The Maturational Refolding of the β-Hairpin Motif of Equine Infectious Anemia Virus Capsid Protein Extends Its Helix α1 at Capsid Assembly Locus

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Background: The function of the maturational refolded N-terminal β-hairpin in retroviral capsid remains unknown.

Results: Folding the β-hairpin of equine infectious anemia virus (EIAV) capsid extends its downstream helix α1 at the N-terminus, which forms the oligomerization core of retroviral capsids.

Conclusion: The β-hairpin structures helix α1, which could be necessary for capsid assembly.

Significance: Solution NMR revealed the function of the puzzling β-hairpin motif in retroviral capsid.

A retroviral capsid (CA) protein consists of two helical domains, CA N and CA C, which drive hexamer and dimer formations, respectively, to form a capsid lattice. The N-terminal 13 residues of CA refold to a β-hairpin motif and the reversible quaternary packing of CA domains, the 17-kDa N-terminal (CAN) and the 9-kDa C-terminal (CA C), which could be necessary for capsid assembly.

Viruses in the family Retroviridae, e.g. human immunodeficiency virus (HIV), simian immunodeficiency virus, avian sarcoma virus, and equine infectious anemia virus (EIAV),3 utilize reverse transcription to replicate the genetic information stored in their RNA genome (1). Mutational analyses indicate that the viral capsid plays a critical role in this event because even small structural changes can disrupt the process (2–8). A retrovirus cycles through morphological transitions: viral particle assembly, budding, and maturation occurs in producer cells; the particle is disassembled in the newly infected target cell. At the onset of HIV-1 assembly, the structural precursor polyprotein, Gag, assembles underneath the plasma membrane (9–11). During or after budding, maturational changes proceed wherein the full-length Gag proteins in the assemblages are cleaved by viral-encoded protease (PR) to generate three major products, matrix (MA), capsid (CA), and nucleocapsid (NC). These are arranged to comprise a MA shell underlying the lipid envelope and a conical (e.g. HIV-1, simian immunodeficiency virus, EIAV, i.e. lentivirus subgroup) or spherical (e.g. avian sarcoma virus) CA core that encapsulates the genome, which is bound to NC (12–14). For all retroviruses the N-terminal 12–13 residues of CA, unstructured within Gag, refold into a β-hairpin structural motif stabilized by a Pro3-Asp51 (HIV-1) salt bridge (2). The salt bridge involving the conserved residue proline 1 is essential and any mutations affecting salt bridge stability will unfold the β-hairpin and cause defects in the assembled capsid that eventually result in noninfectious virions (2–7). Correct assembly of native CA proteins during viral maturation ensures protection and integrity of the packaged viral genome. After virion entry into another host cell, the capsid disassembles to allow its genomic RNA to be released and reverse transcribed. This step requires the structural integrity and correct assembly of CA proteins (15, 16).

Successful viral maturation therefore depends on folding of the β-hairpin motif and the reversible quaternary packing of CA proteins. A retroviral CA protein consists of two helical domains, the 17-kDa N-terminal (CA N) and the 9-kDa C-ter-
The Role of the β-Hairpin in EIAV Capsid Structure

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The β-hairpin motif is crucial in the context of high resolution CA quaternary structures. To understand the contribution of the β-hairpin motif to CA assembly, we applied solution NMR spectroscopy to study the CA protein from EIAV, a lentivirus sharing the similar conical capsid core (12) and assembly (43) structure as HIV-1. Different from HIV-1 CA, EIAV CA dimerizes much weaker in solution, which results in sharper NMR resonance peaks and is more suitable for study by solution NMR. Previous results on HIV-1 CA showed that any point mutation, deletion mutation, or N-terminal extension at Pro1 would affect the Pro1-Asp51 salt bridge and unfold the β-hairpin (2). The published crystal structure of EIAV-CA is a N-terminal 3-residue (Pro1-Met5) deletion mutant with an unfolded signature β-hairpin (44). Instead of using this deletion mutant, which lacks resonances from residues Pro1 to Met5, we studied a N-terminal histidine-tagged variant that could also unfold the β-hairpin structure, in addition to the wt full-length EIAV-CA, to identify the structural differences associated with the β-hairpin formation. Our results showed that the maturational refolded β-hairpin induces a coil-helix transition in residues N-terminal to helix α1, Pro17-Gly19. The same region was also identified to be the expanded interface for the sparse CA domain oligomerization. The Thr16-Gly19 sequence is highly conserved in HIV-1 CA as well, and Arg18 of HIV-1 CA has been demonstrated to be essential (45) and form the inner-core of the CA domain hexamer (38). Therefore we propose the function of refolding the β-hairpin, the prerequisite event for retroviral CA assembly, is to extend helix α1 at the N terminus to enhance CA oligomerization for assembly. Our data further suggested the CA domain oligomerization is kinetically slower and uncorrelated to the faster dimerization, previously not known in retroviral capsid assembly events (46).

