The E144 residue of Scherffelia dubia centrin discriminates between the DNA repair protein XPC and the centrosomal protein Sfi1

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Centrins are members of the EF-hand family of calcium-binding proteins, which are highly conserved among eukaryotes. Centrins bind to several cellular targets, through a hydrophobic triad. However, the W1xxL2xxL3 triad in XPC (Xeroderma Pigmentosum Group C protein) is found in the reverse orientation, as in the U1xxL2xxL3 triad in Sfi1 (Suppressor of Fermentation-Induced loss of stress resistance protein 1). As shown by previous NMR studies of human centrin 2 in complex with XPC or Sfi1, the E144 residue of human centrin 2 is in contact with XPC but is pushed away from the triad of Sfi1. We corroborated these findings using site-directed mutagenesis to generate mutants in Scherffelia dubia centrin (SdCen) and by using isothermal titration calorimetry to analyze the binding affinity of these mutants to XPC and Sfi1. We mutated the F109 residue, which is the main residue involved in target binding regardless of triad orientation, and the E144 residue, which was thought to be involved only in XPC binding. The F109L-E144A mutation reduced the binding of SdCen to XPC and Sfi1 and the negative effect was greater upon temperature increase. By contrast, the E144A mutation reduced the binding to XPC but had no effect on Sfi1 binding. The F109L-E144A mutation enhanced the negative effect of the two single mutations on XPC binding. Sfi1 proteins from Ostreococcus lucimarinus and Ostreococcus tauri, which belong to the same clade as S. dubia, were also investigated. A comparative analysis shows that the triad residues are more conserved than those in human Sfi1.

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1. Introduction

Centrins are small (~20 kDa), acidic, Ca²⁺−binding proteins that belong to the EF-hand superfamily. Centrins were first identified in the unicellular green algae *Tetraselmis striata* [1] and *Chlamydomonas reinhardii* as components of the contracting fibers associated with the basal bodies [2,3]. Centrins have also been found in many eukaryotes, including higher plants [4], yeast [5], invertebrates [6] and humans [7,8]. Only one isoform has been found in lower eukaryotes such as unicellular algae and yeast, whereas four isoforms have been found in higher eukaryotes. Even more isoforms have been identified in the unicellular ciliated protozoan *Paramecium tetraurelia* [9]. Purified centrins have been characterized, and algal centrin exhibits calcium-dependent filaments formation [10].

Centrins have been reported to interact with several cellular targets, such as centrosomal proteins, XPC, Sfi1, Sac3 and transducin. Involvement of centrin in cell division has been demonstrated in yeast. Three temperature-sensitive alleles of yeast Cdc31 that contain single point mutations (i.e., E97G, A101T and F105L) induce arrest in the G2/M transition of the cell cycle [11]. Centrins are concentrated in the microtubule organizer centers and are required for the cell cycle. When centrin is absent or non-functional, microtubule organizer center duplication is blocked, resulting in perturbed mitosis [12,13].

Centrins are also involved in several other cellular processes, for example, modulation of homologous recombination [14]. Centrin participates in the first step of nucleotide excision repair, in particular the global genome repair pathway, which involves centrin binding to XPC in concert with Rad23B. This heterodimer plays a key role in the initial phase of DNA-damage recognition by recruiting other factors, including TFIIH [15]. Because of the coupling of transcription, pre-mRNA processing, and the transport of mature RNA from

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**Abbreviations:** SdCen, Scherffelia dubia centrin; C-SdCen, C-terminal domain of Scherffelia dubia centrin; XPC, Xeroderma Pigmentosum group C protein; Sfi1, Suppressor of Fermentation-Induced loss of stress resistance protein 1; ITC, isothermal titration calorimetry; OstTa, Ostreococcus lucimarinus; OstT, Ostreococcus tauri; CID, proximal CTD-Interacting Domain; HCA, hydrophobic amino acids clusters analysis.

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the nucleus to the cytoplasm, interactions between the transcription machinery and the nuclear pore complex are necessary. Thus, proteins involved in this interaction play an important role in the process. In yeast, Sac3 acts in concert with Sus1 and Cdc31 to mediate interactions between the whole transcription machinery and the nuclear pore complex [16]. Experiments using immunoelectron microscopy have shown that centrin and transducin α co-localize in the connecting cilium of mouse rod photoreceptors. In addition, co-immunoprecipitation, centrifugation, and size-exclusion chromatography experiments have demonstrated that transducin β as well as the heterotrimer transducin αβγ binds centrin 1 [17,18]. SFI1, (suppressor of fermentation-induced loss of stress resistance 1), has been identified as an interacting partner for Cdc31 [19]. Furthermore, immunoelectron microscopy experiments revealed that Cdc31 and SFI1 are localized to the spindle pole body. Deletion of SFI1 causes G2 arrest in Saccharomyces cerevisiae [20]. In all these processes, centrin binds to a hydrophobic triad WXxxLXxxW [21] found in its cellular targets. Although this triad corresponds to the centrin-binding motif found in XPC, the centrin-binding motif of other targets (SFI1, Sac3, Transducin β) is in the opposite orientation LXXXLXXXW [22].

