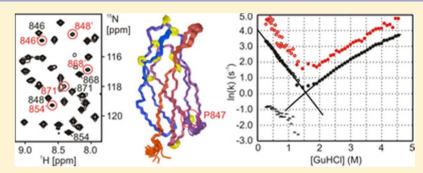
Structure and Unfolding of the Third Type III Domain from Human Fibronectin

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ABSTRACT: Fibronectin is a modular extracellular matrix protein that is essential for vertebrate development. The third type III domain (3FN3) in fibronectin interacts with other parts of fibronectin and with anastellin, a protein fragment that causes fibronectin aggregation. 3FN3 opens readily both as an isolated domain in solution and when part of fibronectin in stretched fibrils, and it was proposed that this opening is important for anastellin binding. We determined the structure of 3FN3 using nuclear magnetic resonance spectroscopy, and we investigated its stability, folding, and unfolding. Similar to most other FN3 domains, 3FN3 contains two antiparallel β -sheets that are composed of three (A, B, and E) and four (C, D, F, and G) β -strands, respectively, and are held together by a conserved hydrophobic interface. *cis*-*trans* isomerization of P847 at the end of β -strand C leads to observable conformational heterogeneity in 3FN3, with a *cis* peptide bond present in almost one-quarter of the molecules. The chemical stability of 3FN3 is relatively low, but the folding rate constant in the absence of denaturant is several orders of magnitude higher than the unfolding rate constants of other FN3 domains investigated to date. This unusually fast rate is comparable to the rate of binding of 3FN3 to anastellin at saturating anastellin concentrations, consistent with the model in which 3FN3 has to unfold to interact with anastellin.

 \mathbf{F} ibronectin is an essential extracellular matrix protein (reviewed in refs 1–3) that plays an important role in cell adhesion and migration and affects cell proliferation, survival, and differentiation. It is required for vertebrate development, and mice that lack fibronectin die during embryogenesis with a shortened anterior-posterior axis, absent notochord and somites, and abnormal heart and vasculature.⁴ Aside from development, fibronectin serves an important function in tissue repair.^{5–7}

Fibronectin is composed of 12 fibronectin type I (FN1), two fibronectin type II (FN2), and 15–17 fibronectin type III (FN3) homologous repeats and is secreted as an ~500 kDa disulfide-linked dimer. Formation of fibrils from soluble fibronectin dimers is not spontaneous and generally occurs only in the presence of cells, which assemble fibronectin fibrils around themselves. The molecular basis of the conversion of soluble fibronectin into insoluble fibrillar aggregates is poorly understood, and the atomic-resolution structure of the fibrils still remains to be elucidated.^{3,8} Some parts of fibronectin have been studied extensively, but relatively little information is available about the third FN3 domain (3FN3) (Figure 1). Together with the second FN3 domain, 3FN3 was proposed to maintain soluble fibronectin dimers in a compact conformation by ionic interactions with the 12th–14th FN3 domains⁹ and the 2nd–5th FN1 domains.¹⁰ 3FN3 was also reported to mediate cell adhesion and migration through binding to a so far unidentified β 1 integrin.¹¹ Finally, 3FN3 and several other FN3 domains were shown to interact with a small fragment of fibronectin called anastellin.^{12,13}

Anastellin exhibits anti-tumor, anti-metastatic, and antiangiogenic properties^{14,15} and requires endogenous fibronectin for these *in vivo* activities.^{16,17} The binding of anastellin to fibronectin leads to conversion of the soluble fibronectin dimers to insoluble aggregates that are reminiscent of

