), buffer B (1 M NaCl, 1% tetrahydrofuran, pH 4.3), and a 1:1 mixture of buffer A and buffer B. The PTH amino acids were eluted by increasing buffer B from 11% to 45% at 2% per minute at a flow rate of 1 ml/min.

HPLC Fractionation

Brevin and brevin digests were fractionated using a Waters HPLC apparatus. DEAE chromatography was done on a Waters DEAE-5PW column, 0.75 × 7.5 cm. The flow rate was 1 ml/min. The solvent was 25 mM Tris-HCl, pH 7.0. 0.1 mM EGTA. A gradient from 0 to 0.4 M NaCl was developed over 30 min; 0.5-ml fractions were collected and analyzed for protein by Bradford assay (1). SDS gel electrophoreses, and using various functional assays as described in the text.

Gel filtration was done using Waters Protein Pak 300SW and 251 gel-filtration columns in tandem. 10-50-μl samples at 0.5-2 mg/ml were injected. The flow rate was 1 ml/min. The solvent was 0.1 M sodium phosphate, pH 7.0. The columns were calibrated using various molecular weight standards, which include beta-galactosidase, 520,000; ferritin, 440,000; catalase, 232,000; phosphorylase, 94,000; bovine serum albumin (BSA), 67,000; ovalbumin, 43,000 and carbonic anhydrase, 30,000.

Protease Cleavage

Brevin at 1 mg/ml was cleaved with bacterial subtilisin (type VII, Sigma Chemical Co., St. Louis, MO) or Na-p-tosyl-L-lysine chloromethyl ketone-treated chymotrypsin (type VII, Sigma Chemical Co.) using various conditions as indicated in the figure legends. The buffer used was 10 mM Tris-HCl, 0.5 mM CaCl₂, 1 mM Na₂SO₄, pH 8.0 at a temperature of 20°C. The reaction was stopped by addition of EGTA to 1 mM and diisopropylfluorophosphate (DIFP) to 5 mM. The DIFP was stored as a 200 mM stock solution in propylene glycol at −20°C.

Results

The results of one experiment in which brevin was digested with subtilisin at a weight ratio of 1:100 enzyme/brevin are shown in Fig. 1a. At early times the two major cleavage products are S45 and S40, 45- and 40-kD polypeptides, respectively. These are then cleaved into two slightly smaller polypeptides; S45 yields S44 and S40 yields S38. We see similar results (Fig. 1b) using chymotrypsin at 1:100 enzyme/brevin (wt/wt) concentrations. Cleavage products, CT45 and CT40, are produced at early digestion times that have molecular weights ~45,000 and 40,000. At later times the 45-kD fragment is cleaved to a 43-44-kD fragment, and the 40-kD fragment is cleaved to pieces with apparent molecular mass of ~30 and 15 kD. Ca²⁺ liganded brevin is more accessible to both subtilisin and chymotrypsin. For comparison, the partial results from one experiment using chymotrypsin in the presence of EGTA are also shown in Fig. 1c. Comparing the extent of hydrolysis at various times, we estimate that Ca²⁺ liganded brevin is about 10 times more sensitive to proteolysis than the unliganded form. Experiments using BSA as the protease substrate show that neither chymotrypsin nor subtilisin have intrinsic calcium sensitivity.

Immunoblotting with 4F8 IgA and 8G5 IgG shows that the epitopes recognized by both monoclonal antibodies are on the 45/43-kD fragments. Selected lanes from the gels in Fig. 1 are shown in Fig. 2. More pronounced proteolysis by subtilisin generates smaller fragments of S44 each of which has only one of the epitopes. S34 is the smallest fragment we have identified that binds to 4F8 IgA, and S37 is the smallest that is recognized by 8G5 IgG (data not shown). Monoclonal antibody binding to all of the fragments is strictly dependent upon the presence of calcium.

Fig. 2 shows data that place the 8G5 epitope on one end of the brevin molecule. In some preparations we see a proteolytic fragment with a molecular weight of ~70,000-75,000. This fragment cross-reacts strongly with 4F8 IgA, but not with 8G5 IgG, indicating that the 8G5 epitope is on the missing 20-kD fragment.