EXPERIMENTAL PROCEDURES

Protein Purifications—The wt EIAV-CA protein consists of 230 residues and starts with proline (Fig. 1A). The cDNA encoding the protein was inserted upstream of the intein gene in the pTXB1 vector from New England Biolabs (Ipswich, MA). The EIAV-CA-pTXB1 construct was transformed into Escherichia coli host strain BL21-DE3. The EIAV-CA-intein fusion protein sequence and the DTT-induced cleavage site can be found in supplemental “Experimental Procedures” and “Results”. The cells were grown in LB medium at 37 °C to an A600 of 0.7 and induced with 0.5 mM isopropyl 1-thio-β-D-galactopyranoside. The expression went on for another 5 h. The E. coli cells were pelleted from a 4-liter culture, dissolved in 500 ml of lysis buffer (20 mM Tris, pH 8, 500 mM NaCl, and 1 mM EDTA), and then passed twice through a high-pressure homogenizer, EmulsiFlex-C3 from Avestin (Ottawa, Canada). The cell lysate was centrifuged for 1 h at 35,000 × g. The supernatant containing the CA-intein fusion proteins was loaded onto a 20-ml gravity column filled with chitin beads (New England Biolabs). The column was then flushed with 60 ml of lysis buffer supplemented with 50 mM DTT. The on-column thiol-induced cleavage took 2 days. The CA proteins were finally eluted and dialyzed into a low-salt buffer (20 mM Tris, pH 9, and 5 mM NaCl). All of the above purification steps were performed at 4 °C. Further purifications were performed on pre-packed FPLC anion exchange (Fast-flow DEAE) and gel filtration columns (Hi-Load Superdex-75) from GE Healthcare. The identity of the purified proteins was confirmed using LC-MS. The measured molecular mass of 26075.8 Da is close to the theoretical value of 26076.9 Da. Isoelectric focusing (IEF) kits from Invitrogen were used to characterize the purified CA proteins. For NMR studies, 2H/15N/13C triple-labeled or 2H/15N double-labeled CA proteins were expressed overnight at 30 °C in E. coli cells grown in 99.8% 2H2O prepared M9 minimal medium. 13NH4Cl and [13C]glucose were the sole nitrogen and carbon sources, respectively. All isotopes were from Cambridge Isotope Laboratories (Andover, MA).

In addition to the wt EIAV-CA, another construct expressing a N-terminal histidine-tagged EIAV-CA was cloned, which had extra amino acid residues of AHHHHHHG added onto the native EIAV-CA N-terminal sequence. The full sequence of the variant can be found in supplemental “Experimental Procedures” and “Results”. The variant His-EIAV-CA protein was purified using a nickel-agarose gravity column first followed by FPLC columns similar to those used for purification of the wt EIAV-CA. The isolate labeling and backbone resonance assignments were obtained as described for the wt protein.