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		A	В	С		D	E	F	G
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	810	820	830	840	850	860	870	880	890 900
3FN3	APDAP	PDPTVDOVDD	TSIVVRWSRPO-	-APITGYRIVY	SPSVEGS	STELNLPE	TANSVTLSDLOP	GVOYNITIYAVEE-	NOESTPVVIOOETTG
1FN3		_	_				_	GVVYEGQLISIQQ-	~ ~~
2FN3	PFSPL	VATSESVTEITA	SSFVVSWVSAS-	-DTVSGFRVEY	ELSEE-GDE	PQYLDLPS	TATSVNIPDLLP	GRKYIVNVYQISE-	DGEQ <mark>S</mark> LILSTSQT <mark>T</mark> AP
7FN3	PLSPP	TNLHLE-ANPDI	GVLT <mark>VSW</mark> ERST-	TPDIT <mark>GYR</mark> ITI	TPTNGQQGN	SL <mark>E</mark> EVVHA	DQSSCTFDNLSP	GLE <mark>Y</mark> NVSVYTVKD-	DKESVPISDTIIPAV
8FN3	AVPPP	TDLRFTNIGP	DTMRVTWAPPP-	SIDLTNFLVRY	SPVKN-EED	VA <mark>E</mark> LSISP	SDNAVVLTNLLP	<mark>GTEY</mark> VVSVSSVYE-	QHE <mark>S</mark> TPLRGRQK <mark>T</mark> GL
9FN3	GLDSP	TGIDFSDITA	-N <mark>SFTVHW</mark> IAPR-	-ATITGYRIRE	HPEHF-SGR	PR <mark>E</mark> DRVPH	SRNSITLTNLTP	GTEYVVSIVALNG-	REE <mark>S</mark> PLLIGQQS <mark>T</mark> VS
10FN3	VSDVP	RDLEVVAATP	-T <mark>S</mark> LLISWDAPA-	-VTVRYYRITY	GETGG-NSP	VQ <mark>E</mark> FTVPG	SKSTATISGLKP	<mark>GVDY</mark> TITVYAVTGR	GDSPAS <mark>S</mark> KPISINYR <mark>T</mark> EI
12FN3	AIPAP	TDLKFTQVTP	-T <mark>S</mark> LSAQ <mark>W</mark> TPPN-	-VQLTGYRVRV	TPKEK-TGP	MK <mark>E</mark> INLAP	DSSSVVVSGLMV.	ATK <mark>Y</mark> EVSVYALKD-	TLT <mark>S</mark> RPAQGVVT <mark>T</mark> LE
13FN3	NVSPP	RRARVTDATE	-TTITIS <mark>W</mark> RTKT-	-ETITGFQVDA	VPANGQT	PIQRTIKP	DVRSYTITGLQP	GTD <mark>Y</mark> KIYLYTLND-	NAR <mark>S</mark> SPVVIDAS <mark>TA</mark> I
14FN3	AIDAP	SNLRFLATTP	-N <mark>S</mark> LLVS <mark>W</mark> QPPR-	-ARITGYIIKY	EKPGSP	PR <mark>E</mark> VVPRPRP	GVTEATITGLEP	GTEYTIYVIALKN-	NQK <mark>S</mark> EPLIGRKK <mark>T</mark> DE
EDA	NIDRP	KGLAFTDVDV	-D <mark>S</mark> IKIA <mark>W</mark> ESPQ-	-GQVSR <mark>YR</mark> VTY	SSPEDG	IH <mark>E</mark> LFPAPDG	EEDTAELQGLRP	GSEYTVSVVALHD-	DME <mark>S</mark> QPLIGTQS <mark>T</mark> AI
EDB	EVPQL	TDLSFVDITD	SSIGLRWTPLN-	SSTII <mark>GYR</mark> ITV	VAAGE-GIP	IF <mark>E</mark> DFVDS	SVGYYTVTGLEP	GID <mark>Y</mark> DISVITLIN-	GGE <mark>S</mark> APTTLTQQ <mark>T</mark> AV

Figure 1. Structure-based sequence alignment of FN3 domains from human fibronectin. The sequences are colored by percent identity.⁵⁷ The sequence of 3FN3 and a diagram of its secondary structure are shown at the top. EDA and EDB stand for the alternatively spliced extra domain A and B, respectively. The 4th–6th, 11th, and 15th FN3 domains are not included because their structures are not available.

Table 1. Experimental Restraints and Structural Statistics for 3FN3

no. of peaks in NOESY spectra					
3D ¹⁵ N-edited NOESY	1057				
3D ¹³ C-edited aliphatic NOESY (without sensitivity enhancement)	1722				
3D ¹³ C-edited aliphatic NOESY (sensitivity-enhanced)	906				
3D ¹³ C-edited aromatic NOESY	64				
2D NOESY in H ₂ O	1912				
2D NOESY in D_2O	1343				
2D ¹³ C- and ¹⁵ N-filtered, ¹³ C- and ¹⁵ N-edited NOESY	25				
no. of experimental restraints					
NOE distance restraints in the last ARIA iteration					
intraresidual	873				
sequential	454				
medium-range $(2 \le i - j \le 5)$	171				
long-range $(6 \le i - j)$	758				
ambiguous	1179				
dihedral angle (Φ and Ψ)	168				
hydrogen bond distance	60				
no. of experimental restraint violations					
distance (NOE and hydrogen bond) violations >0.5 Å	0.2 ± 0.4				
dihedral angle violations >5°	1.1 ± 0.9				
root-mean-square deviation (rmsd) from experimental restraints					
distance (NOE and hydrogen bond) restraints (Å)	0.040 ± 0.002				
dihedral angle restraints (deg)	0.9 ± 0.1				
rmsd from idealized geometry					
bonds (Å)	0.0053 ± 0.0001				
angles (deg)	0.63 ± 0.01				
impropers (deg)	1.75 ± 0.06				
rmsd of residues 811–901 from mean coordinates					
backbone atoms (N, Ca, C') (Å)	0.33 ± 0.08				
heavy atoms (Å)	0.73 ± 0.06				
distribution of Φ and Ψ dihedral angles of residues 811–901 in the Ramachandran plot 56 (%)					
most favored regions	83.6				
additional allowed regions	16.1				
generously allowed regions	0.1				
disallowed regions	0.3				

fibronectin fibrils deposited by cells.^{18,19} It is not known how anastellin interacts with its target FN3 domains and why this interaction causes aggregation of fibronectin. It was proposed that transient opening of the target FN3 domains is important for binding to anastellin.^{12,13,20}

