Biophysical Journal

Article



The cooperative folding of annexin A2 relies on a transient nonnative intermediate

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ABSTRACT Annexins (Anxs) are a family of highly homologous proteins that bind and aggregate lipid vesicles in the presence of calcium. All members of the family contain a variable N-terminus determining specific functions, followed by a conserved core region responsible for the general calcium-dependent lipid-binding property. The core structure consists of four homologous domains $(D_I - D_{IV})$, each consisting of a right-handed super-helix of five α -helices. We present data from a combination of sitedirected mutagenesis, NMR, and circular dichroism showing that the G25-D34 region of the N-terminus as well as the contacts between residues D38A, R63A, and Q67A of AnxA2-D₁ are crucial for the autonomous folding and stability of D₁ of AnxA2. However, we also show that the folding of the full-length protein is very robust in that mutations and truncations that disrupted the folding of AnxA2-D₁ did not abolish the folding of full-length AnxA2, only lowering its thermal stability. This robustness of the folding of full-length AnxA2 is likely to be mediated by the existence of at least one transient nonnative intermediate as suggested by our kinetic data using stopped-flow fluorescence experiments. We also show that hydrophobic amino acids in AnxA2-D₁ involved in interfacial contacts with AnxA2-D_{IV} are important for the cooperative folding and stability of the full-length protein. Mutating all of the V57E-V98R-G101Y residues in AnxA2-D₁ did not affect the folding of the domain, only its stability, but prevented the cooperative folding of the full-length protein. Our collective results favor a highly cooperative and robust folding process mediated by alternative intermediate steps. Since AnxA2 is a multifunctional protein involved in several steps of the progression of cell transformation, these data on structure and folding pathways are therefore crucial to designing anticancer drugs targeting AnxA2.

SIGNIFICANCE Annexin A2 (AnxA2) is associated with tumorigenesis, neo-angiogenesis, metastasis, and chemoresistance. Since it is a multifunctional protein, function-targeted peptides/domains of AnxA2 corresponding to specific ligand-interaction sites can and have been used as biological competitors of endogenous AnxA2. To develop such competitors further, it is important to understand the folding process of AnxA2 and its isolated domains to generate more soluble forms of the protein. More generally, but as importantly, AnxA2, like other members of the annexin family, represents an interesting model to study the determinants governing cooperative folding of a purely α -helical protein. Thus, the identification of an obligatory transient intermediate in the folding pathway adds greatly to our understanding of the folding of AnxA2 and possibly other annexins.

INTRODUCTION

Annexins are defined as a family of α -helical proteins that interact with vesicles containing negatively charged phospholipids in a Ca²⁺-dependent manner (1). Annexin A2 (AnxA2)

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Editor: Doug Barrick.

https://doi.org/10.1016/j.bpj.2022.10.043

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is a multifunctional protein implicated in numerous cellular processes such as signal transduction, membrane trafficking, membrane/cytoskeleton interactions, and mRNA transport and translation (1–6). Its involvement in multiple different functions is regulated by posttranslational modifications (7). AnxA2 is associated with tumorigenesis, neo-angiogenesis, metastasis, and chemoresistance, processes all related to a poor prognosis (8). Inhibitors of angiogenesis have been associated with increased metastasis (9,10), so AnxA2 appears to be an ideal target for anticancer drugs, since the protein is Hollås et al.

involved in both processes (11). Initial attempts to employ isolated D_I or D_{IV} or peptides derived from these domains in in vitro assays resembling VEGF-induced angiogenesis were promising (12,13). In addition, a hexapeptide harboring the tissue plasminogen activator-binding site in the AnxA2 N-terminus competed successfully with endogenous AnxA2 for tissue plasminogen activator, thereby lowering the generation of plasmin, a protease important in tumor progression (14).

All annexins contain a conserved core region of about 300-310 residues harboring the Ca²⁺- and membrane-binding sites. The general structure of the annexin core consists of four domains (repeats) of about 75 residues each (denoted D_I to D_{IV}) that form a planar-cyclic arrangement (15,16). Each domain contains five α -helices (denoted A to E) arranged in a right-handed super-helix, in which helices A, B, D, and E form a bundle closed on one side by helix C lying perpendicular to the other helices. Within the bundle, helix pairs A-B and D-E are each arranged in an antiparallel mode. The Ca²⁺-binding sites are located in interhelical loops situated on the convex membrane-binding surface of the annexin core.

The N-terminal "tail" preceding the core region has a highly variable sequence and confers specific properties to each particular annexin (1). In the obtained wild-type AnxA2 structures (1,13,17–19), the first 20 residues could never be observed. The maximal observable portion of the tail spanned residues P21 to F33 (the next residue, D34, is the capping residue for helix A of AnxA2-D_I). The extreme N-terminus of the visible part of the tail (residues 21–26) interacts with D_{IV} and runs in the direction of D_I. This conserved region in most annexins including AnxA2 was found to function as a velcro-like bridging segment in AnxA11 stabilizing the core structure (20). V27 (counting M as residue 1) and S26 both insert in between D_I and D_{IV} of AnxA2 (19).

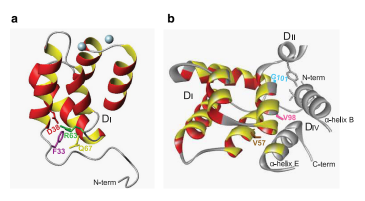
Despite their overall structural similarity, each of the four domains of the annexin core displays some specific structural features. Accordingly, sequence divergence between the four domains within any given annexin is higher than that observed for a given domain between different annexin cores (21). In particular, the interfacial contacts between the domains are not evenly distributed. The domains D_I and D_{II} are engaged in tight hydrophobic contacts with D_{IV} and D_{III}, respectively, resulting in two modules, D_I-D_{IV} and D_{II}-D_{III}, which interact with each other via weaker, largely electrostatic interactions (Figs. 1 and S1) (16). There are also variations in the ability of the annexin domains to acquire a native tertiary structure when expressed separately from the rest of the protein. It has been suggested that the folding process of the domains of full-length AnxA1 follows an ordered and sequential series of events (22,23). In the proposed model, D_I folds first, whereas D_{II} and D_{III} remain partly unfolded until D_{IV} docks on D_I, thereby initiating the complete folding of the protein (22–24). In this model, the establishment of the hydrophobic interface between already folded D_I and D_{IV} is a prerequisite for the completion of the folding of $D_{\rm II}$ and $D_{\rm III}.$ In AnxA2, $D_{\rm IV}$ could be solubilized only at the expense of mutating several hydrophobic amino acids involved in interfacial contacts with D_I (25). However, the resulting construct was only partly folded, indicating the importance of the hydrophobic contacts between D_I and D_{IV} in stabilizing the folding of D_{IV}.

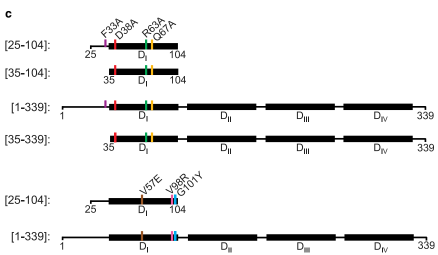
In the present work, we decided to investigate whether, as proposed for AnxA1 (22,23), AnxA2-D_I forms an autonomous folding unit that subsequently induces the cooperative folding of the full-length AnxA2. To this aim, we produced wild-type and various mutated constructs of the AnxA2-D_I as well as of full-length AnxA2. The solubility, the temperature-dependent stability, and the conformational status of the different constructs were studied by circular dichroism (CD) and proton-nitrogen heteronuclear single quantum correlation nuclear magnetic resonance (¹H-¹⁵N HSQC NMR) spectroscopies. Furthermore, we provide a thorough characterization of the folding and unfolding kinetics of full-length AnxA2 in comparison to the mutant proteins. We observed that several mutations/truncations that disrupted the autonomous folding of AnxA2-D_I did not disrupt the capacity of the full-length AnxA2 to fold. Conversely, some mutations that preserved the autonomous folding of AnxA2-D_I strongly disrupted the thermostability of full-length AnxA2. However, these mutations at the AnxA2 D_I-D_{IV} interface render the full-length protein less stable, which does not mean that the folding pathway is affected. Therefore, we propose a new model of annexin folding, in which the autonomous folding of AnxA2-D_I is not a precondition for the folding of the full-length AnxA2 protein. Of additional interest, we provide evidence for the presence of a previously undetected obligatory transient intermediate in the folding of AnxA2. Analysis of the effect of the mutations on the stability of the intermediate versus that of the native state suggests this intermediate to be misfolded, with residues of AnxA2-D_I involved in nonnative contacts. In fact, we observed that most mutations have a more pronounced effect on the stability of the intermediate compared with that of the native state. Thus, the results presented suggest the folding of AnxA2 to be a cooperative process, rather than a stepwise reaction, undergoing transient misfolding events, possibly involving AnxA2-D_I. This does not preclude the possibility that AnxA2-D_I must be folded during the folding events of full-length AnxA2.

