Destabilizing an interacting motif strengthens the association of a designed ankyrin repeat protein with tubulin

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Affinity maturation by random mutagenesis and selection is an established technique to make binding molecules more suitable for applications in biomedical research, diagnostics and therapy. Here we identified an unexpected novel mechanism of affinity increase upon in vitro evolution of a tubulin-specific designed ankyrin repeat protein (DARPin). Structural analysis indicated that in the progenitor DARPin the C-terminal capping repeat (C-cap) undergoes a 25° rotation to avoid a clash with tubulin upon binding. Additionally, the C-cap appears to be involved in electrostatic repulsion with tubulin. Biochemical and structural characterizations demonstrated that the evolved mutants achieved a gain in affinity through destabilization of the C-cap, which relieves the need of a DARPin conformational change upon tubulin binding and removes unfavorable interactions in the complex. Therefore, this specific case of an order-to-disorder transition led to a 100-fold tighter complex with a subnanomolar equilibrium dissociation constant, remarkably associated with a 30% decrease of the binding surface.

The selection of binders for applications where high affinity and specificity are needed has long been based on the use of the immune system. In the last ca. 15 years, as an alternative to natural and synthetic antibodies, libraries of artificial proteins have been developed as a resource of potential binders¹,². These libraries are typically based on a conserved, well-defined, protein scaffold and on a variable binding surface. Such binders often extend the possibilities of antibodies, e.g. with cysteine-free scaffolds that are able to work under reducing conditions. Using molecular display technologies (e.g. ribosome display), binders for a particular target are selected from the library. Once first generation binders have been identified, steps of diversification and selection may be applied to obtain higher affinity ones¹, a process reminiscent of the maturation of antibodies during the immune response.

The binding of a ligand, where the ligand may be a macromolecule (protein, nucleic acid…) or a small molecule, can affect a protein in many ways¹. In addition to small local changes, domain movements may occur in an induced fit mechanism. More dramatic rearrangements like disorder-to-order transitions are also frequently observed¹. The reverse case, an order-to-disorder transition, has also been documented⁶. In this case, the binding surface is typically hidden in the folded protein and becomes more accessible after melting of a secondary structural element or of a domain⁶. As to the unfolding of a protein motif that is part of a protein—ligand interface, it is expected to lead to the destabilization of the assembly.

In contrast to this last expectation, we report a case where a gain in affinity results from the destabilization of a two-helix-containing motif. We previously selected artificial binders to the αβ tubulin heterodimer (tubulin) from a library of designed ankyrin repeat proteins (DARPins)⁸,⁹. Here we applied cycles of randomization alternately with selection steps in order to identify higher affinity binders. This strategy led to evolved DARPins that

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bind tubulin two orders of magnitude stronger than the parental one, resulting in complexes with subnanomolar dissociation constant. Biochemical and structural characterizations demonstrated that the affinity increase is coupled with the destabilization of the ankyrin C-terminal capping motif (C-cap) of the DARPin, which in the parent DARPin interacts with tubulin but needs to undergo a rotation to avoid a clash. This remarkable order-to-disorder transition illustrates a new mechanism for affinity maturation.

**Results**

**DARPins with an improved affinity for tubulin.** In the course of the study of microtubules, we have selected DARPins that bind tubulin as multipurpose tools and in particular as crystallization chaperones. Whereas these DARPins have proven instrumental to crystallize tubulin and its complexes with interacting proteins (e.g. see ref. 10), the tubulin–DARPin affinity is within the range of values usually observed for interactions that are transient on a time scale of seconds to minutes, the measured equilibrium dissociation constant ($K_D$) being in the 100 nM range. Beyond their use as crystallization chaperones, which usually occur at high concentrations, other applications, for example in studies within living cells, would benefit from higher affinity binders.

To this end we evolved the tubulin-binding DARPin named D1 by applying cycles of error-prone PCR-based diversification followed by selection using ribosome display. The selected variants were further screened in an ELISA-based assay. In this assay, tubulin covalently linked to a biotinylated stathmin-like peptide was immobilized on a neutravidin-coated plate, which was then incubated with detached DARPins to immobilized tubulin. The DARPins were ranked as low-, medium- or high-affinity binders. D1 is the parent DARPin whereas off7 is an unrelated one, not binding to tubulin and used as a negative control. Error bars are standard deviations from duplicate experiments. AU, absorbance unit.

![ELISA analysis of D1 variants after two rounds of error-prone PCR followed by selection using ribosome display.](image)

**Table 1.** Equilibrium and kinetic constants of tubulin complexes with DARPin D1 and its variants. n.d.: not determined.
Figure 2. The tubulin-DARPin interaction monitored by fluorescence spectroscopy and surface plasmon resonance (SPR). (a) Fluorescence variation of 100 nM (left) or 15 nM (right) acrylodan-labeled DARPin as a function of tubulin concentration. The curve is the fit of the experimental points with Equation 1, from which the $K_D$ is extracted. Error bars correspond to standard deviation from duplicate experiments. a.u., arbitrary units. (b) Dissociation of acrylodan-labeled DARPin from tubulin. In the case of D1 (left), 5.5 μM unlabeled D1 was added to a 100 nM labeled D1 and 0.5 μM tubulin mixture. The decrease in fluorescence signal was
monitored in a stopped-flow apparatus (30% of the data points are shown). In the case of A-C2 (right), 2 μM unlabeled A-C2 was added to a 20 nM labeled A-C2 and 40 nM tubulin mixture and the fluorescence signal was monitored in a spectrophotometer. The curve is the fit of the experimental points with a mono-exponential decay equation (Equation 2). (c) Determination of the association rate constant by fluorescence. Tubulin at the indicated concentrations was added to a fixed concentration (50 nM) of labeled D1 (left) or A-C2 (right) in a stopped-flow apparatus (20% of the experimental points are displayed). The data were fitted according to Equation 3. The variation of k_obs as a function of tubulin concentration is shown in inset, from which the kon (slope of the curve) is derived. (d) The tubulin-DARPin interaction monitored by SPR. D1 (left) and TM-3 (right) were immobilized through their His-tag on the sensor chip. Tubulin at the indicated concentration was applied at time zero for 60 s (D1) or 180 s (TM-3), followed by a washing buffer flow. The black curves are the fit of the experimental data using the Langmuir analysis, from which the kon and koff are extracted. In the case of D1, plotting the values at the plateau as a function of the tubulin concentration provided an estimate of the K_D (Supplementary Fig. 5) that is very similar to the k_off/k_on ratio.