We employed nuclear magnetic resonance (NMR) spectroscopy, equilibrium denaturation, and stopped-flow methods to characterize 3FN3 and its opening behavior. We determined the solution structure of 3FN3, and we identified a minor 3FN3 conformer that is populated under native conditions. We also measured the stability of 3FN3 and investigated its folding and unfolding kinetics, and we show that the unfolding rate constant in the absence of denaturant is significantly higher than the unfolding rate constants reported to date for other isolated FN3 domains. This unusually fast unfolding rate is in the same range as the rate of binding of 3FN3 to anastellin at

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saturating anastellin concentrations, consistent with the model in which 3FN3 has to unfold to interact with anastellin.

EXPERIMENTAL PROCEDURES

Expression and Purification of 3FN3. The pET15b expression vector encoding 3FN3 was provided to us by T. Ohashi and H. Erickson (Duke University, Durham, NC).^{12,13} The His-tagged fusion protein was expressed in Escherichia coli BL21-CodonPlus(DE3)-RIPL cells (Agilent) and purified by affinity chromatography on a HisTrap FF column (GE Healthcare). The N-terminal His tag was cleaved off with thrombin, and 3FN3 was separated from thrombin and the His tag by affinity chromatography on HiTrap Benzamidine FF (high sub) (GE Healthcare) and HisTrap FF columns, respectively. 3FN3 was then further purified by size-exclusion chromatography on Superdex 75 resin (GE Healthcare). The recombinant 3FN3 protein consisted of six extraneous amino acids at the N-terminus, residues 808-905 of human fibronectin (P02751), and two extraneous amino acids at the C-terminus (gshmgtTTAP…PRSDgt).

NMR Spectroscopy. Samples for NMR spectroscopy typically contained 0.6 mM unlabeled, uniformly ¹⁵N-labeled, or uniformly ¹³C- and ¹⁵N-labeled 3FN3 in phosphate-buffered saline (PBS) (pH 7.5) supplemented with 10% ²H₂O. Most NMR data were acquired at 25 °C on a Varian 600 MHz NMR System that was equipped with a triple-resonance probe or a ¹³C-enhanced salt-tolerant cold probe. NMR experiments that were used to assign ¹H, ¹³C, and ¹⁵N chemical shifts and to generate restraints for structure calculations included twodimensional (2D) ¹H-¹⁵N heteronuclear single-quantum correlation (HSQC), ¹H-¹³C aliphatic and aromatic HSQC, double-quantum-filtered correlation spectroscopy (COSY), total correlation spectroscopy (TOCSY), nuclear Overhauser effect spectroscopy (NOESY), HBCB(CGCD)HD, ¹H-¹⁵N heteronuclear multiple-quantum correlation (HMQC) optimized for histidine imidazole rings, three-dimensional (3D) HNCACB, CBCA(CO)NH, C(CO)NH, H(CCO)NH, HNCO, HN(CA)CO, CCH-TOCSY, HCCH-TOCSY, ¹⁵Nedited TOCSY, ¹⁵N-edited NOESY, ¹³C-edited aliphatic NOESY (with and without sensitivity enhancement), and ¹³C-edited aromatic NOESY. In addition, a 2D ¹³C- and ¹⁵Nfiltered, ¹³C- and ¹⁵N-edited NOESY spectrum²¹ was acquired on a Varian 800 MHz NMR System equipped with a salttolerant ¹³C-enhanced cold probe for a sample containing 0.9 mM ¹³C- and ¹⁵N-labeled 3FN3 and 0.9 mM unlabeled 3FN3. To evaluate backbone conformational flexibility, the steadystate ¹H-¹⁵N nuclear Overhauser effect (NOE) was measured using 2D heteronuclear correlation experiments. All data were processed with NMRPipe²² and analyzed with CcpNmr Analysis version 2.2.2.²³ Φ and Ψ dihedral angle restraints were generated with TALOS-N.²⁴

Calculation of 3FN3 Structure. Structure calculations were performed using Aria 2.3^{25} in conjunction with CNS 1.21,²⁶ with peaks from the NOESY spectra and dihedral angle restraints from TALOS-N as input (Table 1). Manual assignments of the NOESY peaks were retained. ¹H–¹⁵N NOE values and TALOS-N predictions indicated that the N-and C-terminal tails were dynamic, and no medium- or long-range connectivities were identified for residues preceding A810 or following T901 during manual inspection of the NOESY spectra. Aria calculations nevertheless tended to create restraints between the tails and the rest of the protein even

when other, more likely assignments were available. To prevent the generation of such artifactual restraints for the tails, only chemical shifts for residues T809–P902 were included in the calculations. In addition, manually identified intraresidual and sequential NOESY peaks stemming from the dynamic residues outside of the T809–P902 region were omitted from the input.