Several structures of centrin alone or in complex with cellular targets have been solved. Centrins are composed of two relatively independent domains, each one possessing two Ca2+ binding sites. Experiments using microcalorimetry have demonstrated that the C-terminal domain of centrin is sufficient for target binding. However, for algal centrins, the N-terminal domain also binds targets, but only in the presence of calcium [21,22]. The structures of the N-terminal domain of C. reinhardtii [21] and human centrins have been solved by NMR [23].

Two structures of full-length and truncated HsCen2 in complex with an XPC peptide have been solved by X-ray crystallography [24,25], and one structure of the C-terminal domain of HsCen2 in complex with XPC peptide in solution has been solved by NMR [26]. The structure from [24] shows that only residues in the C-terminal domain of HsCen2 are involved in the interaction with the XPC peptide. Nine non-polar residues of HsCen2 (L133, L112, M145, F113, M166, L126, A109, E105 and V129) form contacts with the residues of the XPC peptide. The F113 and M145 residues form the pocket where the W residue of XPC binds. The Y172 residue of HsCen2, which interacts with L9 of the XPC peptide, is located near the S170, which can be phosphorylated and is reported to have a functional role [27].

The structure of CID region (proximal CTD-Interacting Domain) of Sac3 in complex with Sus1 and Cdc31 has been solved by X-ray crystallography [28]. In the complex, the W802 residue of Sac3 appears to play a central role in the interaction with the hydrophobic cavity of the centrin Cdc31. The F105 residue of Cdc31 in turn plays an important role in the association with Sac3.

A crystal structure of centrin Cdc31 in complex with truncated SFI1 has been solved by X-ray crystallography [29]. Both the C-terminal and the N-terminal domains of Cdc31 make contact with SFI1. The structure also reveals several centrin–centrin interactions. These interactions suggest that a filament of several centrin molecules is formed by centrin–centrin interactions and that this filament is stabilized by SFI1 through SFI1 repeat–centrin interactions.

An NMR structure of human centrin 2 in complex with the R71-SFI1 peptide has been solved [30]. Comparison of this structure with superimposed with the NMR structure of the C-terminal domain of human centrin 2 in complex with either peptide-P17-XPC peptide [26] led to the proposal that the centrin 2 E148 residue discriminates between XPC and SFI1 [30]. Although the helix dipoles of the XPC and SFI1 peptides are reversed, their W residues are embedded in the hydrophobic cavity containing the centrin F113 residue. For SFI1, this requires a rotation of the W residue and a slight translation of the peptide within the cavity. The carboxyl group of E148, which faces the positive pole of the XPC helix, establishes a hydrogen bond with W residue of XPC. By contrast, if E148 faces the negative pole of the SFI1 helix, it is pushed away from the tryptophan leading to the disruption of hydrogen bonds.

We used site-directed mutagenesis to mutate the phenylalanine residue that is important for target binding as well as the glutamate residue that discriminates between XPC and SFI1 targets. We also used isothermal titration calorimetry (ITC) to assess the thermodynamic parameters of the binding reaction. The role of the phenylalanine residue in binding was first established using temperature-sensitive mutants in S. cerevisiae, thus we examined the effects of temperature on the association between centrin and two targets, XPC and SFI1. The E148 residue is conserved among centrins. Because there is only one centrin in unicellular green algae, the same centrin isoform interacts with all cellular targets regardless of the centrin-binding motif orientation. We used the Scherfelia dubia centrin to investigate the role of the E144 residue (which corresponds to E148 residue in human centrin) in the interaction with XPC and SFI1. We also retrieved sequences for SFI1 and XPC homologues protein from databases and analyzed their centrin-binding motifs.

2. Results

2.1. The centrin targets SFI1 and XPC are found in green algae

A single centrin is found in S. dubia that must interact with all cellular targets expressed in this unicellular green alga. Sequences of centrins have been found in several other green algae genomes using human centrins as query sequence. Several of the green alga genomes that were investigated contain only one centrin gene (data not shown), and these algal centrins are conserved (Fig. S1). The unique centrin sequences from S. dubia (SdCen) and Ostreococcus lucimarinus (OICen) were aligned with each other and with human centrins using the ClustalW program (Fig. 1). The C-terminal domain of SdCen (M93–F168) has 100% identity with OICen. Residues involved in the interactions with XPC, SFI1, and Sac3 (E105, A109, and F113 in human centrin) are conserved among the centrins from green algae as is the E148 residue specific for XPC binding (Fig. S1).