Assessing the contribution of the residues mutated in A-C2. Several mechanisms may come into play to account for the affinity increase of a protein for a ligand. An obvious one is a substitution at the protein—ligand interface that strengthens the interactions with neighboring chemical partners. To determine the contribution of this mechanism to the affinity maturation process of D1, we compared the sequence of A-C2 with that of D1 (Fig. 3a). There were changes at 11 amino acid positions, two of them (I122V and N158S) concerning residues that interact directly with tubulin in the tubulin–D1 complex, taking a 4 Å distance cutoff (Fig. 3b). To investigate the contribution of these two residues as well as those of the other 9 more distant ones, we prepared 11 single mutants of D1, each having one of the 11 substitutions found in A-C2. We also prepared four double or triple mutants comprising mutations that are close in the sequence and are more likely to operate in synergy. We evaluated these constructs for their tubulin-binding ability by ELISA and compared them with D1 and A-C2 (Fig. 3c). The ELISA conditions were the same as those initially used to screen the randomized D1 variants, i.e. the DARPin variants which give the highest signal in the ELISA conditions. First, we quantified by fluorescence spectroscopy the interaction with tubulin of both D1 and A-C2 using acrylodan-labeled cysteine variants. The stepwise addition of tubulin to labeled DARPin led to an increase of the acrylodan fluorescence signal (Fig. 2a). Fitting the data with a quadratic equation (Equation 1, see “Methods”) yielded the value of K_D and indicated a change by two orders of magnitude of this value to less than 1 nM in the case of A-C2, down from ~125 nM in the case of D1 (Table 1), the latter value being in excellent agreement with the one determined previously by fluorescence anisotropy. We also determined the association (k_on) and dissociation (k_off) rate constants of the tubulin-DARPin complexes. A chase of labeled DARPin from tubulin by the unlabeled counterpart yielded the k_off value (Fig. 2b) whereas the k_on value was derived from association kinetics under pseudo-first-order conditions (Fig. 2c). In all cases, data were fitted with monoeXponential equations (see “Methods”) without any indication of more complex reaction schemes. As expected, the data indicated a dissociation of A-C2 from tubulin two orders of magnitude slower than that of the parent DARPin D1 (Table 1). In contrast, the k_on values of the tubulin complexes with A-C2 and with A-C2 were similar. This scheme is typical for affinity maturation processes and results in this case in a subnanomolar K_D.

Structural investigation of the affinity maturation process. Since the substitutions at the protein—ligand interface did not seem to have much of an effect on dissociation rate, another mechanism may account for the higher affinity of an evolved binder, constituting the stabilization of the DARPin's unbound form in a...
“ready-to-interact” conformation, the one adopted in the complex. This scenario implies that the structure of the isolated parent binder is not optimized for interaction. In this case either an induced fit occurs upon ligand binding or different conformations of the binder coexist and only a sparsely populated one is able to interact. To investigate this possibility, and despite the fact that DARPins are generally very rigid proteins, we first determined the crystal structure of uncomplexed D1 (Supplementary Table 1) and compared it with that of D1 in complex with tubulin. This comparison indicated that this DARPin indeed changes conformation upon binding to the ligand.

Figure 3. Assessing the contribution of the D1 residues mutated in A-C2. (a) Sequence alignment of D1, A-C2 and the optimized TM-3 DARPin (color code: N-cap, green; 1st internal repeat, bright blue; 2nd internal repeat, dark blue; 3rd internal repeat, cyan; C-cap, yellow; mutated residues relative to D1, red). (b) Structure of tubulin–D1 with the Cα position of the residues mutated in A-C2 highlighted as red spheres. D1 is colored according to panel A. (c) ELISA analysis (binding and off-rate estimation) of the 11 D1 single mutants, 2 double mutants (DM-1: H118R, I122V; DM-2: E127K, V131A) and of 2 triple mutants (TM-1: N74S, I76S, I78L; TM-2: F150Y, I152T, N158S). The experimental conditions were as in Fig. 1. (d) ELISA analysis of the DM-3 double mutant (I152T, N158S) and of the TM-3 triple mutant (with the additional H118R substitution) together with the corresponding single mutants. The experimental conditions were as in panel C, except that the concentration of biotinylated peptide-coupled tubulin used for coating was decreased 5-fold (down to 2 nM).
Figure 4. Conformational changes of D1 upon tubulin binding. (a) Overview. Uncomplexed D1 (in orange, C-cap in yellow) has been superimposed on D1 in tubulin–D1 (D1 in green with the C-cap in brighter color, β tubulin in beige), taking the N-cap and the internal repeats as a reference. (b) Close-up (only β tubulin and uncomplexed D1 are shown). Without a C-cap rotation, its residues would clash with tubulin. Five distances shorter than 2 Å are highlighted as black solid lines.