The calculation protocol employed spin-diffusion correction,²⁷ a quadrupled number of cooling steps during simulated annealing,²⁸ and otherwise mostly default Aria parameters and options. Hydrogen bond connectivity in the β -sheets was established on the basis of NOE patterns, delayed ${}^{1}H-{}^{2}H$ exchange, and the presence of the hydrogen bonds in preliminary structures. Distance restraints were then introduced in subsequent Aria runs to reinforce the expected hydrogen bonds. In the final run, a total of 80 structures were calculated and 25 of them were refined in explicit water. The statistics for the 25 refined structures are summarized in Table 1. Figures depicting the structures were prepared with the molecular graphics program Molmol 2K.1.29 The coordinates and chemical shifts were deposited in the Protein Data Bank (PDB entry 2N1K) and the Biomagnetic Resonance Bank (BMRB entry 25564).

Equilibrium Denaturation Studies of 3FN3. Samples for equilibrium denaturation studies contained 1 μ M 3FN3 and 0– 6 M guanidine hydrochloride (GuHCl) in PBS (pH 7.4). After being mixed, the samples were allowed to equilibrate at room temperature for at least 2 h, and their fluorescence at 350 nm was then measured in a JASCO FP-8200 spectrofluorometer at 24 °C, using excitation at 280 nm. For each GuHCl concentration tested, fluorescence intensities from triplicate samples were averaged. The concentration of GuHCl at which half of the protein is unfolded, [GuHCl]50%, and the proportionality constant between the free energy of unfolding and the GuHCl concentration, m_{D-N} , were obtained by fitting eq 11 from ref 30 to the denaturation data. The fitting was done in Matlab version 2012a (MathWorks Inc., Natick, MA). The free energy of unfolding in the absence of denaturant was calculated as $\Delta G_{D-N}^{H_2O} = m_{D-N} [\text{GuHCl}]_{50\%}$.

Folding and Unfolding Kinetics. The changes in fluorescence that accompany folding and unfolding of 3FN3 were monitored in an Applied Photophysics SX20 stopped-flow spectrometer at 23 °C, using a wavelength of 280 nm for excitation and a 320 nm cutoff filter to detect emission. Unfolding was initiated by rapidly mixing 1 volume of 10.6 μ M 3FN3 in PBS (pH 7.4) with 10 volumes of PBS containing 1.40-5.00 M GuHCl. To initiate refolding, 1 volume of 10.6 µM 3FN3 and 3.00 M GuHCl in PBS was mixed with 10 volumes of PBS containing 0-1.14 M GuHCl. Five 5 s fluorescence traces were collected and averaged for each final GuHCl concentration. A double-exponential function was then fitted to the unfolding traces and a triple-exponential function to the refolding traces. A 3 min trace that was recorded for folding in 0.60 M GuHCl was fitted with a quadrupleexponential function. The fitting was performed with the Pro-Data Viewer 4.1.18 software provided with the stopped-flow instrument. The unfolding and refolding rate constants in the absence of denaturant, $k_u^{H_2O}$ and $k_f^{H_2O}$, the proportionality constants between $\ln(k_u)$ or $\ln(k_f)$ and the GuHCl concentration, m_{ku} and m_{kp} and a constant describing the curvature of the unfolding rate, c, were obtained by fitting eqs 1 and 2 from ref 31 to the rate constants for the major unfolding and refolding phases, respectively. The fitting was conducted in Matlab.

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RESULTS AND DISCUSSION

3FN3 is very soluble and stable, and its NMR spectra are of high quality (Figure 2). We were therefore able to assign the

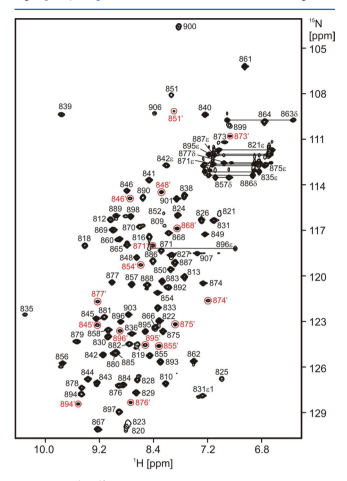


Figure 2. 2D 1 H $^{-15}$ N HSQC spectrum of 3FN3. The assignments are indicated. Cross-peaks from the minor species that do not adjoin their respective major peaks are highlighted in red and denoted by an apostrophe. Signals from T825 and R842 ε are aliased from 97.76 and 83.67 ppm, respectively.

majority of its ¹H, ¹³C, and ¹⁵N chemical shifts, and we determined its solution structure on the basis of 2D and 3D homo- and heteronuclear NMR experiments.