Separation of Brevin Fragments by DEAE-Chromatography

Brevin and the 40- and 45/43-kD fragments are readily separated by DEAE-chromatography. Fig. 3a shows the resolution of a chymotrypsin digest of 3 mg of brevin. CT40 is not retained by the DEAE column in 25 mM Tris-HCl, 0.1 mM EGTA, pH 7.4 at 2.5-5 min. Brevin elutes at 0.19 M at ~29-30 min, and CT45/43 elutes at 0.24 M NaCl at ~34 min. The results suggest that one half of the brevin molecule, the 45-kD fragment, is considerably more acidic than the other half. Fig. 3b illustrates the purity of the three major peaks and shows the cross-reactivity with the two monoclonal antibodies.

Sequence Data

We have placed the 40-kD fragment on the amino half of the parent brevin molecule by sequencing a small fraction of S38 which is derived from this polypeptide. The results are shown in Scheme I. The S38 sequence begins with residue 24 of brevin (22). For this sequencing we used a preparation that
Figure 1. Proteolysis of brevin. Panel a illustrates the digestion of human brevin using bacterial subtilisin at a 1:100 wt/wt ratio. The enzyme was added and the reaction stopped by addition of EGTA to 1 mM and DIFP to 5 mM at various times. The first lane contains marker proteins with known molecular weights; phosphorylase b, 94,000; BSA, 67,000; ovalbumin, 45,000; carbonic anhydrase, 30,000; soybean trypsin inhibitor, 21,000; and lysozyme, 14,300. The digestion times were 0, 5, 10, 15, 20, 30, 45, and 60 min in lanes 2–9, respectively. There is an accumulation of four peptides. Panel b illustrates a similar digestion using chymotrypsin at a 1:100 wt/wt ratio. From left-to-right the reaction times were 0, 1, 3, 5, 10, 30, 60, 90, and 120 min. At early times we see predominately two peptides with apparent molecular weights of 40,000 and 45,000. The 45,000 species gives rise to a third peptide at 43,000; the 40,000 species gives rise to one at about 38,000 that is progressively degraded to fragments of ~30,000–31,000 and 15,000. Panel c shows a 120-min digestion time point with chymotrypsin in the presence of EGTA. Compare with the last lane in b. Solution conditions: 10 mM Tris-HCl, 0.5 mM CaCl₂, 1 mM NaN₃, pH 8.0 for the digestions in a and b. The digestion in c was done in the same buffer plus 2 mM EGTA. The brevin concentration was 1 mg/ml in each digestion mixture; the temperature was 20°C.

was greater than 90% pure S38. Although we have not sequenced S40, we make the tentative conclusion that S38 is generated from S40 by cleavage at residue 24.

**Activity of Cleaved Products**

To estimate the activity of the fragments relative to intact brevin, we have compared the severing activity of brevin with an unfractionated mixture of polypeptides from a subtilisin digest equivalent to the 30-min time point shown in Fig. 1a, lane 7. The results are shown in Fig. 4a. Equal amounts of brevin were present in the digested and undigested samples. A 20-μl aliquot of pyrene-F-actin at 3 μM was diluted into 1,500 μl of 100 mM KCl, 3 mM MgCl₂, 0.5 mM ATP, 2 mM Tris-HCl, pH 8, to a final actin concentration of 40 nM. The EGTA samples contained 1 mM EGTA; the Ca²⁺ samples contained 0.2 mM CaCl₂. The brevin samples contained 1 nM undigested brevin; the digested samples contained an equivalent amount of digested brevin. In the presence of Ca²⁺, brevin markedly increases the rate of depolymerization (open triangles). This effect is attenuated 8-to-9-fold in EGTA (closed triangles) and closely resembles the control without added brevin (circles). The digested brevin retains a substantial severing activity in Ca²⁺ (open squares) and shows no modulation upon reducing the Ca²⁺ concentration (filled squares). It is difficult to quantitate the severing activity precisely, but the initial rate of depolymerization with digested brevin is approximately one-half that of the parent molecule. A similar result is seen for nucleation. Fig. 4b shows the results of initiating assembly of 5 μM actin (% pyrene-labeled actin, 95% unlabeled actin) in the presence of 50 nM brevin or an equivalent amount of digested brevin. The degree of brevin digestion was equivalent to that used in the severing experiment. Brevin nucleates rapid assembly in the presence of Ca²⁺ (open triangles) and, under these conditions, slows assembly in the presence of EGTA (closed triangles). Digested