MATERIALS AND METHODS

Construction of wild-type and mutants of fulllength AnxA2, truncated Δ 1-34AnxA2, and AnxA2-D_I

The bovine anxA2 cDNA (RefSeq: NM_174716.1) for the coding region in the pGEM-3Zf(+) vector (Promega Corporation, Madison, USA) was used as the template for the construction of wild-type (wt) and mutant full-length AnxA2, truncated $\Delta 1$ -34AnxA2, and AnxA2-D_I (for details, see Fig. 2). This was done by BbsI seamless cloning (26) using TA cloning of the





cDNA fragments into the TOPO vector (Thermo Fisher Scientific, Waltham, USA) as an intermediate step or by QuickChange Site-Directed Mutagenesis (Agilent Technologies, Santa Clara, USA). The full-length and the $\Delta 1$ -34 truncated AnxA2 forms were expressed with a 6His tag from the pETM10 vector, whereas the wild-type and mutant forms of AnxA2-D_I were expressed from the pETM41 vector fused to 6HisMBP, and subsequently separated by Tobacco Etch Virus (TEV) cleavage of a TEV-protease sensitive linker region (ENLYFQG). Both vectors were kindly provided by Gunter Stier (University of Heidelberg, Germany). Oligonucleotide primers, 38-59 oligomers, were obtained from Sigma-Aldrich (Saint-Louis, USA). The final constructs were verified by DNA sequencing at the Haukeland University Hospital.

Protein expression and purification

The expression and purification of the AnxA2 wild-type and mutants of full-length, $\Delta 1-34$ truncated and D_I forms were done as described previously (25), with some modifications. The sonicated homogenate was centrifuged in 2-mL tubes at 16,000 gav for 30 min at 4°C, and the supernatant was filtrated through a sterile 0.2-μ-pore-size filter (Merck Millipore, Burlington, USA) before loading onto a column (Bio-Rad Laboratories, Hercules, USA) with equilibrated His-Select Nickel Affinity Gel (Qiagen, Hilden, Germany). The resin and the supernatant were mixed and incubated for 15 min, until the resin settled in the column and the supernatant was eluted by the force of gravity. The flow-through was again mixed with the resin before elution of protein. Subsequently, the 6HisMBP- AnxA2-D_I fusion FIGURE 1 (A) Ribbon plot of domain I (D_I) of AnxA2 (from Pro21) showing the four conserved residues that appear to constitute a structural tetrad at the N-terminus of D_I. The four conserved residues are F33, D38, R63, and Q67, counting the first M as residue 1. (B) Ribbon plot of D_I of AnxA2 (from P21) indicating the side chains of the three mutated hydrophobic residues (V57, V98, and G101) at the interface of D_I with α -helices in D_{II} and D_{IV} of AnxA2. The figures were prepared using Biovia Studio Visualizer (http://accelrys.com/ products/collaborative-science/biovia-discoverystudio/visualization.html) and based on the PDB structures 1XJL and 4X9P. (C) Schematic representation of wild-type and different mutated/truncated forms of AnxA2-D_I and full-length AnxA2. To investigate the role of a folded AnxA2-D_I in the folding of the full-length AnxA2, single mutations (F33A, D38A, R63A, and Q67A) were introduced in the context of [25-104] and [35-104] (not F33A) AnxA2-D_I as well as in [1-339] full-length AnxA2 and [35-339] Δ 1-34-truncated AnxA2 (not F33A). Furthermore, a multiple mutation (F33A-D38A-R63A-Q67A) was introduced in [1-339] full-length AnxA2. Subsequently, to investigate the role of interfacial hydrophobic amino acid residues in AnxA2-D_I for the folding of fulllength AnxA2, single mutations (V57E, V98R, and G101Y) and a multiple mutation (V57E-V98R-G101Y) were introduced in the context of [25-104] AnxA2-D_I and [1-339] full-length AnxA2 as indicated in the figure. The numbers represent the number of the particular amino acid in the bovine and human AnxA2 sequence, and the four domains are shown as black boxes, denoted D_I-D_{IV}. The color code for the respective mutated amino acids as indicated in (C) is also used in (A) and (B).

proteins were cut by TEV and purified again as described (25,27). The TEV cleavage site results in four additional residues (GAMG) at the N-terminus of the peptide constructs.

Proteins used in NMR experiments were expressed in M9 ¹⁵N minimal medium (4.3 mM Na₂HPO₄ x 2H₂O, 22 mM KH₂PO₄, 8.6 mM NaCl, 9.2 mM ¹⁵NH₄Cl, 30 μM FeCl₃ x 6H₂O, 134 μM EDTA, 6.2 μM ZnCl₂, 1 μ M CuCl₂, 0.4 μ M CoCl₂ x 6H₂O, 1.6 μ M H₃BO₃, 68.6 μ M MnCl₂ x 2H₂O, 0.4% glucose, 1 mM MgSO₄, 0.3 mM CaCl₂, 1 μg/mL biotin, 1 μg/mL thiamine, and 35 μg/mL kanamycin). All reagents were from Sigma-Aldrich (Saint-Louis, USA). The labeled proteins were finally eluted with 20 mM Tris (pH 7).

Solubility assays

Cultures (25 mL) were expressed at 37°C for 3 h. 1 mL of the expressed cultures was pelleted and resuspended in 500 µL breakage buffer (50 mM Na₂HPO₄ (pH 8.0), 0.5 M NaCl, 10 mM imidazole, 5% (w/v) glycerol, 0.5 μg/mL DNase I, 0.25 μg/mL RNase A, protease inhibitor cocktail (1 tablet/50 mL) (Complete, EDTA-free, Roche Diagnostics, Basel, Switzerland) and 1 mM DTT). All reagents were from Sigma-Aldrich (Saint-Louis, USA) if not otherwise stated. The bacteria were disrupted by sonication cycles at 4°C (4 x 20 pulses) with a thin probe. Sonicated cells (T) (100 μL) were centrifuged (16,000x g for 15 min) to obtain a supernatant (S) and a pellet (P) fraction. Proteins in equal volumes of total extract (T), supernatant fraction (S), and the pellet (P) (resuspended in 100 μL breakage buffer) were separated by 10% SDS-PAGE.

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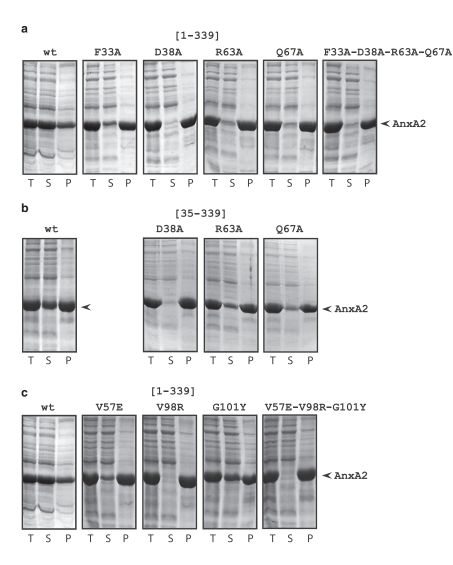


FIGURE 2 Characterization of the solubility of wild-type and mutated (A) [1-339] (wild-type, F33A, D38A, R63A, Q67A, and F33A-D38A-R63A-Q67A), (B) [35-339] (wild-type, D38A, R63A, Q67A), and (C) [1–339] (wild-type, V57E, V98R, G101Y, V57-V98R-G101Y) AnxA2 forms as determined by 10% SDS-PAGE analysis. The 6His-tagged AnxA2 forms were expressed for 3 h at 37°C after induction with 1 mM IPTG. Subsequently, aliquots were retrieved and sonicated in breakage buffer (total, T). After centrifugation at 16,000 g for 15 min, aggregated proteins were recovered as the pellet (P) fraction and the soluble proteins as the supernatant (S) fraction. The protein bands were visualized by Coomassie Brilliant Blue staining. AnxA2 is indicated by an arrowhead to the right.