The structure of 3FN3 is shown in Figure 3. Similar to most other FN3 domains, $^{32-43}$ 3FN3 contains seven β -strands (labeled A-G) that form a three-stranded antiparallel β -sheet (β -strands A, B, and E) and a four-stranded antiparallel β -sheet (β -strands C, D, F, and G); β -strands A and G each contain a bulge (Figure 3A,B). The two β -sheets are held together by an extensive hydrophobic interface that includes several residues highly conserved among FN3 domains, in particular P814, W831, Y841, L870, Y876, and T898 (Figure 3C,D). The hydroxyl groups of Y841 and T898 are buried, and the signals from the hydroxyl hydrogens can be observed in several NMR spectra (Figure 4). The loops between β -strands A and B, D and E, E and F, and F and G are well-ordered, but the intersheet loop connecting β -strands B and C and the intrasheet loop connecting β -strands C and D exhibit increased mobility, as indicated by the ${}^{1}H-{}^{15}N$ NOE data (Figure 5).

The 2D $^{1}H-^{15}N$ HSQC spectrum contains a set of secondary, weaker cross-peaks that stem from a minor 3FN3

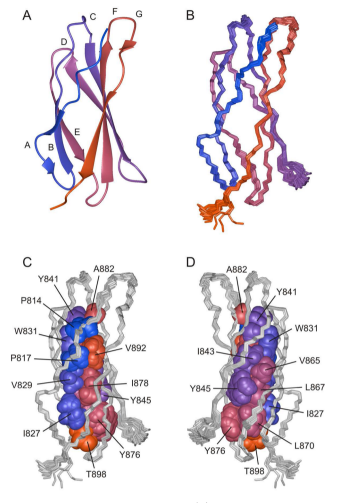


Figure 3. Solution structure of 3FN3. (A) Ribbon model of 3FN3 (residues 811–901). The color changes smoothly from blue at the N-terminus to red at the C-terminus, and the individual β -strands are labeled. (B) Twenty-five superimposed backbone traces of 3FN3. The coloring scheme and orientation are the same as in panel A. (C and D) Hydrophobic core in 3FN3. To illustrate packing at the interface between the two β -sheets, the side chains of buried residues are shown in a space filling representation and are colored as in panels A and B. The backbone traces are colored gray. The orientation in panel C is the same as in panels A and B, and the view in D is related by a 180° rotation around the vertical axis.

species (Figure 2). The population of this species is $23 \pm 3\%$ as judged from peak volumes. The residues whose backbone amide chemical shifts differ the most between the minor and major forms are clustered in the C-terminal half of the fourstranded β -sheet (Figure 6A). There are a total of 11 prolines in 3FN3, and we therefore investigated whether proline cis-trans isomerization might be responsible for the heterogeneity. The majority of the prolines are located at the N-terminal end of the domain, and only two, P847 and P872, are in the part of 3FN3 for which the heterogeneity in NMR spectra is observed (Figure 6A). The $C\alpha$, $C\beta$, and $C\gamma$ chemical shifts of P872 in the minor species are virtually identical to those in the major form (Figure 6D). In contrast, the chemical shifts of P847 in the minor species (C α at 62.48 ppm, C β at 34.93 ppm, and C γ at 24.59 ppm) are significantly different from those in the major form (C α at 62.70 ppm, C β at 32.08 ppm, and C γ at 27.75 ppm) (Figure 6C), and they are indicative of the S846-P847 peptide bond in the minor species being in a cis

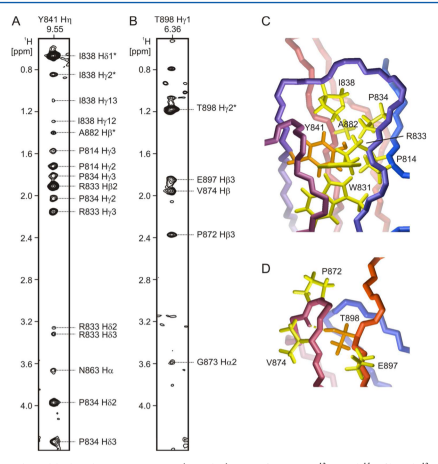


Figure 4. NMR signals from buried hydroxyl groups in 3FN3. (A and B) Strips from a 2D 13 C- and 15 N-filtered, 13 C- and 15 N-edited NOESY spectrum of 3FN3 (mixing time of 200 ms) with cross-peaks between Y841 H η (panel A) or T898 H γ 1 (panel B) and nearby aliphatic hydrogens. The assignments for all observed signals are indicated. The NOESY spectrum also contains cross-peaks between Y841 H η and W831 H ϵ 1 and between T898 H γ 1 and V874 HN (not shown). (C and D) Close-up views of Y841 and T898, respectively, in the 3FN3 structure. The backbone is colored as in panels A and B of Figure 3, and the side chains of Y841 and T898 are colored orange. The side chains of residues that are near the hydroxyl hydrogens according to the NOESY data are also shown and are colored yellow. T898 H γ 1 (D) forms a hydrogen bond (yellow dashed line) to the backbone carbonyl oxygen of Q871.