**Scheme I.** Comparison of amino terminal sequence for brevin and S38.

Brevin          ALA-THR-ALA-SER-ARG-GLY-ALA-SER-GLN-ALA-
                 10
GLY-ALA-PRO-GLN-GLY-ARG-VAL-PRO-GLU-ALA-
XXX-PRO-ASN-
                 20

Brevin          XXX-MET-VAL-VAL-GLU-HIS-PRO-GLU-PHE-LEU-LYS-ALA-
S38            XXX-MET-VAL-VAL-GLU-XXX-PRO-GLU-PHE-XXX-LYS-ALA.

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Figure 2. Cross-reactivity of brevin fragments with conformation-specific monoclonal antibodies. Panels a and b show immunoblots stained using 8G5 IgG and 4F8 IgA, respectively. The gels are from lanes 3–7 of the chymotrypsin digest shown in Fig. 1b. Both antibodies recognize the 45,000- and 43,000-mol-wt fragments, but not the 40,000-mol-wt fragment, whose position is indicated by the open arrow. Lane 1 in c shows the Coomassie Blue profile of a brevin preparation with a 75,000-mol-wt peptide, U75, marked by the small arrow, that is produced by an unknown protease in our preparations. This peptide reacts strongly with 4F8 IgA in lane 2, c, but not with 8G5 IgG in the last lane. We infer that the 8G5 epitope is on a 15,000-20,000-mol-wt piece at one end of the brevin molecule. The larger arrowhead marks the position of the parent brevin molecule.

brevin shows some nucleation in both Ca²⁺ and EGTA with no apparent modulation. The initial rate of assembly of the samples with digested brevin was 10–12% of the control brevin rate in calcium. Both the digested and undigested molecules appear to cap the barbed filament ends as shown by a reduction in the final fluorescence values. The effective nucleation by the brevin digest is strongly dependent upon the concentration used. At low concentrations, less than 1 digested brevin per 100–200 actin molecules, we see a decrease in the rate of assembly compared with controls, similar to that observed with gelsolin and gelsolin–actin complexes (11). We interpret this as capping of some of the barbed ends with an apparent decrease in the overall rate of polymerization. Higher concentrations of the active fragment(s) increase the apparent rate of assembly above the control rate. The control polymerization curve in EGTA has been omitted for clarity, but lies between the brevin plus Ca²⁺ curve and the brevin digest values.

**Identification of the Major Severing Fragment**

The severing activity of the DEAE purified chymotryptic fragments is shown in Fig. 5. The open circles show the depolymerization of the control F-actin diluted to 40 nM. Intact brevin at 1.5 nM in the presence of Ca²⁺ increases the rate of depolymerization ~25-fold (open triangles), but has little effect in EGTA (data not shown). The 40-kD fragment, at 1.5 nM, increases the rate of depolymerization ~12-fold (open squares), about one-half the value for brevin. The activity of the purified 40-kD fragment shows no attenuation in 1–10 mM EGTA (filled squares). The 45/43-kD piece has no detectable effect at 15 nM, 10 times the concentration of the 40-kD fragment as shown by the closed circles that are equivalent to control values.