Figure 5. The C-cap motif of the high affinity DARPin is mobile in the crystal structure. (a) The H118R mutation destabilizes the C-cap. The TM-3 structure (blue) was superimposed to uncomplexed D1, colored as in Fig. 4a. The Arg118 side chain adopts two alternate conformations in TM-3 (Supplementary Fig. 6). Both would clash with the C-cap if it were folded as in D1 (3 distances shorter than 1.5 Å are highlighted as black solid lines), as would the many other Arg118 conformations we modeled. (b) Comparison of the complexes of tubulin (beige) with D1 (green) and with A-C2 (blue). The β tubulin subunits have been superimposed. Some side chains are drawn to help visualize the shift between the DARPin. (c) View of the tubulin – D1 interface colored by electrostatic potential. (Left) Electrostatic potential surface of β tubulin (red, negative; blue, positive) with bound D1 (green). The side chains of two acidic residues of the C-cap that are close to an acidic part of the surface of tubulin are shown. The Cαs of residues 148 and 149 are also highlighted as magenta spheres. Subtilisin cleaves the affinity-improved DARPin preferentially after these two residues (see text and Supplementary Fig. 2). (Right) Electrostatic potential surface of D1 centered on the tubulin-interacting surface. Note that the two panels are not to scale.
(Fig. 4a). With respect to the N-terminal and internal ankyrin repeats, the 2 C-terminal helices rotate by 25° in the complex, thereby preventing a clash with tubulin (Fig. 4b).

We next determined the structure of a high-affinity DARPin (namely the TM-3 triple mutant), not bound to tubulin (Supplementary Table 1). These data should allow us to establish whether the evolved DARPin binds to tubulin with a conformation different from that of D1, as expected from the SPR data, as immobilized DARPin host a higher affinity for tubulin. D1 (Figs 3b and 4a, and 5b), the total surface buried at the tubulin–DARPin interface calculated from the structures went from ~1825 Å² in the D1 complex down to ~1260 Å² in the case of the A-C2 complex. Therefore, going from tubulin–D1 to tubulin–A-C2, an order-to-disorder transition led to a gain in affinity associated with a ~30% decrease of the buried surface area.

A conformational change upon complex formation, as is the case of D1 upon tubulin binding (Fig. 4a), is expected to limit the association rate constant \( k_{\text{on}} \). In contrast to this expectation, the \( k_{\text{on}} \) value of acrylodan-labeled D1 was intriguingly high and moreover close to that of the labeled evolved DARPin A-C2 (Table 1). To ascertain the kinetic parameters determined by fluorescence, we used an orthogonal method and studied the tubulin–DARPin interaction by surface plasmon resonance (SPR). DARPin were immobilized via their N-terminal his-tag on a Ni²⁺-activated tris-NTA sensor chip. The association rate constants were evaluated by injecting different concentrations of tubulin while the subsequent injection of buffer provided an estimate of the dissociation rate constants (Fig. 2d). Whereas the absolute values differed slightly, the SPR results confirmed the trends observed with the fluorescence data. They confirmed in particular that the affinity for tubulin of A-C2 and TM-3 is two orders of magnitude higher than that of D1, and that the main contribution in this affinity increase comes from the \( k_{\text{on}} \) term (Table 1). Even though the \( k_{\text{on}} \) values are one order of magnitude lower than those estimated by fluorescence, the SPR data also confirmed that D1 and the affinity-improved DARPin share similar association rate constants for tubulin binding. The reasons for this last observation are not clear at present. One possibility is that the C-cap in D1 is associated more weakly than normal due to long-range repulsive interactions with neighboring molecules in the TM-3 crystal if it were folded as in D1, leaving the possibility that different conformations of TM-3 coexist in solution but that only the one with a destabilized (mobile) C-cap is compatible with this crystal packing.

Therefore, we also determined the structure of the evolved DARPin A-C2 in complex with tubulin (Supplementary Table 1). The comparison of this structure with that of tubulin–D1 indicated that the structure of the complex did not change much with respect to the DARPin considered (root mean square deviation (r.m.s.d.) between tubulin–D1 and tubulin–A-C2: 0.62 Å (967 C-alpha compared)). Only a slight difference was detected in the DARPin orientation: once the tubulin 3 subunits of tubulin–A-C2 and of tubulin–D1 have been superimposed, an additional 3 to 4° rotation was needed to superimpose the DARPin moieties of the complexes (Fig. 5b).

Actually, the main difference between the two complexes is that in tubulin–A-C2 the two C-terminal helices forming the C-cap motif were not defined in TM-3. However, in contrast to the structure of isolated TM-3, an ordered C-cap would generate only few clashes with symmetry-related molecules in the tubulin–A-C2 crystal; these clashes could be avoided by a different side chain rotamer of two residues.