CD measurements

Thermal transition curves $(25^{\circ}\text{C}-75^{\circ}\text{C}, 30^{\circ}\text{C}-80^{\circ}\text{C}, \text{or }25^{\circ}\text{C}-95^{\circ}\text{C}$ temperature ranges) were determined by monitoring the decrease in ellipticity at 222 nm at a scanning rate of 100°C/h using a 1-mm pathlength cell in a Jasco J-810 spectropolarimeter equipped with a Peltier temperature control unit. The proteins were diluted to $15-40~\mu\text{M}$ in 20 mM Tris (pH 7.0). Since we purified several constructs at the same time to standardize conditions as much as possible, it was important to determine an apparent T_m as soon as possible after purification. Therefore, the scanning rate was 100°C/h for all constructs, so the profiles and apparent T_m s were comparable, providing data sufficiently informative for drawing qualitative conclusions about the folding status and the thermostability of the constructs. All samples were freshly prepared, and the CD measurements were performed with single runs on the same day to obtain the relative differences between the apparent T_m values.

NMR spectroscopy

The NMR spectra were acquired using sample concentrations of 150–700 μ M at 25°C and 700 MHz (1 H frequency) on a Bruker DRX700 spectrometer equipped with a z-gradient triple resonance cryoprobe, in 20 mM Tris (pH 7.0). 1 H- 15 N HSQC were recorded with spectral widths set to avoid

folding of ¹⁵N resonances. In all spectra, the water signal was suppressed using a WATERGATE sequence (28). Typically, 256 t₁ increments were acquired and quadrature was achieved (29). Spectra were processed and analyzed with TopSpin Bruker Software Package Version 2.1 (RRID: SCR_014227).

Stopped-flow measurements and analysis of kinetic data

The kinetics of unfolding and folding of AnxA2 were detected by monitoring the fluorescence emission after excitation at 280 nm, using a cutoff filter at 320 nm on a stopped-flow instrument (Applied Photophysics, Leatherhead, UK). For each experiment, an average calculated from three to five independent data acquisitions was fitted to the following equation:

$$y = A \cdot exp \left(-k_{obs} \cdot t \right) + c, \tag{1}$$

which allows to calculate the observed rate constant ($k_{\rm obs}$) at different denaturant concentrations for each variant analyzed (Eq. 1). The dependence of the $k_{\rm obs}$ as a function of denaturant concentration was fitted using an equation implying a three-state scenario, with the presence of an intermediate accumulating in the unfolding reaction:

$$k_{\text{obs}} = k_f^0 \exp\left(-m_f [UREA]/RT\right) + \frac{k_u^0 \exp(m_u [UREA]/RT)}{1 + K_{N-I} \exp(m_{N-I} [UREA]/RT)}, \quad (2)$$

where k_f^0 and k_u^0 represent the folding and unfolding microscopic rate constant in the absence of denaturant, and K_{N-I} is the equilibrium rate constant between the native and intermediate state (Eq. 2) (30).

The difference in free energy between the denatured and native state for each mutant analyzed was calculated as

$$\Delta G_{D-N} = RT \ln \left(k_f^0 / k_u^0 \right) \tag{3}$$

$$\Delta G_{D-I} = \Delta G_{D-N} - \Delta G_{N-I} \tag{4}$$

and

$$\Delta \Delta G_{D-N} = \Delta G_{D-N}^{wt} - \Delta G_{D-N}^{mut}$$
 (5)

$$\Delta \Delta G_{D-I} = \Delta G_{D-I}^{wt} - \Delta G_{D-I}^{mut}$$
 (6)

The beta Tanford (β_T) value for the intermediate state was calculated as

$$m_{D-N} = m_f + m_u \tag{7}$$

$$\beta_T = 1 - (m_{I-N} / m_{D-N}) \tag{8}$$

Accession numbers

Bovine AnxA2 was used in these experiments; UniProt: P04272, GenBank: BT021010.1, MMDB: 126161, and PDB 1: 4X9P.

RESULTS

Design of mutated AnxA2-D_I, Δ 1-34AnxA2, and full-length AnxA2 forms

In a former study, we showed that calculation of the cumulated surface hydrophobicity of each isolated AnxA2 domain indicated that AnxA2-D_I is the most hydrophilic domain (25). In the AnxA2 crystal structure (PDBs: 1XJL (18) and 4X9P (13)), we observed that four highly conserved amino acids (F33, D38, R63, and Q67; counting the first M as residue 1) appear to form a "structural tetrad," which stands on the periphery of AnxA2-D_I, yet establishes a network of specific mutual contacts (Fig. 1 A). The side chain of F33 is shielded from the solvent by the side chains of D38 and R63, which extend toward each other due to a favorable charge interaction. In addition, the aromatic ring of F33 establishes an antiparallel interaction with the side chain of Q67. The residue Q67, in turn, establishes, via its side chain, specific contacts that connect the N-terminal tail to AnxA2-D_I. The side chain carboxamide oxygen of Q67 makes a bidentate H-bond, one to the main-chain amino group of residue Y30 that belongs to the N-terminal tail, and the other one to the main-chain amino group of S64, a residue situated at the connection between helix B and helix C of AnxA2-D_I. Based on these observations, we hypothesized that the network of contacts established by the F33/D38/R63/Q67 tetrad could be important for stabilizing the folding of AnxA2-D_I and thus of full-length AnxA2. The four positions were therefore mutated into alanine residues, either individually (mutations F33A, D38A, R63A, Q67A) or simultaneously (mutation F33A-D38A-R63A-Q67A) (Fig. 1 C). These mutations were introduced in the context of a construct of D_I, corresponding to residues [25– 104] of AnxA2 encompassing the five helices of D_I preceded by 10 residues of the N-terminal tail (Fig. 1 C), exactly like in the previously published AnxA1-D_I construct (22,23). The same mutations were also performed in the context of fulllength AnxA2 (construct [1–339]).

The C-terminal part of the N-terminal tail that is visible in the crystal structures (residues 25-34) is involved in several contacts with D_I. Thus, a deletion of the N-terminal tail (residues 1–34) would be expected to disrupt the capacity of D_I to fold cooperatively. Such a deletion was therefore performed both in the context of D_I (construct [35–104]) and in the context of full-length protein (construct [35-339]). We also generated constructs that combined the N-terminal deletion G25-D34 with the single-point mutations D38A, R63A, and Q67A, both in the context of isolated D_I and of the full-length protein.

AnxA2-D_I is engaged in strong contacts with other domains within the annexin core. To identify the amino acid residues that are involved in the interaction of AnxA2-D_I with the rest of the protein, we examined the crystal structures of AnxA2 (PDBs: 1XJL and 4X9P) and found that its helices B and E are involved in the interfacial contact with the rest of AnxA2 (Fig. 1 B). Key residues at the interface are V57, H94, L95, V98, G101, L102, and K104. However, in the isolated AnxA2-D_I, L95 and L102 appear to be too buried to be mutated, since it is likely that mutation of these residues will affect the folding of AnxA2-D_I. H94 and K104 are not conserved in AnxA2 (Fig. 1 B in Aukrust et al. (25)). By contrast, V57 (at the interface between D_I and D_{IV}), V98 (interface between D_I and D_{IV}), and G101 (interface between D_I and D_{II} and D_{IV}) are quite conserved and exposed in the isolated AnxA2-D_I. We therefore mutated these three positions to polar residues (V57E and V98R) and bulky, yet polar residue (G101Y) in the context of [25–104] AnxA2-D_I and full-length AnxA2 (Fig. 1 C).