S. dubia is a unicellular green alga of the order Chlorodendrales that belongs to the clade of Prasinophyceae. Unfortunately, the genome of S. dubia has not been sequenced yet. However, the genomes of two species of Mamiellales, O. lucimarinus (OstLu) and Ostreococcus tauri (OstTa), which belong to the same clade as S. dubia, have been sequenced and analyzed.

A BLAST search using SFI1 from both human and S. cerevisiae as query sequences identified SFI1 homologues in the genomes of two other green algae, namely O. lucimarinus (SFI1 containing 1315 residues) and O. tauri (SFI1 containing 1370 residues). We identified centrin-binding motifs using a sequence alignment of human and algal SFI1 (Table 1) as well as the hydrophobic cluster analysis program HCA. Fig. S2 shows that the hydrophobic clusters identified by HCA are composed of the same residues identified by the sequence alignment (Table 1). Similar to human and yeast SFI1, SFI1 homologues from green algae are long molecules composed of 28 and 22 centrin-binding motifs for OstLu and OstTa, respectively. Using the protein secondary prediction programs SOPMA and GOR4, we found that SFI1 proteins from both green algae are composed of 70 to 90–93% of α-helices similar to SFI1 found in other organisms. We compared the hydrophobic triad LXXXLXXXW that forms the centrin-binding motif of SFI1 proteins from OstLu, OstTa, and human (Table 1). We found that the W residue in position 1 is absolutely conserved except for an L residue found in one repeat in human SFI1. By contrast, several other residues replace W in S. cerevisiae SFI1 (data not shown). In algal SFI1, an F residue is absolutely conserved in position 4 with the exception for a V residue in one repeat of OstTa SFI1; however, in other organisms, L, W or F residues are found in position 4. Similar to SFI1 homologues from other organisms, an L residue is mainly found at position 8, with a basic residue at position 7. Thus, the hydrophobic
affinity for P17-XPC than for R18-Sf1 by a factor of 7 at 30 °C and by a factor of 10 at 37 °C (Table 2). However, the change in enthalpy (ΔH) differed only by 3 kcal/mol, and the change in entropy (ΔS) differed by 1 kcal/mol. Thus, for both targets the interaction is enthalpically driven.

The effect of temperature increase on the binding of C-SdCen to P17-XPC and R18-Sf1 was measured by ITC in the absence of calcium (with EDTA) at two temperatures, 30 °C and 37 °C. Increasing the temperature from 30 °C to 37 °C decreased the binding of C-SdCen to P17-XPC by a factor of 3 and decreased the binding to R18-Sf1 by a factor of 4 (Figs. 3 and 4). The change in enthalpy (ΔH) decreased by a factor of 1.2, becoming more negative for both targets and indicating that the binding reaction was more exothermic (Table 2). The change in entropy (ΔS) also decreased with the temperature by a factor of 1.3, becoming more negative. However, increasing temperature did not change the binding reaction mode, which is still enthalpically driven.

2.3. Mutating phenylalanine 109 to leucine decreases the C-SdCen affinity for P17-XPC as well as for R18-Sf1

The F109 residue of SdCen corresponds to the F113 residue of human centrin 2, which has been shown to interact with several residues of XPC as well as Sf1, specifically the W1, L4 and L8 residues of the conserved hydrophobic triad. In the absence of calcium, the binding reaction of the C-SdCenF109L variant to P17-XPC as well as to R18-Sf1 was exothermic (ΔH negative) with a binding stoichiometry of one (Figs. 3 and 4). Mutating F109 to leucine decreased the binding of C-SdCen to P17-XPC as well as to R18-Sf1 but to different extents. The binding constant for the C-SdCenF109L-P17-XPC interaction was reduced by a factor of 15 at 30 °C and the same ratio was observed at 37 °C (Table 2). However, the binding constant for the C-SdCenF109L-R18-Sf1 interaction was reduced by a factor of 6 at 30 °C, and the same ratio was observed at 37 °C. The decrease in binding was accompanied by an increase in change in enthalpy (ΔH) for P17-XPC by a factor of 1.7.
Table 1
Sfi1 and Rad4 centrin-binding motifs.