Taken together, these results indicate that a destabilized C-cap is a characteristic of the evolved DARPin that contributes to the higher affinity for tubulin. In addition to different interaction strengths that might result from the slight DARPin reorientation on tubulin (Fig. 5b), the main structural feature that may explain this affinity difference is an unfavorable contribution of the C-cap in tubulin–D1. Indeed, the tubulin–D1 interface colored by electrostatic potential suggests that there are repulsive charge interactions involving the C-cap: the two C-terminal helices of D1 comprise 5 acidic residues, two of them (Asp155 and Glu159) being close to tubulin and in the vicinity of a mostly acidic cavity (Fig. 5c). The faster dissociation of D1 from tubulin compared to that of the evolved DARPin (Table 1) is well consistent with this scheme. Remarkably, because the C-cap interacts with tubulin in tubulin–D1 (Figs 3b and 4a, and 5b, c), the total surface buried at the tubulin–DARPin interface calculated from the structures went from ~1825 Å² in the D1 complex down to ~1260 Å² in the case of the A-C2 complex. Therefore, going from tubulin–D1 to tubulin–A-C2, an order-to-disorder transition led to a gain in affinity associated with a ~30% decrease of the buried surface area.

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The C-terminal capping motif of the high affinity DARPin is disordered in solution. Because the structural data indicated that the C-cap of the high-affinity DARPins is mobile in the crystals (Fig. 5a,b), we performed an ensemble of experiments to characterize the C-cap behavior of the DARPin in solution. Gel filtration analysis showed that the elution volume of A-C2 and of TM-3 is lower than that of D1 (Fig. 6a), indicating a higher hydrodynamic radius. This result is expected for proteins of similar molecular weights but with different folding states. An alternative explanation, that of a different oligomeric state of the evolved DARPins compared to that of D1, was ruled out by size exclusion chromatography coupled to multi-angle laser light scattering (SEC-MALLS) analysis: both D1 and TM-3 remained monomeric under the conditions of the gel filtration experiment (Supplementary Fig. 1). We also submitted the DARPins to limited proteolysis. In the presence of subtilisin at a 1:2000 protease:DARPin molar ratio, in the case of A-C2 and of TM-3, a polyacrylamide gel electrophoresis analysis indicated that the intensity of the band corresponding to full length DARPin decreased rapidly whereas that of a band of slightly lower molecular weight increased sharply (Fig. 6b). Under the same conditions, uncleaved DARPin remained the main species in case of D1. Mass spectrometry analysis of a subtilisin-treated TM-3 sample identified a mixture of two species of about 15.8 kDa with a mass difference of 71 Da (Supplementary Fig. 2). Only a TM-3 fragment of limits 1-149 was compatible with the higher mass whereas the lower mass matched with that of fragments of limits 1-148 (one Ala residue shorter than 1-149) and 11-160. The latter solution involving two proteolytic cuts implied that, combined with the cut after residue 149, two
additional fragments (1-160 and 11-149) should exist (Supplementary Fig. 2). Because these fragments were not detected by mass spectrometry, we conclude that subtilisin cleaved the TM-3 DARPin preferentially after residues 148 or 149. These cleavage sites, at the C-terminal end of the loop connecting the last internal ankyrin repeat to the C-cap helical motif (Fig. 5c), were less accessible in D1. These results point to evolved DARPins with a flexible C-cap, in agreement with the structural data. Circular dichroism analysis further indicated a lower helical content of these DARPins compared to that of D1 (Fig. 6c), therefore suggesting that their C-cap is not only mobile but also disordered.

In order to validate the result of the proteolysis experiment, we made DARPin constructs terminating after residue 149 both within the D1 framework and that of the evolved A-C2 and TM-3. Interestingly, the circular dichroism signal of the TM-3 1-149 construct was similar to that of TM-3 and of A-C2 (Fig. 6c), indicating a similar helical content and further supporting the hypothesis that the C-cap is disordered in the evolved full length DARPins. Remarkably, an ELISA analysis showed that the removal of the C-cap of D1 led to a high-affinity DARPin, similar to A-C2 or TM-3 and their 1-149 variants (Fig. 6d), again suggesting that the C-cap interaction is detrimental to tubulin−D1 assembly. Alternatively, because the 1-149 constructs were only marginally stable and prone to aggregation, the higher signal in the ELISA experiment might result from an avidity effect of multimeric (aggregated) 1-149 constructs. However, because of the similar signal obtained with the 1-149 constructs and with evolved full length DARPins, and because in the latter case the DARPins are monomeric (Supplementary Fig. 1), this mechanism seems less likely. Moreover, it has been shown previously that C-cap-truncated DARPins remain monomeric at the concentrations used in these experiments\(^2\), even though the C-cap is a main safeguard against aggregation. We also studied the interaction with tubulin of DARPin devoid of a C-cap by fluorescence spectroscopy and by SPR. For the fluorescence experiments, because of the difficulties to handle the 1-149 constructs, we generated a fluorescent short construct in one step, by submitting the acrylodan-labeled TM-3 variant to limited proteolysis. The results of these experiments agree with those of ELISA and show that the affinity for tubulin of DARPin either without a C-cap (TM3 1-149 or subtilisin-digested TM-3) or with a disordered one (A-C2 and TM-3) are of the same order of magnitude (Fig. 2a, Table 1). Another important conclusion from these
factors are substantially higher than those of the rest of the molecule (75 Å² for the two C-terminal helices versus
boring repeat (illustrated in Fig. 5a). Consistently, in the structure of tubulin
interactions, including the hydrogen bonds between Asn158 and the carbonyl main chain of two residues of the neigh-
repeat. Upon tubulin binding, the D1 conformational change is accompanied by the loss of many of these interac-
rescence measurements in the subnanomolar range (Table 1).