Cloning, expression, and purification of the wildtype and mutated AnxA2-D_I, Δ 1-34-AnxA2, and full-length AnxA2 forms

Constructs of AnxA2-D_I were fused via a TEV-proteasesensitive linker to the C-terminus of the highly soluble bacterial maltose binding protein. Constructs of full-length AnxA2 were produced with an N-terminal 6His tag.

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The His-tagged constructs of point-mutated and N-terminally deleted full-length AnxA2 all showed decreased solubility upon bacterial overexpression at 37°C, as compared with the wild-type full-length AnxA2 (Fig. 2). Although about 60%–70% of the wild-type construct [1–339] was retrieved in the soluble fraction of the sonicated lysate, no more than 10% of the other constructs was retrieved in the soluble fraction, except for construct AnxA2 [1–339] G101Y, which was almost as soluble as the wild-type construct. These differences in solubility were no more apparent when the expression was performed at 15°C (data not shown). In this condition, the amount of overexpressed protein retrieved in the soluble fraction was comparable for all mutants and for the wild-type AnxA2. The proteins were therefore all expressed at 15°C for further purification.

Lowering the expression temperature from 37°C to 15°C is expected to decrease 1) the rate of ribosomal peptide synthesis, 2) the concentration of newly expressed AnxA2 polypeptide chains exploring their folding at any given time point, and 3) the diffusion rate of these polypeptide chains undergoing folding. The decreased efficiency of folding of AnxA2 mutants at 37°C as compared with 15°C suggests that the corresponding mutations (or deletion) resulted in misfolded intermediates with high propensity to self-associate. Being more concentrated and prone to faster diffusion at 37°C, these misfolded intermediates form a higher proportion of insoluble aggregates at 37°C, whereas at 15°C, they have an increased chance to explore the folding and reach the correctly folded final state.

Role of the N-terminal tail and of the structural tetrad F33-D38-R63-Q67 in the folding and stability of AnxA2-D₁

To explore the role of the "structural tetrad" formed by F33, D38, R63, and Q67, the thermal stability and the

folding status of the purified protein constructs were analyzed by thermal denaturation CD measurements and $^1\text{H-}^{15}\text{N}$ HSQC NMR.

Thermal denaturation was assessed by following the change in ellipticity at 222 nm. The 222-nm signal is mainly contributed by α -helical secondary structure, yet other structures including random coil can also contribute some signal. Moreover, the presence of α -helical secondary structure does not warrant that the domain has acquired a folded tertiary structure. For some constructs, we could only obtain low-concentrated samples, so that the spectral quality was sometimes suboptimal (see Fig. S2), and therefore subsequent determination of the apparent transition temperature $(T_{\rm m})$ was error-prone (see Table 1). However, even in such cases, the profiles were sufficiently informative for drawing qualitative conclusions about the folding status and the thermostability of the domain.

The folded tertiary structure can be tracked by observing a wide distribution of the amide ¹H frequencies in the ¹H-¹⁵N HSQC NMR (Fig. 3), as well as by observing a cooperative transition in the temperature-induced decrease of the absolute value of the negative ellipticity signal in the CD instrument (Fig. S2).

According to these approaches, the isolated wild-type $AnxA2-D_I$ (construct [25–104] wild-type) was fully folded at 25°C (Fig. 3, upper left panel), in agreement with published data on a corresponding construct of $AnxA1-D_I$ (22,23). The thermal transition of wild-type $AnxA2-D_I$ followed a broad temperature range with an apparent transition temperature (T_m) of \sim 65°C (Table 1) as determined from the first derivative of the transition curve (Fig. S2). Among the mutants of isolated $AnxA2-D_I$ constructs, [25–104] F33A and [25–104] R63A were also folded (Fig. 3, upper central and right panels), albeit with a lower T_m (Table 1; Fig. S2), indicating a loss of thermodynamic stability, particularly in the case of the R63A mutant. By contrast,

TABLE 1 Estimated T_ms of [25–104], [35–104], [1–339], and [35–339] recombinant wild-type or mutated AnxA2 forms

AnxA2-D _I		Full-length/truncated AnxA2		
Construct	Estimated T _m	Construct	Estimated T _m	
[25–104] wild-type	65°C	[1–339] wild-type	58°C	
[25–104] F33A	63°C	[1-339] F33A	56°C	
[25–104] D38A	Not folded	[1-339] D38A	51°C	
[25-104] R63A	55°C	[1-339] R63A	54°C	
[25–104] Q67A	Not Folded	[1–339] Q67A	53°C	
		[1-399] F33A-D38D-R63A-Q67A	49°C	
[35–104] wild-type	Not folded	[35–339] wild-type	47°C	
[35–104] D38A	Not folded	[35–339] D38A	41°C	
[35–104] R63A	Not folded	[35–339] R63A	47°C	
[35–104] Q67A	Not folded	[35–339] Q67A	45°C	
[25–104] V57E	58°C	[1-339] V57E	52°C	
[25–104] V98R	64°C	[1–339] V98R	41°C	
[25-104] G101Y	56°C	[1–339] G101Y	50°C	
[25–104] V57E-V98R-G101Y	62°C	[1–339] V57E-V98R-G101Y	Not folded	

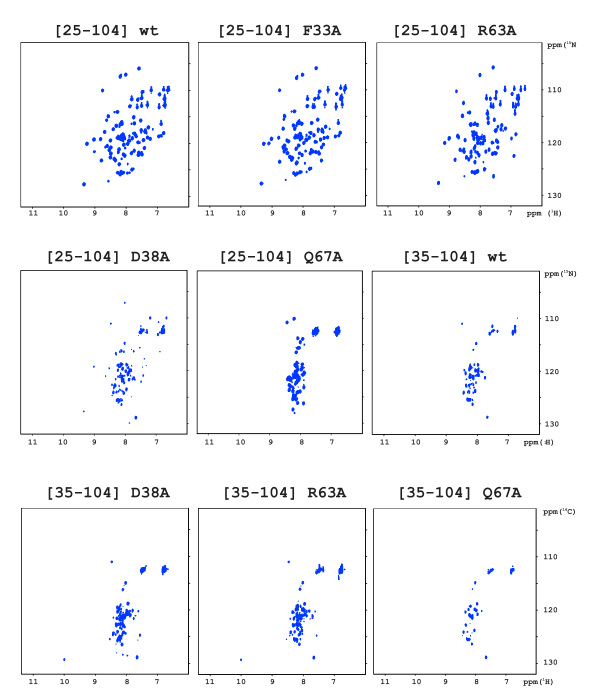


FIGURE 3 Impact of mutating amino acid residues (F33, D38A, R63A, Q67A) or deleting the N-terminus on the folding of AnxA2-D_L. ¹H-¹⁵N HSQC NMR spectra of [25-104] AnxA2; folded wild-type (600 µM), F33A (550 µM) and R63A (150 µM) (upper row), and misfolded D38A (275 µM), Q67A (275 µM) as well as misfolded [35-104] AnxA2; wild-type (50 µM), D38A (700 µM), R63A (700 µM), and Q67A (700 µM), were recorded at pH 7, 25°C, and 700 MHz (¹H frequency). To see this figure in color, go online.

the other mutants of isolated AnxA2-D_I, [25–104] D38A and [25-104] Q67A, were not folded (Fig. 3, middle-left and central panels; Fig. S2). This was corroborated by data from ellipticity scans at 25°C, which showed a high contribution of random coil and thus low helical structure (results not shown). Furthermore, all the wildtype and mutated [35-104] AnxA2-D_I constructs, corresponding to the strict five-helix AnxA2-D_I without any N-terminal tail, were also not folded at 25°C (Fig. 3, middle-right panel and all lower panels). Altogether, these results point to the importance of the G25-D34 region of the N-terminal tail, as well as of the structural tetrad F33-D38-R63-Q67 in the autonomous folding of the isolated AnxA2-D_I.

