Structural determinants of coiled coil mechanics

Patricia López-García, a Melis Goktas, a Ana E. Bergues-Pupo, a Beate Koksch, c Daniel Varón Silva b,c and Kerstin G. Blank a

The natural abundance of coiled coil (CC) motifs in the cytoskeleton and the extracellular matrix suggests that CCs play a crucial role in the bidirectional mechanochemical signaling between cells and the matrix. Their functional importance and structural simplicity has allowed the development of numerous applications, such as protein-origami structures, drug delivery systems and biomaterials. With the goal of establishing CCs as nanomechanical building blocks, we investigated the importance of helix propensity and hydrophobic core packing on the mechanical stability of 4-heptad CC heterodimers. Using single-molecule force spectroscopy, we show that both parameters determine the force-induced dissociation in shear loading geometry; however, with different effects on the energy landscape. Decreasing the helix propensity lowers the transition barrier height, leading to a concomitant decrease in the distance to the transition state. In contrast, a less tightly packed hydrophobic core increases the distance to the transition state. We propose that this originates from a larger side chain dynamics, possible water intrusion at the interface as well as differences in solvation of the hydrophobic amino acids at the transition state. In conclusion, the different contributions of helix propensity and hydrophobic core packing need to be considered when tuning the mechanical properties of CCs for applications.

Coiled coils (CCs) are self-assembled, superhelical motifs that are naturally found in numerous proteins in the cytoskeleton and the extracellular matrix. CCs consist of two (or more) α-helices, each characterized by a repetitive pattern of seven amino acids, called heptad (abcdefg)n (Fig. 1a). Positions a and d form the hydrophobic core of the superhelical structure; e and g are mostly charged amino acids, which participate in interhelical salt bridges; b, c and f are solvent-exposed, often polar amino acids, which contribute to the helix propensity of the individual helices.

Utilizing this simple design, CCs serve as model systems for studying protein folding and stability. As a result, they are increasingly used as templates for protein design and sequences with a pre-determined thermodynamic stability can now be synthesized de novo. Such sequences find application in peptide-based hydrogels and protein origami structures as well as in biosensors and drug delivery systems. Considering their natural role as a mechanical scaffold, surprisingly little information is available about the sequence-structure-mechanics relationship of CCs. With the goal of introducing CCs as nanomechanical building blocks, we have characterized three different CC heterodimers with atomic force microscopy (AFM)-based single-molecule force spectroscopy (SMFS; Fig. 1b). We show that hydrophobic core packing and helix propensity
affect the thermodynamic stability in similar ways; however, the underlying changes to the energy landscape are different.

Using the thermodynamically and mechanically well-characterized CC A₄B₄ as the starting point (Fig. 1c),⁷,¹⁶ we used the standard rules of CC design⁵,⁷ to obtain one sequence with a reduced helix propensity and a second sequence with a different hydrophobic core packing. Throughout the manuscript, we define hydrophobic core packing as the combination of hydrophobicity and side chain packing of the amino acids in positions a and d. To reduce the helix propensity, Ala in position b was substituted with Ser in all heptads¹⁷ (A₄B₄S; Fig. 1c). Hydrophobic core packing was altered using another β-branched amino acid in position a (Val instead of Ile; A₄V₄B₄V).

The Asn in the third heptad was not replaced to maintain heterospecificity.¹⁸,¹⁹ For A₄V₄B₄V, it needs to be considered that this substitution also decreases the helix propensity, which is lower for Val than for Ile (Fig. 1c). To define the points of force application, Cys was introduced at the desired termini for coupling the CC to the surface and the AFM cantilever. For A-peptides, Cys was located at the N-terminus, while it was placed at the C-terminus of B-peptides, thus establishing a shear pulling geometry (Fig. 1b).

To validate the design and to compare mechanical and thermodynamic stability, the CCs were first investigated with circular dichromism (CD) spectroscopy. Wavelength scans, showing an α-helical signature with minima at 208 nm and 222 nm and thermal denaturation experiments are displayed in Fig. S1 (ESI†). As expected, A₄B₄ shows the highest melting temperature Tₘ, while A₄B₄S and A₄V₄B₄V were significantly destabilized (Table 1). The corresponding free energy difference between the folded (F) and the unfolded (U) state (ΔGₐₔ) was obtained from van ’t Hoff plots (Fig. S2, ESI†). In comparison to A₄B₄, both modified CCs also show a lower ΔGₐₔ, which follows a similar trend compared to other CCs reported in the literature.⁵,²⁰