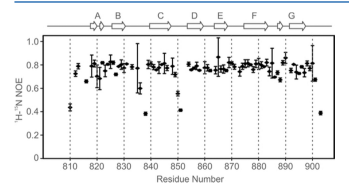


Figure 5. Backbone amide dynamics in 3FN3: ${}^{1}H{-}{}^{15}N$ NOE data. The secondary structure of 3FN3 is outlined at the top. The ${}^{1}H{-}{}^{15}N$ NOE values for D905, g906, and t907 are negative and are not included in the figure.

conformation.⁴⁴ The largest backbone amide chemical shift differences between the two forms are observed for S848, which immediately follows P847, for T854 in β -strand D, which flanks P847 on one side, and for Y876 and N877 in β -strand F, which flank P847 on the other side (Figure 6B). Hence, all our data point to the *cis*-*trans* isomerization of the S846–P847 peptide bond as the primary source of the conformational heterogeneity in 3FN3.

cis—*trans* isomerization of peptidyl—prolyl bonds in the context of a native state was observed previously in other folded proteins and, in some cases, was shown to be functionally important and to act as a molecular switch.^{45–49} The functional significance of the *cis*—*trans* isomerization of P847 in 3FN3, however, is not clear at this point, and the heterogeneity may just be a consequence of the relatively low stability of 3FN3 (see below) and the inability to stabilize one isomer strongly over the other.^{48,49} Interestingly, a number of the FN3 domains in fibronectin contain a proline residue at the position equivalent to P847 (Figure 1), but a population of molecules with a *cis* peptide bond has so far not been identified in any of the structures determined by NMR spectroscopy⁴⁰ or X-ray crystallography.^{35,37,42,43}

3FN3 interacts with anastellin, and it was proposed that the rate of binding at saturating anastellin concentrations (~0.02 s⁻¹) is determined by the rate at which 3FN3 spontaneously opens and exposes an otherwise cryptic binding site.¹³ However, the rates of unfolding that have been reported so far for other FN3 domains under comparable conditions are significantly slower than 0.02 s⁻¹ [(7.21 ± 0.26) × 10⁻⁵ s⁻¹ for the 3rd FN3 domain from tenascin⁵⁰ and (2.3 ± 1.2) × 10⁻⁴ s⁻¹ for the 10th FN3 domain from fibronectin (10FN3)⁵¹]. Experiments probing the accessibility of engineered buried cysteines in FN3 domains suggested that 3FN3 opens readily,

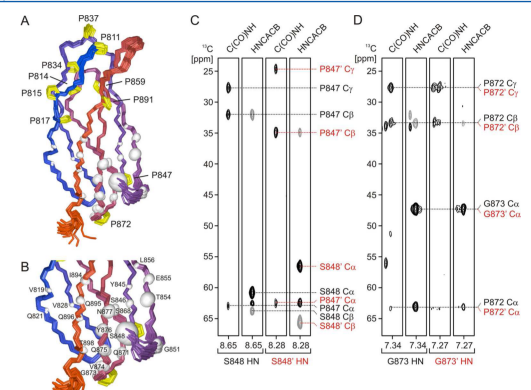


Figure 6. Conformational heterogeneity in 3FN3. (A) Backbone amide chemical shift heterogeneity mapped onto the 3D structure of 3FN3. The backbone amide nitrogen of each residue with a minor cross-peak in the $2D \ ^{1}H^{-15}N$ HSQC spectrum (Figure 2) is displayed as a white sphere. The volume of the sphere is proportional to the chemical shift difference between the cross-peaks of the major and minor form. The backbone traces are colored as in panels A and B of Figure 3. Proline side chains are colored yellow and labeled to help visualize their positions relative to the residues whose chemical shifts differ between the two forms. P902, which is located in the dynamic C-terminal tail, is not shown. The view is related to the orientation in Figure 3A–C by a 40° rotation around the vertical axis. (B) Close-up view with the residues labeled. (C and D) ^{13}C chemical shifts of P847 and P872, respectively, as observed in the 3D C(CO)NH and HNCACB spectra. The left two strips in each panel are for the major conformer, and the right two strips are for the minor conformer. Positive contour levels are colored black and negative levels gray. The chemical shift assignments for the minor conformer are denoted with an apostrophe and are colored red, and the assignments for the major conformer are colored black.

but the opening and closing frequency was not measured.⁵² This lack of information about the opening rates and opening equilibrium prompted us to undertake a rigorous investigation of the unfolding and refolding behavior of 3FN3.

To evaluate the stability of 3FN3, we measured the intrinsic fluorescence of the single tryptophan, W831, as a function of GuHCl concentration. The aromatic ring of W831 is buried in the hydrophobic interface between the two β -sheets (Figure 3) and becomes exposed to solvent when 3FN3 unfolds; this is accompanied by a large increase in fluorescence emission at 350 nm (Figure 7). Fitting of the fluorescence denaturation curve

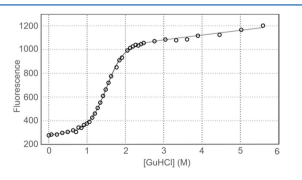


Figure 7. Equilibrium denaturation of 3FN3 in GuHCl. The black curve represents the equation fitted to the data.

yielded a concentration of GuHCl at which half of 3FN3 is unfolded, $[GuHCl]_{50\%}$, of 1.51 ± 0.02 M and a proportionality constant between the free energy of unfolding and GuHCl concentration, m_{D-N} , of 2.65 ± 0.17 kcal mol⁻¹ M⁻¹. The free energy of unfolding in the absence of denaturant, $\Delta G_{D-N}^{H_2O}$, was then calculated from these values to be 4.00 ± 0.26 kcal mol⁻¹. This corresponds to ~1 of every 880 molecules being unfolded under native conditions. No deviation from a two-state transition was apparent, suggesting that the stabilities of the major and minor conformer are comparable.