NMR Spectroscopy and Resonance Assignment—A 0.15 mM 2H/15N/13C triple-labeled EIAV-CA sample was used for resonance assignment. The same buffer of 20 mM potassium Pi at pH 6.7, 0.01% NaN3, and 7%2H2O was used in nearly all NMR experiments. The protein concentration was measured using a UV absorbance in 6 M guanidinium HCl (ε280 = 22460 M−1 cm−1). Temperature for all NMR experiments was 27 °C. The backbone resonance assignments were obtained from transverse relaxation optimized spectroscopy (TROSY) (47, 48) version of three-dimensional experiments HNCO (49), HN(CA)CO (50), HNCACB, and HN(CO)CACB (51). The through space three-dimensional experiments of NOESY-TROSY-HSQC (52) and HMQC-NOE-TROSY-HSQC (in house written) were collected with a NOE mixing time of 150 ms. All data were collected on Bruker Avance 600, 800, and 900 spectrometers equipped with cryogenic probes and Z-axis pulse field gradients. Chemical shifts were externally referenced. The TROSY-measured amide 15N and 1H chemical shifts were offset to true chemical shift values by 15NH2/2. High resolution TROSY spectra, which contained 512 complex points on 15N dimension, were collected on four 15N/2H-labeled samples at concentrations ranging from 0.05 to 0.4 mM on a 900 MHz spectrometer. All NMR spectra were processed and analyzed using NMRPipe (53) and SPARKY 3 (T. D. Goddard).
and D. G. Kneller, University of California San Francisco), respectively. In-house written MATLAB (The Mathworks, MA) programs using the Simplex search algorithm were used to fit all NMR data with appropriate models. The chemical shift assignments of the wt and histidine-tagged variant EIAV-CA have been deposited into the Biological Magnetic Resonance Bank (BMRB) data base with entry numbers 18421 and 18815, respectively.

Residual Dipolar Coupling and Structure Refinement—Both high resolution TROSY and HSQC spectra were collected on samples of the 15N/2H-labeled wt or variant EIAV-CA proteins on a 900 MHz NMR spectrometer. The differences in peak locations along the 15N dimensions of the two spectra yielded half of the $J_\text{N-H}$ coupling constants. Repeating the same measurements on the sample with an added $5–10$ mg/ml of Pf-1 phage co-solvent (54) (ASLA Biotech) yielded summed coupling constants ($J_\text{N-H} + D_{\text{N-H}})/2$, where $D_{\text{N-H}}$ is the residual dipolar coupling (RDC) of N-H bonds. Subtraction and doubling yielded measured $D_{\text{N-H}}$ and experimental errors were taken from duplication reproducibility. A set of five alignment tensor parameters, alignment order $D_\alpha$, rhombicity $R$, and Euler angles $\alpha$, $\beta$, and $\gamma$, were used to fit measured $D_{\text{N-H}}$ to domain structural coordinates. The Q factor was calculated to check the agreement between crystal structure coordinates and $D_{\text{N-H}}$ measured in solution.

The refinement of the $\text{C}^\text{A}$ backbone structure included adding coordinates for missing residues and simulated annealing using XPLOR-NIH (55). In addition to force-field-related restraints within XPLOR-NIH (56), specific restraints are 90 RDCs, 155 amide $^{1}J_\text{H-15N}$ NOEs, 226 $\Phi/\Psi$ dihedral angles derived from TALOS-plus (57), 85 empirical H-bond restraints, and root mean square deviation penalties for deviation away from helical residue coordinates of the original crystal structure. The structure with the lowest energy out of 500 calculated structures was chosen to model the EIAV-CA quaternary structure.

Analytical Ultracentrifugation and Data Analysis—Sedimentation velocity experiments were performed on the wt EIAV-CA protein at 0.4 mM concentrations and pH values of 4, 6, and 8. Buffer components were 100 mM NaCl for all pH values, and 20 mM NaAc for pH 4, 20 mM KPi for pH 6, and Tris-HCl for pH 8 (20 mM). A Beckman Optima XL-I analytical ultracentrifuge and a four-place AN-Ti rotor were used. Samples of 0.4 mM protein concentration were used. Centrifuge cells fitted with double-sector centerpieces and sapphire windows were filled with 0.4 ml of the protein sample and the dialysate buffer reference. After reaching thermal equilibrium at 20.0 °C at rest, the rotor was accelerated to 50,000 rpm. Interference and absorbance scans at 297 nm were started immediately after the rotor reached the set speed and collected until no further sedimentation boundary movement was observed. The apparent sedimentation coefficient distributions were analyzed by Lamm equation modeling using the SEDFIT software of Schuck (58). Positions of menisci and bottoms as well as fractional ratios were optimized during the fitting procedure. The final accepted fits had a root mean square deviation less than 0.006. To obtain information about the amount of oligomers present in samples, absorption and interference data were analyzed globally with SEDPHAT using the hybrid model. In this procedure, monomer, and dimer were fitted as global discrete species with fixed molecular weights and higher molecular weight species were modeled with a continuous sedimentation coefficient distribution.