|          | Tₘ (°C) | ΔGₐₔ (kJ/mol) | Fₚ (pN) | Δxₚₜₛ (nm) | kₐₐₕ (s⁻¹) | ΔGₜₛₜₛ (kJ/mol) |
|----------|---------|---------------|---------|------------|-------------|-----------------|
| A₄B₄     | 77.0 ± 0.3 | 14.2 ± 0.3   | 43.1 ± 0.2 | 1.3 ± 0.2  | (3.2 ± 2.1) x 10⁻⁴ | 29.2 ± 1.4      |
| A₄B₄S    | 54.3 ± 0.3 | 5.3 ± 0.2    | 23.6 ± 4.8 | 0.9 ± 0.1  | (2.8 ± 1.1) x 10⁻⁴ | 21.3 ± 0.2      |
| A₄V₄B₄V  | 59.0 ± 0.6 | 7.1 ± 0.6    | 28.2 ± 0.1 | 1.7 ± 0.0  | (2.4 ± 1.7) x 10⁻₃ | 26.6 ± 0.7      |

Most probable rupture force Fₚ determined at a retract speed of 400 nm s⁻¹. All values are depicted as mean ± standard error of the mean (SEM).

This provides a first indication that the substitutions also lower the mechanical stability of the CCs.

Subsequent dynamic SMFS, performed over a range of loading rates (r = dF/dt)²¹ from approximately 15 pN s⁻¹ to 110 000 pN s⁻¹, revealed that both modified CCs possess a lower mechanical stability over the complete range of loading rates (Fig. S4–S6, ESI†); however, their dependence on the loading rate is different. Fitting the data to the Bell-Evans model²¹ (Fig. 2b) allows for obtaining more detailed information about the energy landscape of the CC two-state system (folded CC vs. random coil peptides): kₐₐₕ, the force-free dissociation rate, and Δxₚₜₛ, the distance from the folded to the transition state (TS). Using the Arrhenius equation, also the barrier height between the folded and the transition state (ΔGₜₛₜₛ) can be calculated, provided that the Arrhenius pre-factor is known. Here, we use an Arrhenius pre-factor of 5 x 10⁹ s⁻¹, which was estimated for the dimeric GCN4 leucine zipper.²² Table 1 summarizes the obtained fit values, as well as the calculated ΔGₜₛₜₛ values.

In addition to the results shown in Table 1, where the mean ± standard error of the mean (SEM) was calculated from three independent experiments, a Peacock test²³ was performed to determine if the modified CCs A₄S₄B₄S and A₄V₄B₄V are significantly different from the reference sequence A₄B₄. This test shows p-values smaller than 0.01 (see ESI† for details).

Comparing the kₐₐₕ values presented in Table 1 shows that both modifications (A₄S₄B₄S and A₄V₄B₄V) lower the height of the transition state barrier with respect to the reference sequence A₄B₄ (higher kₐₐₕ and lower ΔGₜₛₜₛ). This lower barrier height is correlated with the different thermodynamic stabilities (Table 1) and helix propensities of the three CCs (Fig. 1c). This suggests that a reduced helix propensity lowers the barrier height, thereby affecting both the thermodynamic and mechanical stability of the CCs. Interestingly, the Δxₚₜₛ values do not correlate with the thermodynamic stabilities. The Ala-Ser modification (A₄S₄B₄S) reduces Δxₚₜₛ whereas the Ile-Val modification (A₄V₄B₄V) increases Δxₚₜₛ. This suggests that modifications in the solvent-exposed residues affect the energy landscape of the CC interaction differently when compared to hydrophobic core modifications.

In contrast to cooperative dissociation and unfolding upon increasing temperature, CCs respond to an applied axial stretching force in three phases.¹⁶,²⁴–²⁶ Initially, the force increases linearly with extension and the helices remain intact (phase I). At a strain of 10–20%, the individual helices start uncoiling at an almost constant force (phase II). In long CCs, the force increases sharply after the helices are uncoiled and the resulting structure is extended further (phase III). For CCs
destabilizes an already less stable CC further, an effect that was
stability. Uncoiling of only small parts of helical structure thus
a lower helix propensity is correlated with a lower thermodynamic
less force is required to uncoil the individual helices. In addition,
CCs to shear forces. For CCs with a lower overall helix propensity
phase II. This is a direct result of the force-induced chain
separation mechanism. At loading rates typically used in SMFS,
the applied force causes the uncoiling of helical structure at the
points of force application. This, in turn, destabilizes the CC
thermodynamically and facilitates the subsequent dissociation
of the CC chains (uncoiling-assisted dissociation). This mecha-
ism allows for explaining the effects of helix propensity and
hydrophobic core packing on CC shearing.