|                | S. lucinmarinus | O. tauri | H. sapiens |
|----------------|-----------------|----------|------------|
| Sfi1 (SP: A488c7) | 1793 residues   | 1513 residues | 1242 residues |
| Sfi1 (SP: Q00F17) | 1513 residues   | 1242 residues | 1242 residues |
| Sfi1 (SP: A6K8F3) | 1242 residues   | 1242 residues | 1242 residues |
| **Rad4 (SP: A488c7)** | **637 residues** | **637 residues** | **637 residues** |
| **Rad4 (SP: Q00F56)** | **1122 residues** | **1122 residues** | **1122 residues** |
| **Rad4 (SP: A488c7)** | **637 residues** | **637 residues** | **637 residues** |
| **RKL** | **1242 residues** | **1242 residues** | **1242 residues** |
| **WTRAAL-586** | **846** | **846** | **846** |

at 30°C and by a factor of 1.4 at 37°C. Similarly, the change in enthalpy for R18-Sfi1 increased by a factor of 1.4 at 30°C and 37°C. Thus, mutating F109 to leucine resulted in a change in enthalpy (ΔH) that was less negative than that observed for the wild-type SdCen for both targets. Furthermore, the binding reaction of C-SdCenF109L to both targets was still enthalpically driven. The increase in the change in enthalpy (ΔH) was greater (10 kcal/mol) than the corresponding increase in binding free energy (ΔG) (less than 2 kcal/mol) (Table 3). Consequently there is a compensating gain in entropy of binding (10 kcal/mol) (Table 2).

2.4. Mutating glutamate 144 to alanine decreases the affinity of C-SdCen for P17-XPC but has no effect on R18-Sfi1 binding

The E144 residue corresponds to the E148 residue of HsCen2, which has been shown to interact with XPC, in particular with the W1 residue of the triad (Fig. 1). In the absence of calcium the binding of the C-SdCen E144A variant to P17-XPC was exothermic (ΔH negative) with a binding stoichiometry of one (Fig. 4). Mutating E144 to alanine decreased the affinity of C-SdCen for XPC by a factor of 3 at 30°C and by a factor of 4 at 37°C. However, the change in enthalpy
The ITC experiments were conducted at 30 °C or 37 °C in buffer (MOPS 50 mM, NaCl 100 mM, EDTA 2 mM or Ca²⁺ 2 mM pH 7.5). N was the stoichiometry of binding. NB means no binding. The standard deviations of the fitted parameters are given in parenthesis.

Table 2
Thermodynamic parameters of wild-type and mutant SdCen variants binding to XPC and Sfi1.

| Protein C-SdCen | Ligand    | Ca²⁺ | N   | Ka (error) (10⁶ M⁻¹) | ΔH (error) (kcal/mol) | TΔS (kcal/mol) | ΔG (kcal/mol) |
|-----------------|-----------|------|-----|----------------------|-----------------------|----------------|---------------|
| WT              | P17-XPC 30 °C | +    | 0.95  | 59.4(9.10)              | −23.0(0.16)             | −12.63         | −10.77        |
| 30 °C           | −         |      | 1.02  | 30.2(1.2)               | −33.8(0.07)             | −23.39         | −10.41        |
| 37 °C           | +         |      | 1.02  | 73.2(7.0)               | −23.36(0.08)            | −12.18         | −11.18        |
| 37 °C           | −         |      | 1.04  | 10.6(0.4)               | −29.6(0.12)             | −29.94         | −10.66        |
| F109L           | P17-XPC 30 °C | +    | 1.02  | 67.1(5.0)               | −15.63(0.09)            | −4.78          | −10.85        |
| 30 °C           | −         |      | 1.33  | 2.02(0.07)              | −20.6(0.09)             | −11.88         | −8.72         |
| 37 °C           | +         |      | 1.23  | 37.8(5.02)              | −16.97(0.10)            | −6.20          | −10.77        |
| 37 °C           | −         |      | 1.05  | 6.02(0.01)              | −28.96(0.2)             | −20.27         | −8.63         |
| E144A           | P17-XPC 30 °C | +    | 1.02  | 95.6(15.6)              | −20.44(0.12)            | −9.36          | −11.08        |
| 30 °C           | −         |      | 1.15  | 10.1(0.5)               | −29.5(0.13)             | −19.78         | −9.72         |
| 37 °C           | +         |      | 1.01  | 107(11.5)               | −25.17(0.62)            | −13.76         | −11.41        |
| 37 °C           | −         |      | 1.03  | 2.49(0.08)              | −38.4(0.17)             | −28.66         | −9.74         |
| F109L-E144A     | P17-XPC 30 °C | +    | 1.07  | 32.7(2.65)              | −14.11(0.06)            | −3.69          | −10.42        |
| 30 °C           | −         |      | 1.0   | 0.43(0.01)              | −22.2(0.2)              | −14.39         | −7.81         |
| 37 °C           | +         |      | 1.01  | 29.1(2.12)              | −17.19(0.06)            | −6.60          | −10.59        |
| WT              | R18-Sfi1 30 °C | −    | 1.12  | 4.5(0.1)                | −31.5(0.08)             | −22.24         | −9.26         |
| 37 °C           | −         |      | 1.15  | 1.23(0.02)              | −37.09(0.1)             | −27.88         | −9.31         |
| F109L           | R18-Sfi1 30 °C | −    | 1.17  | 0.77(0.01)              | −23.2(0.1)              | −14.96         | −8.24         |
| 37 °C           | −         |      | 1.05  | 0.23(0.01)              | −30.4(1.0)              | −22.3          | −8.10         |
| E144A           | R18-Sfi1 30 °C | −    | 1.19  | 4.64(0.08)              | −30.3(0.06)             | −21.06         | −9.25         |
| 37 °C           | −         |      | 1.12  | 1.38(0.04)              | −36.2(0.17)             | −26.88         | −9.34         |
| F109L-E144A     | R18-Sfi1 30 °C | −    | 1.17  | 0.66(0.02)              | −24.4(0.16)             | −16.33         | −8.07         |
| 37 °C           | −         |      | 1.04  | 0.29(0.01)              | −26.4(0.3)              | −18.24         | −8.19         |