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Role of the N-terminal tail and of the structural tetrad F33-D38-R63-Q67 in the folding and stability of full-length AnxA2

Next, we introduced the same mutations, namely F33A, D38A, R63A, Q67A, and Δ 1-34, separately or combined, in the context of full-length AnxA2. The thermal unfolding of wild-type full-length AnxA2 (Fig. S2) showed a sharp transition ($T_{\rm m}=55.4^{\circ}{\rm C}$) over a range of less than $10^{\circ}{\rm C}$, characteristic of a strong cooperative transition from a specific native tertiary structure to an unfolded state. Interestingly, all our constructs of AnxA2 bearing the abovementioned mutations were still able to fold according to their ¹H-¹⁵N HSQC NMR spectra acquired at 25°C (Fig. 4). They also retained the highly cooperative thermal unfolding behavior of wild-type AnxA2 (Fig. S2), albeit with variable decrease in thermodynamic stability, illustrated by lower melting temperatures (T_m) (Table 1). Therefore, although mutations $\Delta 1$ -34, D38A, and Q67A abolished the folding capacity of isolated AnxA2-D_I, they did not prevent full-length AnxA2 to acquire a folded conformation, demonstrating that the folding of the core four-domain structure can occur in the presence of N-terminal deletions or mutations that prevent the folding of the isolated D_I domain. It remains possible that D_I – D_{IV} docking and subsequent formation of a rigidified D_I – D_{IV} module would be the first step of AnxA2 folding, as proposed earlier in the sequential folding model (22,24).

Nonetheless, although the full-length F33A and R63A mutants only have slightly lowered T_m s, the full-length D38A and in particular the multiple full-length AnxA2 mutant (F33A-D38A-R63A-Q67A) show decreased thermodynamic stability (Table 1; Fig. S2). This is in agreement with the decreased solubility, upon bacterial overexpression at 37°C, of the mutated full-length AnxA2 constructs compared with the wild-type full-length AnxA2 (Fig. 2). This suggests that the full-length AnxA2 mutants, which harbor folding-defective $D_{\rm I}$, display slower kinetics of folding than the wild type.

The importance of hydrophobic interfacial amino acids of D_I in the folding of full-length AnxA2

Subsequently, we focused on several amino acids located in AnxA2-D_I, which were targeted to disrupt the folding of full-length AnxA2, but not of the isolated AnxA2-D_I. Thus, a new series of mutations, this time targeting

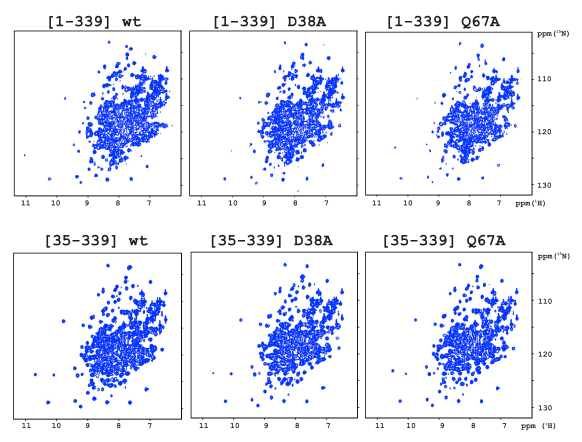


FIGURE 4 Impact of mutating amino acid residues (D38A, Q67A), causing collapse of the folding of AnxA2-D_I, and of deleting the N-terminus on the folding of full-length AnxA2. 1 H- 15 N HSQC NMR spectra of [1–339] AnxA2; wild-type D38A and Q67A as well as [35–339] AnxA2, all at 200 μ M were recorded at pH 7, 30°C, and 700 MHz (1 H frequency). To see this figure in color, go online.

hydrophobic amino acids involved in interfacial contacts with other domains of AnxA2, was performed: three single-point mutants V57E, V98R, and G101Y and the triple mutant V57E-V98R-G101Y in the full-length protein. Neither the single-point mutations nor the combined triple mutation altered the capacity of the isolated AnxA2-D_I domain to acquire its folded tertiary structure (Fig. 5). However, all mutations affected the thermodynamic stability of the isolated domain to various degrees (Table 1; Fig. S2). When the same interfacial mutations were introduced into full-length AnxA2, all mutants could be purified in a soluble form and retained, to some extent, a cooperative thermal unfolding process (Fig. S2). However, the unfolding profile of the multiple mutant (V57E-V98R-G101Y) of full-length AnxA2 was seriously altered (Fig. S2), suggesting a less cooperative process than that observed for the wild-type protein. Furthermore, the T_m s of the three single mutants of full-length AnxA2 decreased significantly. The multiple mutant (V57E-V98R-G101Y) of full-length AnxA2 was also found to precipitate when prepared for ¹H-¹⁵N HSQC NMR spectroscopy measurements, so that no spectrum could be obtained. Altogether, these data point to the contribution of the interfacial residues V57, V98, and G101 of AnxA2-D_I, mainly interacting with interfacial residues in AnxA2-D_{IV}, to the stability of the full-length AnxA2 protein.

The folding and unfolding kinetics of AnxA2

Prompted by the detailed characterization of the unfolding stability of AnxA2 and its site-directed mutants reported above, we resorted to analyze quantitatively their folding and unfolding kinetics by stopped-flow fluorescence. In particular, we carried out 11-fold dilution experiments in which 1) the native protein diluted in buffer (25 mM Tris-HCl, 150 mM NaCl, 2 mM EDTA, 2 mM Tris(2-carboxyethyl)phosphine (TCEP), pH 8.0) was rapidly mixed with guanidinium-HCl (GdnHCl) at different concentrations (unfolding), and 2) the protein in denaturing conditions (buffer with added GdnHCl) was mixed with refolding buffer to reach different final concentrations of GdnHCl. Samples were excited at 280 nm, and the fluorescence emission was collected using a 320-nm cutoff filter. Typical folding and unfolding time courses are reported in Fig. 6. Interestingly, although in both cases the apparent decay appears consistent with a single exponential function, both folding and unfolding correspond to a decrease of fluorescence versus time, suggesting the rapid accumulation of a burst phase intermediate in the dead-time of the stopped-flow in either folding or unfolding (31,32).

A semilogarithmic plot of the observed (un)folding rate constants (chevron plot) for wild-type AnxA2 is shown in Fig. 7. A visual inspection of the observed dependence might appear consistent with a V-shape chevron plot, a typical signature of two-state folding (33). Fitting the observed rate constant to a simple two-state equation, however, indicates a folding m value of 2.2 ± 0.1 kcal $\text{mol}^{-1} \text{ M}^{-1}$ and an unfolding m value of 0.9 \pm 0.05 kcal $\text{mol}^{-1} \text{ M}^{-1}$, with a total m value of 3.1 \pm 0.1 kcal mol^{-1} M^{-1} . The m value, defined as $\delta \Delta G/\delta$ [denaturant], represents a quantitative measure of the cooperativity of the (un) folding reaction, and it reflects the change in accessible surface area upon denaturation (34). The m value calculated from the two-state analysis of wild-type AnxA2 appears to be much lower than that expected from a protein of 339 amino acids, which should be around 6 kcal mol⁻¹ M⁻¹ (34). This result suggests the presence of additional undetected states along the reaction pathway that influence the calculation of the structural transition occurring between native and denatured state. Therefore, the unexpected lower m value suggests the presence of (at least) one intermediate.

Keeping in line with the stability experiments reported above, we subjected the mutants reported in Table 2 to kinetic experiments. The chevron plot of all the mutants in comparison to that of wild-type AnxA2 is reported in Fig. 8. A pronounced curvature of the unfolding limb of the chevron plot is clearly observed in all cases, strongly indicating the presence of an intermediate. Therefore, the stopped-flow experiments clearly point to the accumulation of a reaction intermediate in the folding of AnxA2, for the wild-type protein as well as all for its mutants. Quantitative analysis of the chevron plots presented in Fig. 8, by fitting kinetic data with a three-state model (Eq. 2), allows to calculate the m_f, m_u, and m_{I-N} parameters, which represent the

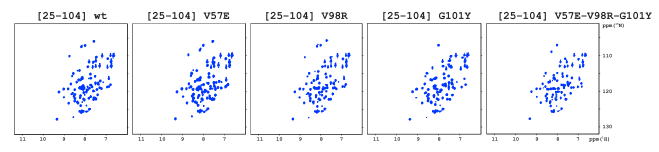


FIGURE 5 Mutating amino acid residues (V57E, V98R, G101Y, V57E-V98R-G101Y) in AnxA2-D₁ involved in interfacial contact with other domains does not unfold this domain. $^{1}\text{H-}^{15}\text{N}$ HSQC NMR spectra of [25–104] AnxA2; wild-type (600 μM), V57E (250 μM), V98R (250 μM), G101Y (250 μM), and V57E-V98R-G101Y (150 μM) were recorded at pH 7, 25°C, and 700 MHz (¹H frequency). To see this figure in color, go online.