47 Å² for the N-cap motif and the three internal repeats up to residue 149) 9. Therefore, the C-cap goes from a
that there is more than one way to destabilize the C-cap to yield high-affinity binders.
A-G2 does not contain any of the mutations introduced in the TM-3 optimized DARPin, these results also imply
that the affinity for tubulin is inversely correlated with the stability of the C-cap. Because the high-affinity DARPin
of these two substitutions was found in the low- and medium-affinity DARPins. Collectively these results suggest
binders, two of them had the H118R mutation and I152T was found three times (Supplementary Fig. 4c). None
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DARPins, the general trend was a mixture of full-length protein and cleaved fragments with bands of similar
intensity on SDS-PAGE (Fig. 7b), with the exception of one DARPin (B-B7) which was less resistant to subtilisin.
Finally, all the high-affinity binders were susceptible to subtilisin cleavage and behaved mainly as A-C2 (Fig. 7c).
We also sequenced all these DARPins. The low-affinity ones, which were also the most resistant to proteolysis,
had residue substitutions that were spread all along the sequence (Supplementary Fig. 4a). In the case of the
medium-affinity DARpins, several mutations were clustered in the third internal repeat and in the C-cap (framed
in Supplementary Fig. 4b), in the region expected to influence the C-cap stability. In the case of the high-affinity
binders, two of them had the H118R mutation and I152T was found three times (Supplementary Fig. 4c). None
of these two substitutions was found in the low- and medium-affinity DARpins. Collectively these results suggest
that the affinity for tubulin is inversely correlated with the stability of the C-cap. Because the high-affinity DARPin
A-G2 does not contain any of the mutations introduced in the TM-3 optimized DARPin, these results also imply
that there is more than one way to destabilize the C-cap to yield high-affinity binders.

Destabilizing the C-cap is the major route towards higher affinity binders. We then asked whether
the stability of the C-cap also correlates with the affinity for tubulin of the other variants generated from D1. As
a means to characterize the C-cap stability, we recorded the resistance to proteolysis of all the variants studied in
Fig. 1. The DARPinS ranked as low-affinity binders were mostly resistant to limited proteolysis (Fig. 7a). The band
corresponding to full-length protein remained the strongest one in all cases. In the case of the medium-affinity
DARpins, the general trend was a mixture of full-length protein and cleaved fragments with bands of similar
intensity on SDS-PAGE (Fig. 7b), with the exception of one DARPin (B-B7) which was less resistant to subtilisin.
Finally, all the high-affinity binders were susceptible to subtilisin cleavage and behaved mainly as A-C2 (Fig. 7c).

Discussion
In this study, we identified DARPinS with an improved affinity for tubulin associated with the destabilization of
the C-cap. The progenitor of evolution, DARPin D1, shows electrostatic repulsions (Fig. 5c) and it would clash
with the target tubulin within the C-cap, as can be deduced from the structure of the uncomplexed molecule
when docked onto tubulin. The actual complex of D1 with tubulin avoids this clash by a rotation of the C-cap
(Fig. 4). Evolution has brought about mutations which destabilize the interaction of the C-cap with the rest of the
DARPin, leading to an overall affinity increase with the target. Thus, the energetic cost of removing the C-cap
from its normal position has been lowered, translating into an overall better affinity with a K_D estimated by fluo-
rescence measurements in the subnanomolar range (Table 1).

In uncomplexed D1, like in all DARPinS, the C-cap mediates numerous interactions with the last internal repeat. Upon tubulin binding, the D1 conformational change is accompanied by the loss of many of these interac-
tions, including the hydrogen bonds between Asn158 and the carbonyl main chain of two residues of the neighbor-
ning repeat (illustrated in Fig. 5a). Consistently, in the structure of tubulin–D1, the C-cap atomic temperature
factors are substantially higher than those of the rest of the molecule (75 Å² for the two C-terminal helices versus
47 Å² for the N-cap motif and the three internal repeats up to residue 149) 9. Therefore, the C-cap goes from a
well-defined state in D1 to an intermediate, more mobile, state in the tubulin–D1 complex and to a disordered
(Fig. 6) state in the evolved DARPinS. In this last case, the negative charges of the C-cap are remote from a tubulin
negative patch, unlike what is seen in tubulin–D1, providing a rationale for the gain in affinity. It remains however
possible that, in the evolved DARPinS, the C-cap, while too mobile to be seen in the crystal structure, contributes
to the interaction by mediating dynamic contacts with tubulin. But, because the DARPinS devoid of a C-cap
behave as high-affinity tubulin-binders (Figs 2a and 6d, Table 1), the contribution of these putative dynamic
interactions is likely to be low. Nonetheless, the presence of the disordered C-cap protects the protein against
aggregation.

The three mutations introduced to generate the optimized TM-3 variant from the parent DARPin (Fig. 3a)
all contribute to destabilize the C-cap. They do so either by introducing steric conflicts between the last internal
repeat and the C-cap (as is the case for the H118R substitution, Fig. 5a) or by removing potential stabilizing
and 0.1 nM in generations 1 and 2, respectively. After 10 nM of target was used for immobilization, while for off-rate selection rounds its concentration was reduced to 1, 0.5% BSA for 30 minutes. For rounds 1 and 3 (diversification and rescue rounds, respectively) of both generations, the target-bound translated library was incubated with 1, 0.5% BSA for 30 minutes. For rounds 1 and 3 (diversification and rescue rounds, respectively) of both generations, the target-bound translated library was incubated with 1 μM tubulin in WBT-B for 2 hours. After incubation with the