When mechanically loaded in shear geometry, the hydro-
gen bonds stabilizing the individual helices are aligned parallel
to the force vector, whereas the hydrophobic side chains are
arranged almost perpendicularly. The torsional angles and
helical propensities of the individual amino acids, which are
responsible for maintaining stable hydrogen bonds in the
helices, are thus critically determining the resistance of
CCs to shear forces. For CCs with a lower overall helix propensity
less force is required to uncoil the individual helices. In addition,

a lower helix propensity is correlated with a lower thermodynamic
stability. Uncoiling of only small parts of helical structure thus
destabilizes an already less stable CC further, an effect that was
observed earlier when decreasing CC length. Assuming that the
hydrophobic core is not altered, this suggests that chain separa-
tion occurs at smaller extensions. In the case of A_{4S}B_{4S}, a higher
k_{off} value is thus accompanied by a shorter \( \Delta x_{F-TS} \). This result is in
line with the observation that artificial constraints, which stabilize
the helices against uncoiling, lead to an increase in the forces
required for chain separation.\(^{3,11,29}\)

Following this line of argumentation, \( \Delta x_{F-TS} \) for A_{4V}B_{4V}
is expected to lie in between the values obtained for A_{4S}B_{4S}
and A_{4B}; however, A_{4V}B_{4V} shows an increase in \( \Delta x_{F-TS} \). This
suggests that substituting Ile with Val does not only affect the helix propensity. We propose that the increased \( \Delta x_{F-TS} \) originates from the combined effect of side chain hydrophobicity
and packing. A reduced packing is expected to increase side chain
dynamics at the hydrophobic interface, thus facilitating the
rearrangement of the Val side chains in response to the applied
force. In other words, a less densely packed hydrophobic core may
allow the relative displacement of the helices prior to or during helix uncoiling. At the same time, changes in side chain hydro-
phobicity affect the interaction of the CC structure with water.
Shear deformation may lead to water intrusion at the interface
(wetting), followed by force-induced helix uncoiling. Alternatively,
the helices may unfold without significant wetting. Independent
of the presence of wetting, helix unfolding exposes hydro-
phobic amino acids to solvent, which need to be hydrated
(solvation).\(^{20,30–33}\) While both wetting and solvation are primarily
determined by side chain hydrophobicity, also the local
environment\(^{11,32,34,35}\) and protein flexibility\(^{36}\) are expected to play
important roles. As the two-state energy landscape model cond-
enses the contribution of all these factors onto a single reaction
coordinate (Fig. 3), it remains an open question which factor(s)
determine the observed change in \( \Delta x_{F-TS} \).

An increase in \( \Delta x_{F-TS} \) has also been observed when introdu-
cing destabilizing substitutions in the hydrophobic core of the
globular proteins GB1 and protein L.\(^{37,38}\) This effect, termed
mechanical softening, was absent for the titin I27 domain,
however, where mechanical unfolding is determined by

a number of key hydrogen bonds.\(^{39,40}\) This suggests that

![](image)

**Fig. 2** Single-molecule force spectroscopy. (a) Representative rupture force
histograms obtained at a retract speed of 400 nm \( s^{-1} \) with \( n = 284 \) \( A_{4B} \),
\( n = 243 \) \( A_{4S}B_{4S} \) and \( n = 420 \) \( A_{4V}B_{4V} \). The dashed lines show Gaussian fits,
applied to extract the most probable rupture forces. The inset shows
representative force–extension curves for each CC. (b) Dynamic SMFS plot.
Each CC was measured in triplicate using a different cantilever and surface
different shades of the same colour). The solid lines represent fits to the
Bell–Evans model.

![](image)

**Fig. 3** Energy landscape of the CCs. The horizontal line represents the
distance from the folded (F) to the transition state (TS) \( \Delta x_{F-TS} \), while the
vertical solid arrow represents the transition barrier height \( \Delta \xi_{F-TS} \).
The dotted arrow shows the energy difference between the folded and
the unfolded state \( \Delta G_{F-U} \).
substitutions in the hydrophobic core only affect the energy landscape parameters when positioned at a mechanically loaded interface. Clearly, CCs represent an attractive model system to dissect the different contributions of hydrophobicity and side chain packing to the mechanics of hydrophobic interfaces, e.g. via temperature-dependent SMFS experiments.

In summary, the combination of helix propensity and hydrophobic core packing determines the mechanical stability of CCs; however, with different effects on the energy landscape (Fig. 3). Whereas a reduced helix propensity decreases both the barrier height and the distance to the transition state, an increase in the transition state distance is obtained when decreasing the hydrophobic core packing. Clearly, both parameters can be utilized when designing CCs with controlled mechanical properties for future applications. Most interestingly, these parameters can be used to obtain CCs with similar thermodynamic stabilities that possess a different dynamic response to an externally applied force. Such systems will find application as molecular force sensors\(^1\) or as physical hydrogel crosslinks, which show a pre-defined response to mechanical deformation.

**Conflicts of interest**

There are no conflicts to declare.

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