The ITC experiments were conducted at 30 °C or 37 °C in buffer (MOPS 50 mM, NaCl 100 mM, EDTA 2 mM or Ca²⁺ 2 mM pH 7.5). N was the stoichiometry of binding. NB means no binding. The standard deviations of the fitted parameters are given in parenthesis.

The difference in binding enthalpy between the interaction of SdCen wt and P17-XPC and the interaction of SdCen mutants and P17-XPC, as well as for the Sdc-R18-Sfi1 complexes.

The difference in free energy between the interaction of SdCen wt and P17-XPC and the interaction of SdCen mutants and P17-XPC, as well as for the Sdc-R18-Sfi1 complexes.

(ΔH) was the same as for the wild type (ΔH = −30 kcal/mol at 30 °C and −38 kcal/mol at 37 °C) but the change in entropy increased (ΔG of −9.72 kcal/mol at 37 °C as well as at 37 °C). The change in free energy (ΔG) for C-SdCenE144A was an intermediate between that of wild-type C-SdCen and C-SdCenF109L (Table 2).

In the absence of calcium, the binding reaction of C-SdCenE144A to R18-Sfi1 was exothermic (ΔH negative) and enthalpically driven with a binding stoichiometry of one (Fig. 4). In contrast to what we observed with P17-XPC, the E144A mutation had no effect on the binding of C-SdCen to R18-Sfi1 and the SdCenF109L-E144A variant exhibited the same behavior as wild-type SdCen at both temperatures. The affinity of C-SdCenE144A for R18-Sfi1 was almost identical to the value found for wild-type C-SdCen, as were the changes in enthalpy and entropy values (Table 2).

2.5. Both F109L and E144A act in concert to stabilize the binding of C-SdCen to P17-XPC, but the effect of the double F109L-E144A mutation on R18-Sfi1 binding is similar to that of the single F109L mutation.

Because both single mutations had a negative effect on the binding of C-SdCen to P17-XPC, a variant with the F109L-E144A double mutation was constructed and analyzed. In the absence of calcium, the binding reaction of SdCenF109L-E144A to P17-XPC was exothermic (ΔH negative) with a binding stoichiometry of one (Fig. 3). The F109L-E144A double mutation decreased the binding of C-SdCen to P17-XPC more than the single F109L and E144A mutations did. The Ka values were 30 × 10⁶ M⁻¹, 1.0 × 10⁶ M⁻¹, 2.0 × 10⁶ M⁻¹, 0.4 × 10⁶ M⁻¹ for wild-type C-SdCen, SdCenE144A, SdCenF109L, and SdCenF109L-E144A, respectively, at 30 °C. The Ka values at 37 °C were lower (10 × 10⁶ M⁻¹, 2.5 × 10⁶ M⁻¹, 0.6 × 10⁶ M⁻¹, 0.25 × 10⁶ M⁻¹)
(Table 2). The double mutation led to an increase in the change in enthalpy (ΔH) (ΔH of −22.2 kcal/mol at 30 °C) and an increase in the change of free energy (ΔG) (ΔG of −7.81 kcal/mol at 30 °C) with a compensatory effect in the change in entropy (TΔS) (TΔS of −14.4 kcal/mol), which increased twofold compared to wild-type SdCen.