Affinity maturation is a common approach to identify high(er) affinity binders. Potentially every residue can be diversified, including those of the so-called protein framework, i.e. outside of the residues randomized in the original library and chosen to point towards typical targets. As an example, in the case of DARPin selected to bind the human epidermal growth factor receptor 2 (Her2), the mutation of 4 residues of the framework accounted for most of the gain in affinity. Of particular interest, the main beneficial change was due to the substitution H52Y. This residue of the first internal repeat is at the interface with the second internal one. Structural analysis indicated that this substitution led to an opening of the anti-Her2 DARPin with a rotation of the 2nd internal repeat with respect to the first one. More recently, it was shown that this open conformation is retained when this DARPin is in complex with its target and fits it very well. A similar structural change is also found in D1. In this case it is induced upon tubulin binding and involves the interface between the last internal repeat and the C-cap (Fig. 4a). The residue 52 of the anti-Her2 DARPin corresponds to residue 118 of the 3rd internal repeat of D1. Therefore, a mutated residue at this position in a repeat has been selected twice in affinity maturation experiments for two unrelated targets. It either induces a ready-to-interact conformation (anti-Her2 DARPin) or destabilizes the C-cap (anti-tubulin DARPin, Fig. 5a). These two pathways lead to different structural results but in both cases to improved affinities.

Protein–protein association is often accompanied by local structural changes and by domain movements, either induced fits or disorder-to-order transitions. In principle, during a maturation process, evolution may improve the interaction by taking advantage of all these mechanisms. It may replace residues at the protein–protein interface, either to provide the evolved binder with chemical groups ideally positioned to interact with the partner (e.g., ref. 23) or to remove the entropic cost of stabilizing long and flexible side chains. Indeed the charged amino acids Lys, Glu and Asp are found depleted at those interfaces, an observation that has been harnessed to enhance the crystallization ability of proteins through the mutation of lysine and glutamate patches. Evolution may also lead to the stabilization of a ready-to-interact conformation (e.g., ref. 22) or to the optimization of loop length, which is best described in antibody maturation. ORDER-TO-DISORDER transitions have also been reported; in these cases the mechanism relies on the exposure of a previously hidden well-defined interacting region. To the best of our knowledge, the specific mechanism found here is unprecedented. It is an order-to-disorder transition, going from an (ordered) parent binder (with a repulsion to the target) to a mature one (where one motif is disordered and the repulsion is avoided), with two orders of magnitude higher affinity, resulting in a sub-nanomolar dissociation constant. It has been brought out in an in vitro selection experiment. Whether this mechanism also operates in nature remains to be determined.

**Methods**

**Affinity maturation, ribosome display selection and screening of high affinity DARPin.** Affinity maturation of DARPin D1 was performed over two generations using the ribosome display selection procedure as described earlier. Each generation was comprised of three rounds: (i) diversification by error-prone PCR followed by selection of all binders using ribosome display, (ii) amplification followed by incubation with competitor for slow off-rate selection under highly stringent conditions, and (iii) a rescue round to amplify and select the rare high-affinity binders.

**Subcloning and error-prone PCR.** For affinity maturation, DARPin D1 gene cloned in pDST679 was subcloned in pRDV829, which was used as template. Error-prone PCR was performed in the presence of dNTP analogues, dPTP (6-(deoxy-β-d-erythro-pentofuranosyl)-3,4-dihydro-8H-pyrimido-[4,-5-c][1,2]oxazine-7-one-5’-triphosphate) and 8-oxo-dGTP (8-oxo-2’-deoxyguanosine-5’-triphosphate)28. Two PCR reactions were performed using 3 and 9 μM dNTP analogues to obtain PCR products at low and high mutational loads, respectively. PCR products of both reactions were mixed in equimolar amounts and used as template for in vitro transcription.

**Ribosome display selection.** The random mutant library obtained by error-prone PCR was subjected to in vitro transcription and translation. All selection steps were performed at 4°C, except the blocking step which was performed at room temperature. For selection, 96-well MaxiSorp plates (Nunc) were coated overnight with 66 nM neutravidin or streptavidin in 50 mM Tris-HCl pH 8, 150 mM NaCl (TBS). Coated plates were washed 3 times with TBS and blocked for 1 hour with 0.5% BSA in TBS supplemented with 0.05% Tween-20 (TBST). Blocked plates were washed 3 times with TBST then biotinylated stathmin-derived peptide-coupled tubulin was incubated in 20 mM PIPES-K pH 6.8, 150 mM KCl, 50 mM Mg-Acetate, 0.05% Tween-20, and 0.1 mM GTP (buffer WBT) with 0.5% BSA for 30 minutes. For rounds 1 and 3 (diversification and rescue rounds, respectively) of both generations, 10 nM of target was used for immobilization, while for off-rate selection rounds its concentration was reduced to 1 and 0.1 nM in generations 1 and 2, respectively. After in vitro translation, the random mutant library of DARPin D1 was incubated with the target for 1 hour and washed five times with WBT comprising 0.1% BSA (WBT-B). From the second round onward, the translation mix containing the ternary miRNA–ribosome–DARPin complexes were first prepped for 15 min in a well without the biotinylated target. Further, in the off-rate selection rounds the target-bound translated library was incubated with 1 μM tubulin in WBT-B for 2 hours. After incubation with the
with 250 mM imidazole in 50 mM Tris-HCl pH 8, 150 mM KCl. Buffer was exchanged using PD-10 columns (GE Healthcare) equilibrated with 20 mM PIPES-K pH 6.8, 150 mM KCl, 1 mM MgCl₂ and 0.5 mM EGTA (PKM buffer). The concentration of DARPins was estimated by UV absorption at 280 nm, using extinction coefficients calculated from the sequence with the ProtParam webserver (http://web.expasy.org/protparam). For DARPin 1-149 constructs, care was taken to remove any aggregates. In particular, just before use, purified proteins were ultracentrifuged and subjected to an additional gel filtration step. The peak corresponding to 1-149 monomers was collected and the protein concentration was determined using a bicinchoninic acid assay (BCA; Thermo Fisher Scientific) taking DARPin D1 as a reference. DARPin D1 was labeled with acrylodan mainly as described previously. Briefly, the cysteine of Q26C mutants of D1, A-C2 and TM-3 was reduced and then reacted with acrylodan, following which the excess dye was removed by size-exclusion chromatography on a Superdex 200 10/300 GL (GE Healthcare). As for the 1-149 constructs, BCA was used to determine the concentration of labeled proteins.