RESULTS

Sequence Alignment Reveals Conserved Region at the N Terminal of Helix $\alpha$1—The three-dimensional structure-based sequence alignment between HIV-1-CA and EIAV-CA was performed on individual domains (Fig. 1A). The backbone heavy atom root mean square deviation between the two lentiviral CAs are 2.3 Å and 1.4 Å for domains $\text{C}^\text{A}$ and $\text{C}^\text{A}$, respectively. Both proteins share 29 and 57% sequence identity.
The Role of the β-Hairpin in EIAV Capsid Structure

and homology, respectively. The interesting conserved region is Thr<sup>16</sup>-Pro<sup>17</sup>-Arg<sup>18</sup>, corresponding to Ser<sup>16</sup>-Pro<sup>17</sup>-Arg<sup>18</sup> in HIV-1-CA (Fig. 1A), where we identified a conformational transition upon folding its upstream β-hairpin motif in the following NMR studies.

Figuring 2. The Cα/β secondary chemical shift indexes of wt (A) and variant (B) EIAV-CA at pH 6.7. ΔCα or ΔCβ was the difference between the measured <sup>13</sup>C chemical shifts and the values for the same amino acid in the unfolded state (59). The ΔCα,ΔCβ value at residue i was the average of the values at residues i-1, i, and i+1 (79).

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Folded the β-Hairpin Extends Helix α1 at Its N Terminus—
The protein carbon chemical shifts are sensitive to its local secondary structures (59). NMR resonance assignments were performed on <sup>2</sup>H/<sup>13</sup>C/<sup>15</sup>N-labeled wt EIAV-CA protein. At pH 6.7 a total of 186 backbone <sup>1</sup>H-<sup>15</sup>N resonances were assigned, representing 88% of 212 non-proline residues. Residues Ile<sup>2</sup>-Pro<sup>90</sup> corresponding to the cyclophilin A binding site on HIV-1-CA (23, 61), were in secondary structures (59). NMR resonance assignments were performed in parallel. Backbone resonances for nearly all of the residues Thr<sup>16</sup> and Thr<sup>22</sup> adopted helix N-capping (64) and helical conformations, respectively (Fig. 3A and Table 1). Residues between Thr<sup>16</sup> and Thr<sup>22</sup> in the wt CA have <sup>1</sup>H-<sup>15</sup>N resonances line broadening at pH 6.7, hence no δCα measurements. We lowered the pH to 5.2 to reduce line broadening due to any microsecond-millisecond exchange, and assigned additional <sup>1</sup>H-<sup>15</sup>N resonances for Arg<sup>18</sup>, Gly<sup>19</sup>, and Thr<sup>21</sup>. The comparison between δCα of the wt at pH 5.2 and the variant (Fig. 3B) clearly showed the higher helical content for residues Pro<sup>17</sup>, Arg<sup>18</sup>, and Gly<sup>19</sup> within the wt EIAV-CA. Normally an α-helix structure features a stretch of successive strong positive (>1 ppm) secondary Cα chemical shifts (ΔCα). Shown in Table 1 are the ΔCα of residues from Leu<sup>15</sup> to Thr<sup>22</sup>, which showed that helix α1 starts at Pro<sup>17</sup> and Tyr<sup>20</sup> for the wt and variant EIAV-CA, respectively, and overall the wt has a larger ΔCα value, meaning more stable helical structure. In addition, the program TALOS+ (57) predicted the wt Thr<sup>16</sup> dihedral angles Φ and Ψ to be −91 ± 43° and 147.5 ± 29.5°, respectively, which were close to the typical values predicted for helix N-capping residues, −94 ± 15° for Φ and 167 ± 5° for Ψ (65). The wt EIAV-CA therefore has Thr<sup>16</sup> to cap helix α1 that starts at Pro<sup>17</sup>.