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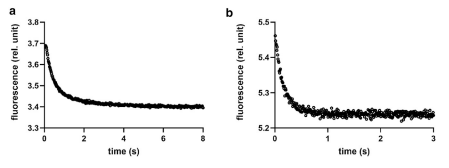


FIGURE 6 Typical unfolding and refolding time courses of AnxA2. (*A*) AnxA2 diluted in buffer (25 mM Tris-HCl, 150 mM NaCl, 2 mM EDTA, 2 mM Tris(2-carboxyethyl)phosphine (TCEP), pH 8.0) was rapidly mixed with GdnHCl to reach 5.4 M final concentration (unfolding); (*B*) the protein in denaturing conditions (buffer with added GdnHCl) was mixed with refolding buffer to reach 0.5 M final concentration of GdnHCl.

dependence of folding and unfolding microscopic rate constants and equilibrium rate constant between the intermediate and the native state, from denaturant concentration. The $m_{\rm I-N}$ value calculated for the intermediate is 2.4 \pm 0.3 kcal $\text{mol}^{-1} \text{ M}^{-1}$, with a total $m_{\text{D-N}}$ value of 5.9 \pm 0.2 kcal mol^{-1} M⁻¹. As mentioned above, these kinetic parameters correlated to the change in accessible surface area upon unfolding and allow to pinpoint the position of the intermediate state along the reaction coordinate. The calculation of the so called Tanford β value (β_T) for the intermediate state, as formalized by Eq. 8, reports a value of $\beta_T = 0.59 \pm 0.06$. This value, typically lying between 0 (for the denatured state) and 1 (for native state), suggests the intermediate to be populated as a relatively late event in the reaction, implying a possible involvement of more than one of the four domains of AnxA2. Importantly, the observed robustness of kinetic m values and consequently of β_{T} values clearly points to the evidence that, although destabilized, all variants explored appear to conserve a similar folding mechanism.

A quantitative analysis of the chevron plots reported in Fig. 8 allows calculation of the native and intermediate stability for each of the measured mutants (Table 2). Of particular interest, it may be noted that the mutations result in a clear destabilization of the intermediate state versus the

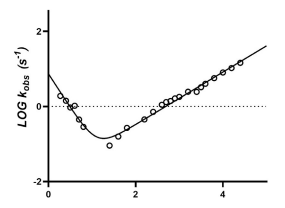


FIGURE 7 A semilogarithmic plot of the observed (un)folding rate constants (chevron plot) for wild-type AnxA2. The chevron plot of the observed rate of folding and unfolding is shown as a function of final denaturant concentration (GdnHCl). The total chevron plot is fit to a two-state equation.

native state. Importantly, in all cases, with the sole exception of the Q67A mutant, the analysis of thermodynamic parameters reports that such destabilization appears to be more evident in the intermediate state compared with the native state. This peculiar effect, whereby a mutation has a more pronounced effect on a transient intermediate state rather than on the native state, represents a characteristic signature of transient misfolding, suggesting that $AnxA2-D_I$ is misfolded in the intermediate state.

DISCUSSION

AnxA2 is a genuinely multifunctional protein (1-5), involved in a variety of cellular processes ranging from membrane-cytoskeleton interactions and actin dynamics to mRNA biogenesis. The latter process involves its binding to specific mRNAs and participation in the regulation of their transport (as exemplified by c-myc mRNA (35)) and/ or translation (as exemplified by p53 (36) and c-myc (6) mRNAs). Moreover, AnxA2 appears to have several roles in the nucleus (37). Furthermore, it functions as an extracellular protein (38,39) and constitutes a marker of exosomes derived from the endocytic pathway (www.exocarta.org/). Thus, it comes as no surprise that AnxA2 is closely involved in tumor progression by participating, for example, in metalloproteinase activation (40), cell migration (41), cell proliferation (42), and epithelial-mesenchymal transition (43). In several tumor types, it even serves as a specific marker for the more advanced/progressive states of the cancer (40,44). It has been proposed that AnxA2 serves as an ideal target for anticancer drugs (45). However, to target the fulllength protein by knockout or knockdown may not be an adequate strategy due to its multifunctionality and/or compensatory effects (46). Thus, a better approach may be to use peptides representing specific regions of AnxA2 to compete with specific cellular ligands that bind to this particular region. The proper design of such peptides highly depends on our understanding of the structure of the fulllength protein and of its individual domains. This also implies understanding the effects of perturbation of the AnxA2 structure on its folding and the underlying kinetics.

When considering the folding and stability of globular proteins or domains, one should distinguish two very distinct

TABLE 2 Kinetic and thermodynamic parameters for [1-339] and [35-339] wild-type AnxA2 and the indicated mutant AnxA2 variants

	$k_{D\text{-}I}\ (s^{-1})$	$k_{N-D} (10^{-5} s^{-1})$	$K_{I-N} (10^4)$	$\Delta \Delta G_{\text{D-N}} \; (\text{kcal mol}^{-1})$	$\Delta \Delta G_{D ext{-}I} ext{ (kcal mol}^{-1})$
[1–339] wild-type	6.5 ± 1.0	1.3 ± 0.7	0.2 ± 0.1		
[1-339] F33A-D38A-R63A-Q67A	1.3 ± 0.2	0.8 ± 3.7	5.5 ± 0.5	0.7 ± 0.2	2.5 ± 0.3
[1-339] D38A	1.1 ± 0.2	1.2 ± 0.6	2.3 ± 0.2	1.0 ± 0.2	2.4 ± 0.3
[1-339] V57E	2.8 ± 0.5	2.1 ± 1.0	6.3 ± 0.6	0.8 ± 0.2	2.7 ± 0.3
[1-339] R63A	3.7 ± 0.6	1.3 ± 0.6	1.9 ± 0.2	0.3 ± 0.2	1.6 ± 0.2
[1-339] Q67A	2.2 ± 0.4	4.8 ± 3.2	0.2 ± 0.1	1.4 ± 0.2	1.4 ± 0.2
[1-339] V98R	2.9 ± 0.5	60 ± 20	0.6 ± 0.1	2.7 ± 0.3	3.3 ± 0.3
[1-339] G101Y	3.0 ± 0.5	1.7 ± 0.8	1.7 ± 0.2	0.6 ± 0.1	1.8 ± 0.2
[35–339] wild-type	1.4 ± 0.2	2.4 ± 1.0	0.9 ± 0.1	1.2 ± 0.2	2.0 ± 0.2
[35–339] R63A	1.4 ± 0.2	2.9 ± 1.3	4.2 ± 0.4	1.4 ± 0.2	3.0 ± 0.3
[35–339] Q67A	1.3 ± 0.2	3.2 ± 1.4	1.8 ± 0.2	1.5 ± 0.2	2.7 ± 0.3

properties. A first property is the ability to adopt a cooperatively folded state, encoded by the protein sequence. The cooperatively folded state displays various characteristics, including a wide distribution of ¹H frequencies and a cooperative denaturation profile. The second property is the thermostability of the folded state, characterized—at least in part—by the temperature of the denaturation mid-point. Some mutations may alter the thermostability of the domain while preserving the ability to adopt a folded state. Other mutations, directed at particular "hot spots," may completely disrupt the ability to adopt a folded state (47). In the present work, the two types of effects have been observed, both for isolated AnxA2-D₁ and for full-length AnxA2.