The kinetics of 3FN3 unfolding and refolding were also monitored using W831 fluorescence. Two unfolding and three refolding phases were resolved in 5 s fluorescence traces at all GuHCl concentrations investigated, and an additional slow refolding phase was detected in a 3 min experiment (Figure 8). Outside of the transition region, the amplitudes of the individual phases are relatively independent of GuHCl concentration. The natural logarithm of the rate constant for the major refolding phase (empty black circles in Figure 8A) varies linearly with GuHCl concentration. A linear equation, $\ln(k_f) = \ln(k_f^{'H_2O}) - m_{kf}$ [GuHCl], was therefore fitted to the data, yielding a refolding rate constant in the absence of denaturant, $k_f^{H_2O}$, of 58 ± 3 s⁻¹, and a proportionality constant, m_{kp} of 2.57 \pm 0.08 M⁻¹. In contrast, the natural logarithm of the rate constant for the major unfolding phase (filled black circles in Figure 8A) displays a nonlinear dependence on

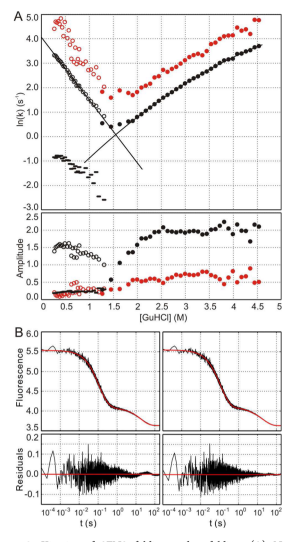


Figure 8. Kinetics of 3FN3 folding and unfolding. (A) Natural logarithm of the rate constants (top) and the amplitudes (bottom) shown as a function of GuHCl concentration for two unfolding (black and red filled circles) and three refolding (black and red empty circles and black dashes) phases. The slowest refolding phase that was detected in only a 3 min experiment is not included in the plot. The solid black lines represent the equations fitted to the data for the major unfolding and refolding phases. (B) Triple-exponential function (left) or quadruple-exponential function (right) fitted to a 3 min refolding trace in 0.60 M GuHCl. The residuals are shown in the bottom panels. The rate constants and amplitudes for the quadruple-exponential function are as follows: $k_1 = 17.5 \pm 2.2 \text{ s}^{-1}$, $a_1 = 0.38 \pm 0.09 (20\%)$, $k_2 = 6.7 \pm 0.3 \text{ s}^{-1}$, $a_2 = 1.06 \pm 0.09 (55\%)$, $k_3 = 0.28 \pm 0.06 \text{ s}^{-1}$, $a_3 = 0.07 \pm 0.01 (4\%)$, $k_4 = 0.047 \pm 0.002 \text{ s}^{-1}$, and $a_4 = 0.40 \pm 0.01 (21\%)$.

GuHCl concentration, and a second-degree polynomial, $\ln(k_u) = \ln(k_u^{H_2O}) + m_{ku}[\text{GuHCl}] - c[\text{GuHCl}]^2$, was used to fit the data. The fitting yielded an unfolding rate constant in the absence of denaturant, $k_u^{H_2O}$, of 0.057 \pm 0.010 s⁻¹, a proportionality constant, m_{ku} , of 2.1 \pm 0.1 M⁻¹, and a curvature, *c*, of 0.15 \pm 0.02 M⁻². The rate of 3FN3 folding is within the range observed for other FN3 domains, but the rate of unfolding is several orders of magnitude higher than those reported previously^{50,51} and is comparable to the rate of binding of anastellin to 3FN3 at saturating anastellin concentrations.¹³

Interestingly, we also observed a minor unfolding and a minor refolding phase whose rate constants were slightly higher than those for the major phase (filled and empty red circles in Figure 8A). The slopes and the positions of the minima in the $\ln(k)$ versus [GuHCl] plot for the minor and the major phases are very similar, and consequently, the plot for these phases has a "nested chevron" appearance (Figure 8A). We attribute the minor unfolding and refolding phase to unfolding and refolding of the minor 3FN3 conformer that contains a cis S846-P847 peptide bond. This interpretation is consistent with the stabilities of the minor and major conformer being indistinguishable in equilibrium denaturation experiments, and with the structure of the minor conformer overall closely resembling that of the major species (\sim 70% of backbone amide chemical shifts are identical in the two forms). The minor kinetic phase accounts for $25 \pm 5\%$ of the unfolding amplitude, in excellent agreement with the population of the minor conformer that was estimated from the NMR data.