The N-terminal extension of helix α1 can be visualized by aligning crystal structures of the native HIV-1-CA<sub>N</sub> (23) and the (Pro<sup>-</sup>-Met<sup>3</sup>)-deletion mutant of EIAV-CA<sub>N</sub> (44) (Fig. 3C). The Pro<sup>1</sup>-Asp<sup>3</sup> salt bridge stabilizes the β-hairpin motif, which in turn stabilizes and extends helix α1 from Tyr<sup>20</sup> to Pro<sup>17</sup>.

Oligomeric Interfaces Include the Extended Helix α1—For HIV-1-CA only at pH values slightly above its pl of 6.6 (17, 66) it could assemble to a mature-like capsid particle (19). In line with such observations we have performed most solution NMR measurements on EIAV-CA at pH 6.7, above its pl value of 6.4, to maximize chances in observing EIAV-CA oligomerization using <sup>1</sup>H-<sup>15</sup>N chemical shift mapping.

For both the wt and variant EIAV-CA high resolution TROSY <sup>1</sup>H-<sup>15</sup>N spectra were collected at pH 6.7 and concentrations of 0.05 and 0.2 mM. If no oligomerization is present...
The Role of the β-Hairpin in EIAV Capsid Structure

TABLE 1
Secondary $^{13}$Cα chemical shift values ($\Delta$Cα, defined in the legend to Fig. 2) of helix α1 N-terminal residues

| Residue | Wild-type $\Delta$Cα | Variant $\Delta$Cα |
|---------|----------------------|-------------------|
|         | pH 6.7,$^{a,b}$ ppm | pH 5.2,$^{a,b}$ ppm |
|         |                      |                   |
| Leu$^{15}$ | -0.01 (55.09)         | -0.02 (55.12)     |
| Thr$^{16}$ | -2.48 (59.62)         | -2.47 (59.63)     |
| Pro$^{17}$ | NA                   | -3.98 (58.12)     |
| Arg$^{18}$ | NA                   | -1.03 (62.07)     |
| Gly$^{19}$ | NA                   | 1.05 (37.15)      |
| Tyr$^{20}$ | NA                   | 0.34 (45.44)      |
| Thr$^{21}$ | NA                   | 3.74 (61.84)      |
| Thr$^{22}$ | 4.54 (66.64)         | 4.03 (66.13)      |
|         | 4.39 (66.49)         | 3.63 (65.73)      |

$^{a}$ Shown in the parentheses are the measured chemical shift values of nuclei $^{13}$Cα.
$^{b}$ Missing chemical shift values were due to $^1$H-$^1$5N line broadening that prohibited $^{13}$Cα assignments.
$^{c}$ NA, not applicable.

FIGURE 3. The N-terminal extension of helix α1 based on Cα chemical shifts ($\delta$Cα). The $\delta$Cα correlations between the wt and variant EIAV-CA$^{N}$ were shown with $\delta$Cα values of the wt EIAV-CA measured at pH 6.7 (A) and 5.2 (B). Residues with significant $\delta$Cα differences (> mean + 1.5 S.D. or < mean - 1.5 S.D.) between the wt and variant are indicated (A and B). C, the overlay of the three-dimensional crystal structures of native HIV-1-CA$^{N}$ (PDB code 1AK4, blue) domain and the (Pro$^{1}$-Met$^{3}$)-deletion mutant EIAV-CA$^{N}$ domain (PDB ID 2EIA, red). The salt bridge of HIV-1-CA$^{N}$ involving the nitrogen atom of Pro$^{1}$ and one side chain oxygen atom of Asp$^{31}$ was shown in magenta and cyan balls, respectively. The difference in the N-terminal residue of helix α1 was indicated.

there should not be any changes in the NMR chemical shifts as a function of protein concentration. Those residues that do show large chemical shift changes most likely will be located in the oligomeric interface. Chemical shift changes in backbone $^1$H-$^1$5N resonances ($\Delta$N,H) were normalized and from that a total of four possible oligomeric interfaces were identified (Fig. 4). The wt and variant proteins had nearly identical perturbation profiles for domain CA$^{C}$, which clustered around the two interfaces, the domain linker, e.g., Asn$^{152}$, and helix α9, i.e., Ile$^{181}$ and Thr$^{188}$. Thus folding the β-hairpin did not affect oligomerization of domain CA$^{C}$, and such oligomerization is in fast nanosecond-microsecond exchange kinetics.