By contrast, the effect of the double F109L-E144A mutation R18-Sf1 binding was similar to the single F109L mutation. In the absence of calcium, the binding reaction was exothermic (ΔH negative) with a binding stoichiometry of one. The binding reaction of C-SdCenF109L-E144A to R18-Sf1 was enthalpically driven (Fig. 4). The double mutant C-SdCenF109L-E144A exhibited almost the same thermodynamic values (Kₐ, ΔH, ΔG, TΔS) as the single mutant C-SdCenF109L (Table 2).

3. Discussion

3.1. In green algae, a single centrin must bind several targets regardless of the orientation of their centrin-binding motif

Although the genome of S. dubia has not been sequenced yet, several other genomes from green algae have been sequenced, including O. lucimarinus and O. tauri. The sequences of the centrin found in these green algae were retrieved from genomes databases using the psiBlast program with the SdCen sequence as the query and aligned with SdCen using the ClustalW program (Fig. 1). Full-length SdCen has 90% identity with OstLu centrin, and the C-terminal domain of SdCen (M93-F168) is identical to the OstLu C-terminal domain (M90-F165). The SdCen sequence was also aligned with the human centrins, HsCen1 and HsCen2. In terms of the C-terminal domain, SdCen has the corresponding K131 residue of HsCen2 but lacks the S158 and N160 residues found in HsCen2. However, the S158 residue is only found in some organisms such as primates, and certain bird and some fish species. This suggests that they appeared through the evolution...
after the appearance of HsCen1 upon retrotransposition of HsCen2. Thus, sequence comparison suggests that the green algal centrin belongs to the HsCen2 family, whereas the yeast centrin Cdc31 belongs to the HsCen3 family. However, in both cases the unique centrin must bind several targets, including XPC, Sac3 and Sf1, which differ in the orientation of the centrin-binding motif.

We were able to identify Sf1 and Rad4 homologues in the genomes of the algae O. lucimarinus and O. tauri (Table 1). We further analyzed the evolution of the hydrophobic triad of these targets. The three important residues W1, L4, and L8 that form the triad of found in Sf1 repeats are more conserved in green algae than in higher eukaryotes. Indeed, in all Sf1 repeats, there is a W residue in position 1 and an F residue in position 4 of the triad. Such a strict conservation at position 4 is not found in higher organisms although the F residue is common at this position.

The centrin residues involved in target binding have been analyzed using various methods. First, the isolation of yeast temperature sensitive mutants allowed the identification of functionally important residues [11], Second, the structures of human centrin 2 [24–26] and yeast Cdc31 [29] in complex with targets have shed light on which residues play a role in target binding. For HsCen2, residues E105, A109, F113, and E148 have been identified as target-binding residues. These residues are highly conserved in centrins across species as well as in isoforms from the same organism.

3.2. General properties of C-SdCen binding to the cellular targets XPC and Sf1 which have opposite centrin-binding motifs

Regardless of the orientation of their centrin-binding motif, the binding of targets to SdCen was exothermic with one binding site in the C-terminal domain of SdCen. For both XPC and Sf1 targets, the binding reaction was enthalpically driven. Calcium was not required for binding although its presence enhanced the binding. ITC measurements performed at 30 °C in the presence of EDTA showed that C-SdCen had a greater affinity for P17-XPC than for R18-Sf1, with binding affinities of 30 × 10^6 M^-1 and 4.5 × 10^6 M^-1, respectively.

The temperature sensitive mutations identified in yeast centrin result in the arrest of cell division [11]; however, which centrin-binding interaction was perturbed was not elucidated. We attempted identify which centrin-target interaction is involved in the arrest of cell division by exploring the effect of temperature on centrin binding to XPC and Sf1. Our ITC experiments showed that a temperature increase...
had a more pronounced effect on Sf1 most likely because several centrin contacts are reduced in the case of Sf1. Indeed the previously published structure of human Cen2 showed that contacts between F113 (corresponding to SdCenF109) and Sf1 were reduced compared to XPC [30].

3.3. The F109 residue of SdCen is the main centrin residue involved in target binding regardless of centrin-binding motif orientation

Structural analysis of HsCen2 has shown that the F113 residue makes contacts with several target residues [30]. The important role of F113 has already been described in yeast where the corresponding mutation F105L in Cdc31 results in cell cycle arrest when the temperature is increased [11]. Mutating F109 to leucine decreased the binding of C-SdCen to both XPC and Sf1. However, the F109L mutation had a more negative effect on the affinity constant for P17-XPC than for R18-Sf1, with a decrease in Ka by a factor of 15 and 6, respectively. In addition, the change in enthalpy \( \Delta H \) was more pronounced in the case of P17-XPC than R18-Sf1 (Table 3). These results are consistent with previous observations made when the structures of Cen2/ P17-XPC [26] and Cen2/R17-Sf1 [30] are superimposed. Indeed, the dipole inversion in Sf1 reduces contacts between the centrin (F113) and the target-triad (L4).