A sequential folding model had been previously proposed for the annexin core structure (22,23). In this model, D_I folds first, whereas D_{II} and D_{III} remain partly unfolded until D_{IV} folds by docking on D_I, which in turns induces the complete folding of D_{II} and D_{III}. This model was primarily based on the observation that, within AnxA1, only the isolated domain D_I was capable of folding independently (22,23). While working on AnxA2, we also observed that AnxA2-D_I was also able to fold independently (12,25). This observation, together with the high conservation of annexin core sequences across the annexin family members, suggested that the folding process of the annexin core structure should be conserved throughout the entire family. We therefore decided to investigate if the proposed annexin core sequential folding model also applies to AnxA2.

As a first step, we identified mutations that are likely to disrupt the autonomous folding of isolated AnxA2-D_I. One possible approach would have been to mutate one or several residues within the central hydrophobic core of the domain, which hold together the five conserved helices. However, mutations within the hydrophobic core sometimes just decrease thermostability, without really disrupting the folding process. By carefully inspecting the structure, we realized that the network of contacts established by the F33/D38/R63/Q67 tetrad constitute a "structural knot," which might represent a critical bottleneck in the folding process of AnxA2-D_I. This hypothesis was indeed confirmed, as we found that the single-point mutations D38A and Q67A, directed at the tetrad, were both sufficient to completely disrupt the folding ability of D_I. In addition, we found that deletion of the N-terminal region of D_I until residue 34 also completely disrupted the folding ability of this domain. This can be explained by the fact that this deletion eliminated both the side chain of F33 and the main chain of Y30, which both provide contacts to the abovementioned "structural knot."

As a second step, we showed that re-introduction of the mutated D_I bearing any of the disrupting mutations (single-point, deletion, or combined) within the full-length annexin did not prevent the folding of the latter. This unambiguously demonstrated that, unexpectedly and in contradiction with the previously proposed folding model (22,23), the ability of D_I to fold independently is not a prerequisite for the folding of the full-length AnxA2.

As a third step, we performed other mutations of D_I, which were not intended to disrupt its folding, but rather aimed at disrupting the interface of D_I with D_{IV}. According to the stepwise folding model previously proposed for AnxA1, the protein can achieve its complete folding only when D_{IV} is allowed to dock onto D_{I} . We generated three mutations of D_I, which replaced interface residues of D_I with bulkier charged or polar residues, expected to perturb the interface. None of these mutations, either single-point or combined in a multiple mutant, disrupted the independent folding of the isolated D_I. Neither did they prevent fulllength AnxA2 to acquire a cooperatively folded structure. Nevertheless, some of these mutations strongly impacted the thermostability of full-length AnxA2, pointing to a role of interfacial residues in holding together the fourdomain cooperative arrangement. Going more in depth, we observed an inverse correlation between the impact of G101Y and V98R mutations on the thermostability of the individual D_I and their impact on the thermostability of the entire protein. G101Y strongly destabilized the domain yet destabilized only slightly the full-length protein. In full contrast, V98R hardly altered the stability of the isolated domain, yet it strongly destabilized the full-length AnxA2 and suggests that V98 holds a key position in the interfacial contacts between the domains (D_I with D_{IV}).

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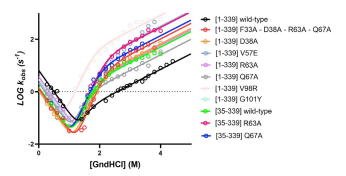


FIGURE 8 The chevron plots of wild-type AnxA2 and the indicated AnxA2 variants. Lines represent the best fit to an equation that describes a three-state folding mechanism, implying the presence of an intermediate along the reaction pathway. Chevron plots were globally fitted by sharing kinetic m values for all datasets (see materials and methods and results sections for details).

Hence, we found no correlation between the folding behaviors of D_I and full-length AnxA2, neither regarding their ability to fold nor regarding their thermostabilities. The ensemble of these data is in favor of a highly cooperative and robust folding process for AnxA2, which may be able to go through various alternative intermediate steps. What about other vertebrate annexins? AnxA7 and AnxA11 both harbor a very long N-terminus (\sim 20 kDa) and are difficult to obtain in a soluble state ((20) unpublished observations). Likewise, AnxA9 is notoriously prone to aggregation. Thus, it is expected that annexins that can easily be obtained in a soluble form also present a robust folding process stabilized by interfacial contacts like that seen in AnxA2. However, other annexins, less stabilized by interfacial contacts, may follow a different folding path.

The analysis of the kinetic data reported in this work appears to reconcile the stepwise mechanism, invoked for AnxA2, with a more cooperative view of folding. In fact, the accumulation of a misfolded intermediate involving more than one domain (with a β_T of 0.59), implies that, although D_I may represent a folding nucleus of AnxA2, it is prone to engage some nonnative interactions with the other domains before folding, as suggested by the analysis of thermodynamic parameters. In fact, although it is not surprising to observe that mutations disrupting the structure of D_I in isolation correspond to a change in stability in the case of the full-length protein, a comparison of $\Delta\Delta G$ values obtained from the analysis of kinetic data highlights a more pronounced destabilizing effect upon mutation on the intermediate state compared with the native state. This finding points to the evidence of the presence of a misfolded intermediate on the folding pathway of AnxA2. Future mutational analysis combined with kinetic (un)folding experiments may provide additional information useful to characterize these transient misfolding events in greater detail. Importantly, mutations affecting the structure of D_I in isolation appear to have no effect on kinetic m values calculated from the analysis of chevron plots. This aspect further corroborates the scenario of a robust cooperative folding reaction and suggests the folding pathway of the protein to be not affected by mutations.

In conclusion, we show that the folding process of full-length AnxA2 is highly cooperative and robust and does not rely on an initially folded $D_{\rm I}$. By contrast, hydrophobic interfacial amino acids of $D_{\rm I}$ are important in the folding of full-length AnxA2. Furthermore, our kinetic data indicate that AnxA2 folds through a transient nonnative intermediate, which is a novel finding. This novel insight into the folding process of AnxA2, and possibly also other annexins, provides the basis to dissect the cooperative folding process of proteins, in particular with repeating subunits.

SUPPORTING MATERIAL

Supporting material can be found online at https://doi.org/10.1016/j.bpj. 2022.10.043.

AUTHOR CONTRIBUTIONS

H.H.: methodology, validation, data curation, formal analysis, investigation, visualization, writing – part of original draft, writing – review & editing. J.R.: methodology, validation, data curation, formal analysis, investigation, writing – review & editing. Y.N.: methodology, validation, formal analysis, investigation, writing – review & editing. C.K.: methodology, validation, formal analysis, writing – review & editing. A.T.: methodology, validation, formal analysis, writing – review & editing. S.G.: methodology, validation, data curation, formal analysis, investigation, resources, supervision, writing – original draft, writing – review & editing. G.T.: conceptualization, funding acquisition, investigation, visualization, project administration, resources, supervision, writing – original draft, writing – review & editing. A.V.: conceptualization, funding acquisition, investigation, project administration, resources, supervision, writing – original draft, writing – review & editing.

ACKNOWLEDGMENTS

The study was supported by grants from the Research Council of Norway (Aurora program; no. 199747) and Helse Vest (no. 911499). We are grateful to Hanne Ravneberg (now Equinor) for excellent technical assistance and to Gunter Stier (Heidelberg University, Germany) for the pETM10 and pETM41 vectors.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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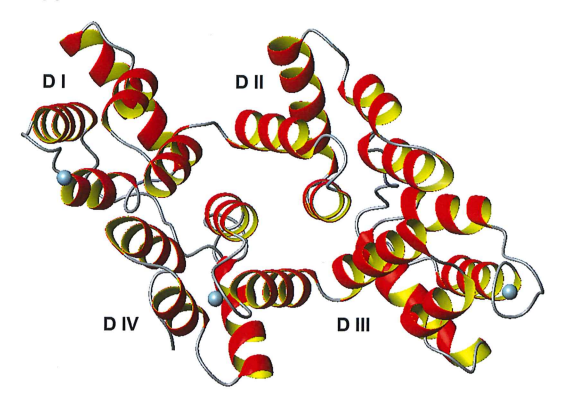
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Supplemental information

The cooperative folding of annexin A2 relies on a transient nonnative intermediate