The other two interfaces on domain CA$^{N}$ were similar between the wt and variant proteins in loci but different in boundaries when the un-assigned exchange-broadened residues were taken into account. Specifically the variant EIAV-CA showed strong chemical shift perturbation around Thr$^{21}$, close to the N terminus of its helix α1 (Tyr$^{20}$) (Fig. 4B); in comparison, wt EIAV-CA showed chemical shift changes and severe line broadening for residues ranging from its helix α1 capping residue Thr$^{16}$ up to Trp$^{23}$ (Fig. 4A) in the middle of the helix. This interface boundary distinction is correlated to the difference in N termini of helix α1, Pro$^{17}$ versus Tyr$^{20}$ in the wt and variant proteins, respectively (Table 1). It is conceivable that only helical residues could participate in the oligomeric interface along helix α1. The other CA$^{N}$ interface is the region spanning helices α2 and α3. Residue Leu$^{40}$, in the middle of helix α2, was readily identified in chemical shift perturbation for both the wt and variant proteins. For the variant residue Leu$^{44}$ and its neighboring exchange, broadened residues Val$^{46}$-Cys$^{48}$ were interface residues (Fig. 4B). For the wt the exchange broadening was expanded up to Asn$^{54}$ (Fig. 4A), in the middle of helix α3. Overall folding of the β-hairpin expanded the domain CA$^{N}$ oligomeric interfaces in helices α1–3 and caused slower exchange kinetics for interface residues, including those that are too severely broadened to be observed (Fig. 4 and supplemental Fig. S1). The $^1$H-$^1$5N exchange broadening in domain CA$^{N}$ (supplemental Fig. S1) caused by microsecond-millisecond kinetics usually reflects the existence of sparsely populated excited states potentially carrying biological interests (60, 67, 68), e.g. the protein folding and unfolding in populations as small as 1% (69). Formation of the N-terminal β-hairpin has
been shown to stabilize the structure of the CA^N domain (70). Therefore, increased line broadening in the presence of the β-hairpin is less likely due to intramolecular processes. It is also worth noting that the pH change from 6.7 to 5.2 did not alter the chemical shifts for the wt EIAV-CA in the region, e.g. Leu^15, Thr^16, and Thr^22 (Table 1), which indicated that the microsecond-millisecond exchange broadening at pH 6.7 for ^1H-^15N resonances was not due to any local structural variations, but to changes in solvent exposure or packing contact due to a shift in the oligomerization equilibrium.

The Monomer-Oligomer Equilibrium—Because the chemical shift mapping and resonance line broadening suggested the presence of both CA^N- and CA^C-mediated oligomerization at pH 6.7, it would be ideal to cross-validate such oligomers at this pH. The sedimentation velocity profiles of the wt EIAV-CA at pH 4, 6, and 8 all showed one major monomeric species that banded at 2.4 S. A secondary peak at 3.5 S, corresponding to the dimer species, was observed at pH 6 (Fig. 5). An incomplete separation of the dimer peak at pH 4, between 2.5 and 3.0 S, suggested fast dissociation and instability of the dimer. The partial CA dimerization is common among retroviral CAs except for the CAs from primate lentiviruses such as HIV-1 that fully dimerize in solution.

The EIAV-CA^N driven oligomer was not detected in analytical ultracentrifugation experiments due to its sparse population, below the general 2% detection limit of analytical ultracentrifugation measurement. Similarly for HIV-1-CA in solution, only the cross-linking method, not sedimentation, could identify oligomers larger than a dimer (17). In fact all crystal structures of CA^N-oligomerized HIV-1-CA were obtained on mutants or specially treated proteins that favor oligomerization (39).