3.4. The E144 residue of SdCen, which corresponds to the E148 residue of HsCen2, discriminates between the DNA repair protein XPC and the centrosomal protein Sf1

The C-SdCenE144A variant allowed us to confirm our hypothesis concerning the role of the E148 residue of HsCen2 in complex with target-derived peptides [30]. The observed effects of dipole inversion in centrin targets presented here can be considered a general effect because it was first elucidated in the human Cen2 structure and is now demonstrated using mutagenesis and thermodynamic experiments with green alga centrin.

The E148 residue of HsCen2 (or SdCenE144), which discriminates between XPC and Sf1, is strongly conserved in organisms that possess either a single centrin or several centrin isoforms. The E148 residue is located at the junction of helix 7 and the calcium-binding loop IV. Could it be required for a calcium-induced conformational change? This does not seem to be the case, because addition of calcium enhanced the affinity of XPC for both wild-type SdCen and the SdCenE144A variant. Thus, the question remains why the E148 residue (in human) is conserved among centrin isoforms from the same organism. Does it mean that all isoforms bind two types of targets regardless of the orientation of the centrin-binding motif? All ITC experiments were performed using target-derived peptides, and the interaction with E148 depends on the charge of peptide terminus facing it. In the future, the binding reaction should be characterized using full-length targets to avoid charge effects that may interfere with contacts involving the E148 residue.

3.5. What is the advantage of the centrin-binding motif orientation?

The ITC experiments presented here and in previous work demonstrated that the affinity of centrin is better for XPC than for other targets such as Sf1 [22,25,30] and transducin \( \beta \) [Grecu and Assairi to be published elsewhere]. This could suggest that the orientation of the centrin-binding motif of XPC is the best orientation for effective centrin binding. However, because calcium is more important for Sf1 binding [25,30] and absolutely required for binding to transducin \( \beta \) [17, Grecu and Assairi to be published elsewhere], the orientation of their centrin-binding motifs may be optimized for calcium-regulated binding.

4. Material and methods

4.1. Materials

The peptides used in this study were: R18-HsSf1, 677-SQHRQRLRGLRRLW-K692, and P17-XPC, 846-NWKLAKGLRERLKR-863. These peptides were purchased from GENE\textsc{CUST}; the peptides were acetylated at their N-terminus and amidated at their C-terminus for increased stability. Oligonucleotides were purchased from Eurogentec; dNTPs, Ndel and Xhol restriction enzymes, T4 DNA ligase and Finzymes Phusion DNA polymerase were purchased from New England Biolabs. The pET24a vector and Escherichia coli NM554, BL21(DE3) strains were purchased from Novagen; DEAE-TSK and phenyl-TSK were purchased from Interchim; Sephadex G25 was purchased from GE-Healthcare.

4.2. Cloning of S. dubia centrin (wild type and variants) and expression in E. coli

The S. dubia integral centrin (SdCen) coding sequence was amplified from a plasmid containing the centrin gene (gift of Michael Melkonian, University of Köl, Germany) by polymerase chain reaction (PCR) using Phusion DNA polymerase, dNTPs, and flanking 5’- and 3’-primers (5’-GGAGGGTCATGATTACGCGAAGGCTGCCC-3’ and 5’-GGCCTTCAGTACGAACAGTGCAGCTCTTTCTCAGT-3’) to create the plasmid pLA17.36.1 encoding the full-length SdCen. The PCR products were digested with the Ndel and Xhol restriction enzymes, and T4 DNA ligase was used to insert them between the Ndel and Xhol restriction sites of the pET24a expression vector. This recombinant plasmid was used as a template for site-directed mutagenesis. Site-directed mutagenesis of SdCen was accomplished via the double PCR method [31], using the full-length wild-type clone as the template, and adding dNTPs, Phusion DNA polymerase, both flanking 5’- and 3’-primers, and mutagenic oligonucleotides, F109L: 5’-CGCTCTTCGTCGTCTAATACGGCCGAAGCCCTT-3’ to create the plasmid pLA17.36.9; or E144A: 5’- GCCATGCCGTCGTCGCCCTCACTACTCCTGCC-3’ to create the plasmid pLA17.36–10. The double mutant E144A-F109L was constructed by using the plasmid pLA17–36–10, containing the mutation E144A, as the DNA template and the mutagenic oligonucleotide F109L: 5’-CGCTCTTCGTCGTCTAATACGGCCGAAGCCCTT-3’ to create the plasmid pLA17.36.11.