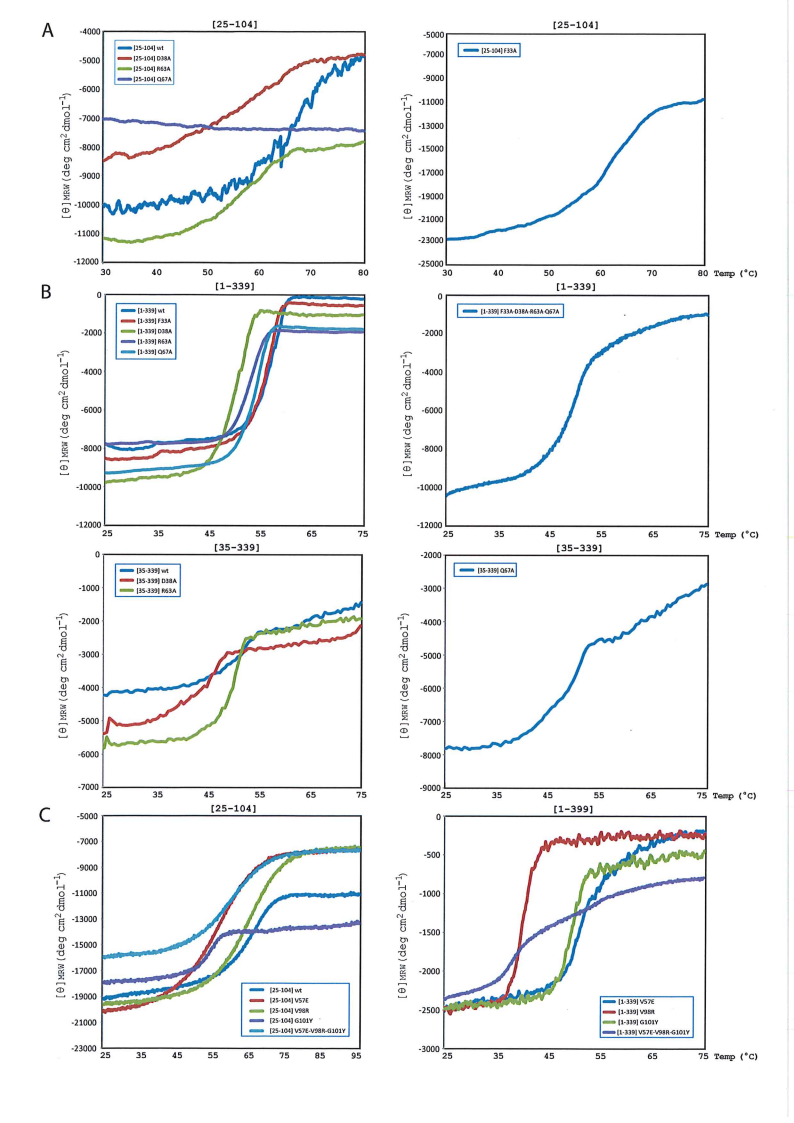
Hanne Hollås, Juan Ramirez, Yves Nominé, Camille Kostmann, Angelo Toto, Stefano Gianni, Gilles Travé, and Anni Vedeler

Supplemental material:



Supplementary Fig 1S.

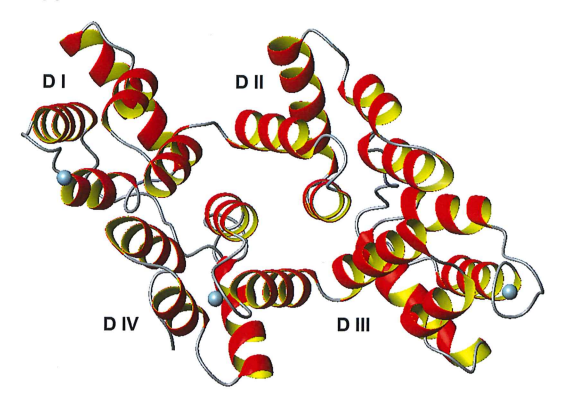
Ribbon plot of full-length AnxA2 (4x9p.pdb) showing the four domains (DI through DIV). Note that DI-DIV and DII-D-III constitute two tightly imbricated sub-modules with large interdomain helix-helix interfaces.



Supplementary Fig. S2

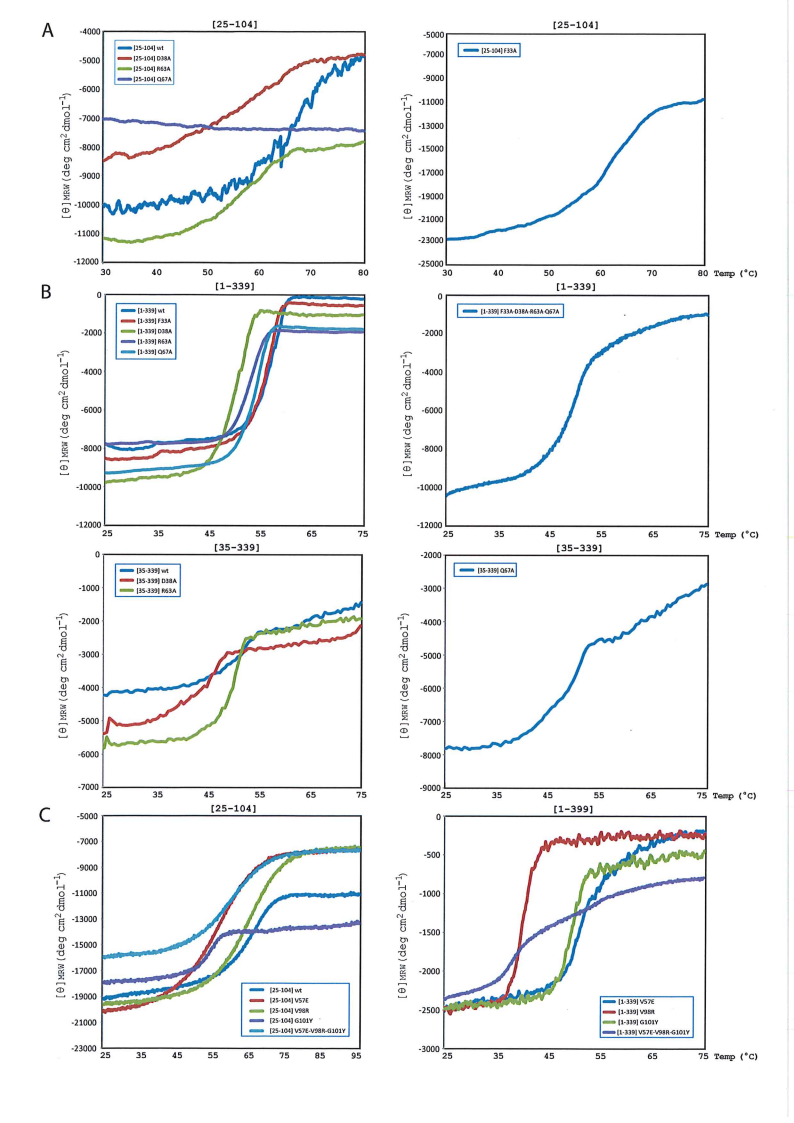
CD-monitored thermal disruption of α -helicity of (A) 10-30 μ M [25-104] (wt, F33A, D38A, R63A, Q67A) AnxA2; (B) 35-40 μ M [1-339] (wt, F33A, D38A, R63A, Q67A, F33A-D38A-R63A-Q67A) and 12-20 μ M [35-339] (wt, D38A, R63A, Q67A) AnxA2; (C) 20 μ M [25-104] and [1-339] (V57E, V98R, G101Y, V57E-V98R-G101Y) AnxA2. The change in ellipticity at 222 nm was measured at pH 7 at a heating rate of 100°C/h. The observed optical activity is expressed as the mean residue molar ellipticity [θ]_{MRW} (deg.cm².dmol⁻¹). The estimated apparent transition temperature (T_m) was determined from the first derivative of the curve.

Supplemental material:



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Ribbon plot of full-length AnxA2 (4x9p.pdb) showing the four domains (DI through DIV). Note that DI-DIV and DII-D-III constitute two tightly imbricated sub-modules with large interdomain helix-helix interfaces.



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