Folding of the β-Hairpin Induces Different Domain Alignment and Oligomerization—For solution NMR studies it is important to validate the overall consistency between protein solution and crystal structures by fitting NMR-measured RDCs to existing crystal structure coordinates (71, 72). A total of 150 resolvable backbone ^1H-^15N RDCs were measured on a 0.15 mM wt sample at pH 6.7. RDC fittings were performed on the individual domains instead. For the CAN and CAC domains, reasonable agreements were obtained, as indicated by Q factors of 31 and 22%, respectively, using the coordinates of chain B in the crystal structure (44) (Table 2 and Fig. 6). Because the crystal structure missed the β-hairpin, refinement of the backbone structure coordinates for domain CA^N was carried out using XPLOR-NIH (55). After
TABLE 2
The domain specific alignment tensors derived from backbone amide 
$^{1}H-^{15}N$ RDCs measured on both the wild-type and the variant EIAV-CA

| Protein   | Domain | $D_{N\cdot H}$ | $R$ | $\alpha$ | $\beta$ | $\gamma$ | $Q^a$ |
|-----------|--------|----------------|-----|-----------|---------|----------|-------|
| Wild-type | CAN$^b$ | −19.6          | 0.081 | 25.0 | 126 | 136 | 31.0% |
|           | CAN$^c$ | −20.5          | 0.0154 | 38.2 | 118 | 14.8 | 11.6% |
|           | CAN$^b$ | 4.71           | 0.397 | 64.1 | 18.7 | 37 | 21.9% |
| Variant   | CAN$^b$ | −17.2          | 0.320 | 25.0 | 117 | 124 | 40.8% |

$^a$ The $Q$ factor was calculated according to the published method (76).

$^b$ The molecular coordinates were from chain b of the EIAV-CA crystal structure (PDB code 2EIA) (44).

$^c$ The molecular coordinates were from the backbone structure refinement.

FIGURE 6. The correlation between the measured backbone $^{1}H-^{15}N$ RDCs and calculated values for domains CAN (A) and CAC (B) of the wt EIAV-CA at pH 6.7. The circles are fits using the crystal structure coordinate. The triangles are the fits using the refined CAN domain coordinates.

this further refinement, the $Q$ factor for the CAN dropped to 12% (Table 2 and Fig. 6).

In addition to validating the domain structure, the $D_{N\cdot H}$ determined alignment tensors could qualitatively report inter-domain motion and domain oligomerization. Larger oligomers, ready to be aligned, contribute more into the observed tensor order ($D_{O}$) and rhombicity ($R$). The presence of inter-domain motions would introduce a difference in $D_{O}$ and $R$ between the two domains. The wt EIAV-CA exhibited the opposite signs of $D_{O}$ with a difference of $−4.4$-fold and $R$ difference of 0.38 (theoretical maximum value of $R$ is 0.67) between CAN and CAC domains (Table 2), which demonstrated that individual domains participated in two different oligomers that involve significant changes in inter-domain orientation. Measurements on the EIAV-CA variant showed the alignment orders were in the same negative sign with a difference of only 1.6-fold, roughly scale with their size ratio, and a rhombicity $R$ difference of 0.14 (Table 2), both differences were much smaller than the wt values, which indicated the oligomerization within the variant monomer-oligomer equilibriums did not change the inter-domain orientation significantly. Based on the $^{1}H-^{15}N$ chemical shift mapping profiles (Fig. 4) the CAC driven oligomerizations were nearly identical for both the wt and variant. Therefore the CAN oligomerization difference should be the dominant cause for domain tensor parameter differences. The close to zero rhombicity of the wt domain CAN$^c$ suggest the presence of CAN$^c$ driven symmetric oligomers, whereas the variant CAN$^c$ domain oligomer lacks symmetry with a medium value for rhombicity of 0.32, indicating less CAN$^c$ oligomers were formed in the absence of the $\beta$-hairpin. This is consistent with the more exchange-broadened residues in the wt CAN oligomeric interfaces (Fig. 4 and supplemental Fig. S1).