The C-terminal domain (residues Met93–Phe168) of wild-type SdCen and its variants was constructed by using the plasmids listed above as the DNA template encoding the full-length SdCen and the flanking primers: 5’-GGAAATCGCATATTGGCCAGCGCAGCTCCC-3’ and 5’-GGCCGTTGGATACGAAAGTCGTCATCTTCTAGT-3’ to create the plasmids pLA17.36.5 (wild type), pLA17.36.9 (F109L), pLA17–36–10 (E144A), pLA17.36.11 (E144A-F109L).

The PCR products were digested with the Ndel and Xhol restriction enzymes, and T4 DNA ligase was used to insert them between the Ndel and Xhol restriction sites of the pET24a expression vector. The resulting plasmids were used to transform E. coli NM554 cells, and the transformed clones were sequenced to verify their integrity and incorporation of the desired mutation. DNA from successfully transformed clones was isolated and used to transform the E. coli BL21(DE3)/pDIA17 strain [32], which is chloramphenicol resistant, in order to express the recombinant proteins.

For overproduction of SdCen and its variants, the transforms containing the centrin-encoding plasmid were grown at 37 °C in 2YT medium (Difco) supplemented with chloramphenicol (30 \( \mu \)g/mL) and kanamycin (70 \( \mu \)g/mL). When the absorbance reached 1.5 at 600 nm, the production of the protein was induced by addition of 1 mM isopropyl-\( \beta \)-D-thiogalactopyranoside, and the growth was continued for an additional 3 h at 37 °C. The cells were then pelleted
by centrifugation and used as a protein purification source.

4.2.2. Purification of the C-terminal domain of SdCen and its variants

Bacterial extracts were heated at 63 °C for 10 min, and the denatured proteins were removed by centrifugation. The wild-type and mutant variants of SdCen C-terminal domain were purified using three successive chromatographic steps, including DEAE-TSK (ion exchange), phenyl-TSK (reversed phase) and G25 (gel filtration). Centrins were then lyophilized and the degree of purity was assessed using SDS–PAGE [33]. The SdCen concentration was determined by UV spectrophotometry using the extinction coefficients of (E) of 1490 and 2860 M⁻¹ cm⁻¹ at 280 and 259 nm, respectively, based on the tyrosine and phenylalanine content [34].

4.2.3. Isothermal Titration Calorimetry (ITC)

The thermodynamic parameters of the molecular interactions between SdCen (C-terminal domain of wild type and variants) and the target peptides (P17-XPC, R18-Sf1) were investigated via ITC using a MicroCal MCS instrument (MicroCal Inc., Northampton, MA). The protein and peptide were equilibrated in the same buffer (50 mM MOPS, 100 mM NaCl pH 7.5) with 1 mM CaCl₂ or 2 mM EDTA. The centrins (10 µM) in the calorimeter cell were titrated with the target peptide (100 µM) by automatic injections of 8–10 µL at 30 °C or 37 °C. The first injection of 2 µL was ignored in the final data analysis. Integration of the peaks corresponding to each injection and baseline correction were performed using the Origin-based software provided by the manufacturer. The data were fit to an interaction model to generate the stoichiometry (n), equilibrium binding constant (Kₐ), and enthalpy of complex formation (∆H). The free energy (∆G) was determined using the following equations: ∆G = −RT ln(Kₐ), where K (1.987 cal/mol) is the gas constant and T (303 K) is the absolute temperature, and ∆G = ∆H − T ∆S. The following units were used: dissociation constant (Ka) M⁻¹; Gibbs energy change (∆G) kcal/mol; enthalpy change (∆H) kcal/mol; entropy change (∆S) kcal/mol.

4.2.4. Identification of Sf1, Rad4 and centrins

Sf1, Rad4, and centrins from various eukaryotes were identified via searches in the National Center for Biotechnology Information (NCBI) database, using the Psi-Blast program [35] with human and yeast homologue proteins as a query sequence.

4.2.5. Determination of the Sf1 repeats and sequence analysis of Sf1 protein

The Sf1 sequences were aligned with HsSf1 and ScSf1 sequences using the ClustalW program [36]. Centrin-binding motifs are known for both HsSf1 and ScSf1, making it possible to determine the centrin-binding motifs of the new Sf1 sequences from the alignments. Hydrophobic cluster analysis (HCA) [37,38] was performed for the new Sf1 proteins, allowing a better determination of their respective repeats. The program for detection of hydrophobic amino acids clusters (HCA) is accessible on the Ressource Parisienne en Bioinformatique (RPBS) server. The programs used to perform protein sequence analyses are accessible on the SWISS-PROT web site [39]. We used GOR4 [40], SOPMA [41] for protein secondary structure predictions.

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Supplementary material

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.jfebsopenbio.2013.11.005.

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