**ELISA experiments.** To rank the DARPin variants according to their affinity for tubulin, ELISA experiments were performed essentially as described above for the analysis of the crude extracts, with minor modifications. For lower screening stringency, 10 nM of biotinylated peptide-coupled tubulin was used for immobilization on neutravidin-coated plates, while to achieve higher stringency the concentration of modified tubulin was reduced to 2 nM. After a washing step, purified DARPin diluted to 30 nM in WBT-B were incubated with the immobilized target for binding. After washings, the target-bound DARPin were incubated for 1 h either with buffer (WBT-B) or with 100 nM tubulin to prevent the rebinding of detached DARPins. Detection was performed as described above.

Equilibrium and kinetic analysis of tubulin–DARPin interaction by fluorescence. The dissociation constant (K_d) at equilibrium was determined using a fixed concentration of acrylodan-labeled DARPin (either 100 nM (D1) or 15 nM (high affinity DARPins)) titrated against increasing concentration of tubulin in 20 mM K-phosphate buffer pH 7.2, 100 mM KCl, 1 mM MgCl₂, 0.5 mM EGTA and 10 μM GDP at 20°C. The increase in fluorescence (λ_ex, 290 nm; λ_em, 510 nm) was monitored in a FluoroMax spectrofluorometer (Jobin Yvon, HORIBA). The data were fitted to a 1:1 binding isotherm using Equation 1.
$$\Delta \text{Fluo} = \text{Fluo}_{\text{Max}} \times \frac{\{D^*\} + \{T\} + K_D - \sqrt{((\{D^*\} + \{T\} + K_D)^2 - 4 \times \{D^*\} \times \{T\})}}{2 \times \{D^*\}}$$

where \(\Delta \text{Fluo}\) is the variation of the fluorescence signal, \(\text{Fluo}_{\text{Max}}\) is the fluorescence at saturating concentration of tubulin, \([T]\) and \([D^*]\) are the concentrations of tubulin and labeled DARPin, respectively, and \(K_D\) is the dissociation equilibrium constant.

The dissociation rate constant \(k_{\text{off}}\) of D1 was determined by adding 2.75 or 5.5 \(\mu\)M unlabeled D1 to a 100 nM labeled D1 and 500 nM tubulin solution (final concentrations) using a SX20 Stopped-Flow spectrometer (Applied Photophysics). The same results were obtained with both concentrations of unlabeled D1 used as a chase. For the high affinity DARPin\(\_\text{S}, 2\ \mu\)M unlabeled DARPin was added to the fluorescent complex made of 20 nM labeled DARPin and 40 nM tubulin, and the fluorescence decay was monitored in the spectrofluorometer. The drop in fluorescence signal was fitted with the following mono-exponential decay function,

$$\text{Fluo} = \text{Fluo}_{\text{Max}} + \Delta \text{Fluo} \times e^{-k_{\text{off}}t}$$

where \(\text{Fluo}\) is the fluorescence signal, \(\text{Fluo}_{\text{Max}}\) is fluorescence at infinite time, \(\Delta \text{Fluo}\) is the amplitude of the fluorescence variation, and \(k_{\text{off}}\) corresponds to \(k_{\text{off}}\) at saturating concentration of DARPin.

The association rate constant \(k_{\text{on}}\) was determined using a Hi-Tech KinetAsyst stopped-flow system (TgK Scientific). At least a 5-fold excess of tubulin was added to a fixed concentration (50 nM) of labeled DARPin, which made the binding reaction pseudo-first-order. The \(k_{\text{obs}}\) value was fitted with the following exponential equation,

$$\text{Fluo} = \text{Fluo}_{\text{Max}} + \Delta \text{Fluo} \times \left(1 - e^{-k_{\text{obs}}t}\right)$$

where \(\text{Fluo}_{\text{Max}}\) is the fluorescence at time 0. \(k_{\text{on}}\) was extracted from the variation of the apparent rate constant \(k_{\text{obs}}\) as a function of the tubulin concentration.

**Kinetic analysis of tubulin–DARPin interaction by SPR.** Surface plasmon resonance was measured using a Proteon XPR36 instrument (Bio-Rad). The measurements were performed at 20 °C in the same buffer as the one used in the fluorescence experiments, but with a lower (20 \(\mu\)M) EGTA concentration and adding 0.005% Tween 20. HTG sensor chips (Bio-Rad) were used to immobilize the DARPin\(\_\text{S}\) through their His-tag. For the determination of kinetic data, tubulin was injected at five different concentrations centered around the expected \(K_D\) value. The kinetic data were fitted according to the Langmuir isotherm using the Proteon Manager software. In the case of D1, although the fit was not as good as that of the data of the evolved DARPin\(\_\text{S}\), the different conditions tested (different amounts of immobilized D1, various tubulin concentrations, different durations of the association phase) led to very similar kinetic values. In addition, the estimation of the \(K_D\) obtained by plotting the RU value at the plateau as a function of tubulin concentration (Supplementary Fig. 5) agreed with the \(k_{\text{off}}/k_{\text{on}}\) ratio value.