DISCUSSION

Here we have studied the EIAV-CA protein, the first lentiviral CA protein characterized using solution NMR in its native full-length sequence. The NMR data suggested a mechanism for the initial EIAV-CA assembly in solution, i.e., upon Gag processing by viral PR, the folding of the $\beta$-hairpin structure extends the N terminus of helix $\alpha_1$, the highly conserved Thr/Ser$^{16}$-Pro$^{17}$-Arg$^{18}$, where the CA core assembly initiates. Such a core interface is within a sparse population of the slower forming domain CAN$^c$ driven oligomer. The conclusion derived from our NMR results is consistent with previously published data.

Cross-validations on Enhanced/Extended Helix $\alpha_1$ Upon Folding of the $\beta$-Hairpin—Our $^{13}Ca$ chemical shift definitions of the N terminus of helix $\alpha_1$ of EIAV-CA in solution were in line with the published crystallography results. Specifically the crystal structure of the HIV-1-CAN domain (23) with the $\beta$-hairpin have helix $\alpha_1$ starting from residue Pro$^{17}$ and the side chain of Ser$^{16}$ forming a hydrogen bond with the amide of Thr$^{19}$ to cap helix $\alpha_1$. In contrast, the crystal structure of the (Pro$^1$-Met$^3$)-deletion mutant EIAV-CA (44), which is missing the $\beta$-hairpin, has a shorter helix $\alpha_1$ starting at Tyr$^{20}$, not Pro$^{17}$ (Fig. 3C).

Recent hydrogen-deuterium exchange mass spectrometry identified stronger protection in HIV-1-CA helices $\alpha_1$–3 in the presence of the $\beta$-hairpin and suggested helix $\alpha_1$ could be re-oriented upon maturation (73). Stronger protection is consistent with the enhanced helix stability.

Extended Helix $\alpha_1$ Constitutes the Essential CA Assembly Core—Early studies of HIV-1-CA assembly suggested that refolding of the $\beta$-hairpin created a new CA-CA interface com-
The Role of the β-Hairpin in EIAV Capsid Structure

FIGURE 7. Models for hexameric (A and C) and dimeric (B) interfaces of EIAV-CA. The side chain bonds of chemical shift mapped residues were colored blue in both hexamer (A) and dimer (B) models, which were generated using rigid body rotation of the refined EIAV-CA hexamer crystal coordinates onto the crystal structure of HIV-1-CA hexamer (PDB code 3GV2)(39) and the solution structure of HIV-1-CA dimer (PDB code 2KOD)(41), respectively. One EIAV-CA molecule was colored red in both oligomer models. The cyan colored ribbons correspond to residues with severe 1H−15N line broadening. C, a close view of the hexamer core. The side chain bonds of the most-inner core residue Arg18 were colored green.

petent for hexamer formation (2, 74). One hypothesis was that residue Pro17 forms the new interface (74). Recently a total of three HIV-1-CA assembly interfaces were identified (38), which are helices α1–3, formed a 18-helix bundle of the CAN hexamer core, the CA dimerization along helix α9, and the inter-molecular and inter-domain contact between helices α4 of the CAN and α10 of the neighboring CA. Among them residue Arg18 of helix α1 was located in the deepest center core of the HIV-1-CA hexamer interface (Fig. 7A). Arg18 is a determinant in HIV-1-CA hexamerization (45), e.g. the R18L mutant increases the hexamer population for crystallization (38). All the above evidences from HIV-1-CA are completely in line with our EIAV-CA NMR data showing an expanded oligomeric assembly phenotype from tubular to spherical particles (74).

In addition to identifying the β-hairpin induced structural effects, our NMR data suggested the two CA domains independently drive oligomerization, i.e. the readily formed domain CA dimerization along helix α9 (Fig. 7B) is in fast kinetics (supplemental Fig. S1), in contrast, the domain CAN has to structure the helix α1 to position the side chain of its Arg18 toward the inner core of the hexamer (Fig. 7C), overcoming a repulsion energy barrier and thus kinetically slower (supplemental Fig. S1). Therefore it is not surprising that the CAN-driven oligomer at the experimental sub-millimolar concentration was a sparse population (probably <1%). However, the CA assembly could still be efficient within a retrovirus where the CA concentration is much higher (75).

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The Role of the β-Hairpin in EIAV Capsid Structure

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