**Identification and Characterization of Actin Binding Fragments**

We have used three approaches to identify and characterize further the half of the molecule that binds actin: chromatography of digests on DNase I-Sepharose, reconstitution of fragments with actin followed by gel-filtration, and enhancement of NBD–actin fluorescence. Chromatography on DNase I was done by adding G-actin to the DIFP-inactivated digestion mixture at 2 mol of actin per mole of brevin, incubating for 30 min at 4°C, and chromatographing on DNase I-agarose as described by Wang and Bryan (20). The 45/43-kD fragments, from either a subtilisin or a chymotrypsin digest, are not retained at high or low ionic strength, in the presence or absence of calcium, or in the presence or absence of magnesium. An EGTA wash, which normally
The actin can be separated from the 40-kD fragment by the brevin molecule. After either quanidine or low pH treatment, the initial binding data for the 40-kD fragment gives a value of 2.4 actin molecules bound/40-kD fragment. The reasons for the somewhat higher value are unclear, but could be due to differences in dye binding or extinction coefficient for bovine brevin (10). The curves for both brevin and the 40-kD fragment are quite similar; the striking difference is that the 40-kD-actin interaction is not Ca\textsuperscript{2+} regulated. A similar extrapolation of the initial binding data for the 40-kD fragment gives a value of 2.4 actin molecules bound/40-kD fragment. The reasons for the somewhat higher value are unclear, but could be due to differences in dye binding or extinction coefficient for the whole versus part of the molecule.

Complex formation was also examined using HPLC gel filtration. The results for CT45/43 are summarized in Fig. 7a. We see no evidence for actin-45/43-kD fragment interaction.
The extrapolated value of 2.4 actin molecules per 40-kD fragment. The titration of actin-binding sites on brevin and the 40-kD fragment using NBD–actin. (a) Titration of 100 nM brevin with NBD–actin. Aliquots of NBD–actin in 2 mM Tris-HCl, 0.2 mM CaCl<sub>2</sub>, 0.5 mM ATP at pH 8.0 were added to 100 nM brevin in the same buffer plus 2 mM MgCl<sub>2</sub> (△) or to the buffer alone (△). The brevin samples show a fluorescence increase equivalent to that described previously for gelisolin (3). The difference binding curve is given by the open circles and shows saturation at ~1.8 actin molecules per brevin. For a different preparation of brevin this estimate has a range of 1.8–2.2 actin molecules/brevin. (b) Binding data for brevin and the 40-kD fragment are compared. The 40-kD fragment gives an extrapolated value of 2.4 actin molecules per 40-kD fragment. The data are normalized to percent of maximum binding given by fluorescence/maximum fluorescence for each sample and are plotted versus the number of actin molecules added per protein. ○, brevin plus 0.2 mM CaCl<sub>2</sub>; ●, are for brevin plus 2 mM EGTA. The corresponding 40-kD data are given (△ and △). The brevin plus EGTA data were calculated using the maximum fluorescence value obtained for brevin plus Ca<sup>2+</sup>. Sample additions were made and fluorescence measurements taken after 15–20 min at 20°C.

actions in the presence or absence of Ca<sup>2+</sup>, using either 0.1 M sodium phosphate, pH 7.0 or 0.15 M NaCl, 10 mM Tris-HCl, pH 7.5. Fig. 7b illustrates the results using CT40. We find complex formation in both Ca<sup>2+</sup> and EGTA and have estimated the stoichiometry of binding by estimating the Stokes radius of the complex by comparison of the elution time with elution times for proteins of known Stokes radii. The Stokes radius of the CT40–actin complex is that expected for a protein of ~125,000 mol wt. In four separate determinations

The fact that the actin binding domain retains a significant fraction of its severing, capping, and nucleating activities, but is unaffected by Ca<sup>2+</sup>, shows that Ca<sup>2+</sup> binding is not required to induce a brevin conformation that will interact with actin. The reverse appears to be the case, the 40-kD fragment appears to have constitutive severing and capping activity that is inhibited in the whole molecule at low Ca<sup>2+</sup> concentrations. Ca<sup>2+</sup> binding appears to induce a conformation change that makes these binding sites accessible to actin. Cleavage with subtilisin separates the actin binding domain from the inhibitory Ca<sup>2+</sup> regulatory domain and also makes these sites accessible.