In addition to the major phase and the fast minor phase described above, we identified two slow minor refolding phases (Figure 8B and black dashes in Figure 8A). In 0.6 M GuHCl, these slow phases represent ~25% of the total refolding amplitude (Figure 8B). We attribute the slow phases to processes in which proline *cis*-*trans* isomerization is the rate-limiting step, i.e., to refolding of denatured 3FN3 in which some of the prolines are in the non-native *cis* conformation.⁵³

The kinetic data are in general consistent with a two-state folding transition, as demonstrated using several wellestablished tests.^{31,50,54} We do not observe a significant burst phase or deviation from linearity in the $\ln(k_f)$ versus [GuHCl] plot for the major refolding phase. In addition, the equilibrium unfolding constant derived from the kinetic data and adjusted for proline isomerization in the denatured and native state corresponds to a free energy of unfolding, $\Delta G_{D-N}^{H_2O}$ of 3.89 \pm 0.12 kcal mol⁻¹. This value compares well with the free energy of unfolding from equilibrium experiments, 4.00 \pm 0.26 kcal mol⁻¹. Finally, the *m*-value calculated from the kinetic data, $m_k = RT(m_{kf} + m_{ku})$, is 2.76 \pm 0.08 kcal mol⁻¹ M⁻¹, in good agreement with the *m*-value obtained from equilibrium experiments, 2.65 \pm 0.17 kcal mol⁻¹ M⁻¹.

We have inspected the sequence and structure of 3FN3 for features that might explain the relatively low stability and high unfolding rate. The structure of 3FN3 is very similar to that of other FN3 domains, and most residues in its hydrophobic core are identical to those in the markedly more stable 10FN3 or are conservatively substituted (Figures 1 and 3). The only residue in the hydrophobic core that significantly differs from 10FN3 and from the FN3 consensus is P817. In 10FN3 and most other FN3 domains, this position is occupied by leucine (Figure 1). In addition to hydrophobic core packing, electrostatic interactions are also known to play a role in protein stability.⁵⁵ 3FN3 is highly acidic, with six aspartates, seven glutamates, three arginines, and no lysines or histidines within the structured region (Figure 9). It is therefore likely that unfavorable electrostatic interactions contribute to the lower stability and faster unfolding rate. The only residue that has so far been shown to destabilize 3FN3, D869, is in fact acidic.¹³ Most other FN3 domains contain a glycine or an asparagine at this position (Figure 1), and a D869G mutation in 3FN3 increases the chemical stability of the domain and decreases the rate of binding to anastellin.

Overall, our results demonstrate that while the structure and folding behavior of 3FN3 are similar to those of other FN3 domains, 3FN3 differs from previously characterized FN3 domains in several aspects. First, 3FN3 is conformationally

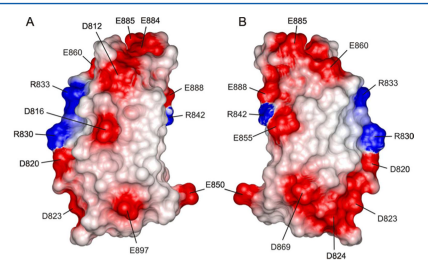


Figure 9. Distribution of charged residues in 3FN3. The solvent accessible surface of 3FN3 is colored according to electrostatic potential, with areas of negative, positive, or neutral electrostatic potential red, blue, or white, respectively. In panel A, the orientation is the same as in Figure 3A–C, and the view in panel B is related by a 180° rotation around the vertical axis.

heterogeneous because of a *cis*-*trans* peptide bond isomerization at the semiconserved P847. Second, 3FN3 displays relatively low stability and an unfolding rate significantly higher than those of other isolated FN3 domains investigated to date. Ohashi et al. proposed that when 3FN3 interacts with anastellin, the rate of binding at saturating anastellin concentrations (~0.02 s⁻¹) is determined by the rate at which 3FN3 spontaneously opens.¹³ We have now shown that the experimentally determined rate of 3FN3 unfolding (~0.06 s⁻¹) is comparable to the rate observed by Ohashi, consistent with the model in which 3FN3 has to open to bind anastellin.

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Notes

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ABBREVIATIONS

2D, two-dimensional; 3D, three-dimensional; 10FN3, 10th fibronectin type III domain from human fibronectin; 3FN3, 3rd fibronectin type III domain from human fibronectin; COSY,

correlation spectroscopy; EDA, extra domain A; EDB, extra domain B; FN1, fibronectin type I; FN2, fibronectin type II; FN3, fibronectin type III; GuHCl, guanidine hydrochloride; HMQC, heteronuclear multiple-quantum correlation; HSQC, heteronuclear single-quantum correlation; NMR, nuclear magnetic resonance; NOE, nuclear Overhauser effect; NOESY, nuclear Overhauser effect spectroscopy; PBS, phosphate-buffered saline; TOCSY, total correlation spectroscopy.

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