**Crystallization and structure determination.** Uncomplexed D1 (not bound to tubulin) was crystallized by vapor diffusion in a buffer consisting of 100 mM Na-acetate pH 4.6, 20 mM CaCl\(_2\) and 30% 2-methyl-2,4-pentanediol. Crystals were flash-cooled in liquid nitrogen and diffraction data were collected at 100 K at the SOLEIL Proxima 1 beamline and processed with XDS\(_\text{34}\) and XSCALE\(_\text{35}\). The structure was solved with Phaser\(_\text{36}\) taking the structure of a 3-variable ankyrin repeat DARPin (PDB code 4DRX\(_\text{9}\)) as a search model. The structure was refined in Phenix\(_\text{38}\), alternating refinement with manual model building in Coot\(_\text{39}\).

To crystallize the tubulin–A-C2 complex, the C-terminal tails of tubulin were first cleaved with subtilisin\(_\text{42}\). A-C2 was added to digested tubulin in 1:5:1 A-C2:tubulin molar ratio. The complex was purified on a Superdex 200 10/300 GL column (GE Healthcare), equilibrated with 20 mM Hepes pH 7.2, 100 mM KCl, 1 mM MgCl\(_2\) and 0.5 mM EGTA. The GDP concentration was adjusted to 0.2 mM, then the complex was concentrated to 200 \(\mu\)M and stored in liquid \(N_2\) until use. Crystals grew spontaneously within 24 h in 25% (w/v) polyethylene glycol 4000 (PEG 4000), 0.1 M tri-sodium citrate pH 5.6 and 0.2 M ammonium sulfate. These crystals were used for micro-crystallization in 2-methyl-2,4-pentanediol. Crystals were flash-cooled in liquid nitrogen and diffraction data were collected at 100 K at the ESRF ID30A-3 beamline. Data were processed with XDS\(_\text{44}\) and Aimless\(_\text{40}\) and the structure was solved with Phaser using tubulin–D1 (PDB code 4DRX\(_\text{9}\)) as a search model. The structure refinement was performed using Buster. Data collection and refinement statistics are reported in Supplementary Table 1. Coordinates and structure factors have been deposited with the Protein Data Bank with accession codes 4DU1 (D1), 5EYL (TM-3) and 5EYP (tubulin–A-C2). Figures of structural models were generated with PyMOL\(_\text{43}\). The electrostatic potential at the tubulin–D1 interface was calculated using APBS\(_\text{44}\) and rendered in PyMOL.

**Characterization of the DARPin\(\_\text{S}\) in solution.** **Analytical Gel Filtration.** Samples of 100 \(\mu\)M purified DARPin\(\_\text{S}\) were injected on Superdex 75 10/300 GL column (GE Healthcare) equilibrated
with 50 mM K-phosphate pH 7.2, 150 mM KCl, 1 mM MgCl₂ and 0.5 mM EGTA. Proteins were resolved at 0.4 ml/min flow rate using an Äkta Purifier system (GE Healthcare).

Limited proteolysis. Proteolysis of DARPins was performed in 20 mM Hepes pH 7.2, 100 mM KCl, 1 mM MgCl₂ and 0.5 mM EGTA. DARPins at 40 μM were mixed with 20 nM subtilisin (Protease Type VIII; Sigma) and incubated at 25°C. Aliquots were removed at 0, 15 and 30 minutes and the reaction was stopped by adding 0.1% (w/v) phenylmethanesulfonyl fluoride and subjected to SDS-PAGE analysis.

Mass Spectrometry Analysis. TM-3 DARPins was subjected to limited proteolysis for 30 minutes as described above. Post cleavage buffer was exchanged to 50 mM ammonium acetate and the sample was analyzed by electrospray ionisation mass spectrometry (ESI-MS).

Circular dichroism. Far-UV circular dichroism spectra of 20 μM purified DARPins (5 μM in the case of TM-3 1-149) in 50 mM Na-phosphate pH 7.2, 150 mM NaCl were recorded from 200 to 260 nm in a 1 mm path length cuvette using a J-810 Spectropolarimeter equipped with a Peltier-controlled cuvette holder (Jasco). Data were acquired with 3 accumulations at a scan speed of 100 nm/min with 0.2 nm pitch and 2 s integration time. The bandwidth was 2 nm. All spectra were buffer-subtracted and normalized and are represented as molar ellipticity. Thermal unfolding of DARPins was performed by heating 10 μM of protein from 20 to 90°C at a rate of 0.5°C per minute while recording ellipticity at 222 nm. Reversibility of thermal unfolding was estimated by cooling the protein samples to 20°C followed by recording the spectra after 20 minutes of incubation. The extent of reversibility for D1 and A-C2 was 95%, respectively.

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Author Contributions
S.A., A.P., M.K. and B.G. designed research; S.A. performed the maturation experiments with the help of B.D, characterized the DARPin biochemically, crystallized the tubulin–DARPin complex and determined its structure; L.P. determined the structure of DARPin D1; D.H. assisted with the stopped-flow experiments; M.A.N. assisted with the SPR experiments; B.G. determined the structure of the TM-3 DARPin from crystal obtained by S.A.; S.A, A.P., M.K. and B.G. analyzed the data; A.P., M.K. and B.G. wrote the manuscript with input from all other authors.

Additional Information
Accession codes: Coordinates and structure factors have been deposited with the Protein Data Bank (accession code: 4DUI (D1), 5EYL (TM-3) and 5EYP (tubulin–A-C2)).
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