Kwiatkowski et al. (12) have reported on the isolation of actin binding domains from brevin using chymotrypsin digestion. They have focused on CT17, a 17K fragment from the amino terminal end of the brevin molecule, that binds to G-actin–agarose and retains ~1% of the severing activity. We assume, but have not specifically demonstrated, that this is the 15–17-kD fragment that appears later in our chymotrypsin digests. We have isolated a similar 17-kD fragment from trypsin digests that retains severing activity and have used monoclonal antibodies to show this comes from the amino terminal of the 40-kD fragment. Kwiatkowski et al. (12) also describe a set of internal peptides, CT52 and CT47, that bind to G-actin in Ca<sup>2+</sup>, but not EGTA, and CT47 has no severing, capping, or nucleating activity. We see no similar major actin-binding fragments in our chymotrypsin digests and note that the 45-kD fragments from the carboxyl half of the molecule do not interact with actin.

In Fig. 8, we have diagrammed the organization of the various peptides. We have defined the order of the two Ca<sup>2+</sup> sites by the following observations. CT40 and S40 are located at the N-terminus and do not react with either antibody. S45/43 and CT45/43 contain both epitopes. The cleavage product, U75, from an unknown protease, in some of our brevin...
Figure 7. Formation of actin-brevin fragment complexes. G-actin at a final concentration of 0.2 mg/ml was mixed with either the 45-kD carboxyl terminal fragment at a final concentration of 0.2 mg/ml (a) or the 49-kD amino terminal fragment at a final concentration of 0.2 mg/ml (b) and incubated for 5 min at 20°C. 25-μl samples of the 2:1 actin/fragment mixtures were injected onto a Waters 1125 column in tandem with a Waters 300 SW column equilibrated with 0.1 M sodium phosphate at pH 7.0. The flow rate was 1.0 ml/min at 20°C. In each panel, curve a is from the G-actin standard. The peak at ~16.5 min is G-actin, the second peak at 22.5 min is from ATP. Curve b is for either the 45-kD fragment (panel a) or the 40-kD fragment (panel b). The c curves are for the mixtures. There is no shift in elution position for the actin + 45-kD mixture, but the actin + 40-kD mixture gives a more rapidly eluting complex with a larger Stokes radius and an apparent molecular weight of 125,000 when compared with standard proteins of known Stokes radius and molecular weight. Four determinations gave a range of 125,000–135,000-mol-wt. We see the same result with the 40-kD fragment + actin in the presence of 2 mM EGTA. The optical densities at 214 nm are arbitrary and have been scaled to allow the curves to be placed over one another. Similar results were obtained using 10 mM Tris-HCl, 150 mM NaCl, 0.2 mM CaCl₂, pH 7.5 as the equilibrating buffer with and without 2 mM EGTA. In this buffer, however, we find that brevin interacts to some degree with the resin in the separating columns.

Figure 8. Diagram of brevin functional domains. This is a schematic representation of the functional domains of the brevin molecule. The arrows indicate sites of protease sensitivity for chymotrypsin, subtilisin, and an unknown protease. Sequencing places the 40,000-mol-wt fragment with severing/capping activity at the amino terminal. We locate Ca site II, the exchangeable Ca²⁺ site, on the carboxyl terminal using the 8G5 IgG monoclonal antibody that binds to brevin and gelsolin or their corresponding actin complexes only if Ca²⁺ is present. Ca site I, the non-exchangeable site in gelsolin-actin and brevin-actin complexes, was located using the 4F8 IgA monoclonal antibody that binds to brevin-actin complexes even in EGTA. We have ordered actin sites I and II arbitrarily and propose that actin site I is located close to Ca site I. Furthermore, we infer that Ca site II and actin site II must be folded into close proximity if the carboxyl portion of the molecule is to inhibit severing in the absence of Ca²⁺.
We infer that Ca site II and actin site II must be in close proximity if the carboxyl part of the molecule is to inhibit actin–brevin interactions in the absence of Ca$^{2+}$. In support of this, we have preliminary evidence that the 75,000-mol-wt fragment, U75, missing only Ca site II, actively severs filaments in EGTA. Finally, we have some evidence that a 70,000–75,000-mol-wt trypsin fragment, missing only the amino terminal, will cap but not sever filaments.

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