A conserved CAF40-binding motif in metazoan NOT4 mediates association with the CCR4–NOT complex

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The multisubunit CCR4–NOT mRNA deadenylase complex plays important roles in the posttranscriptional regulation of gene expression. The NOT4 E3 ubiquitin ligase is a stable component of the CCR4–NOT complex in yeast but does not copurify with the human or Drosophila melanogaster complex. Here we show that the C-terminal regions of human and D. melanogaster NOT4 contain a conserved sequence motif that directly binds the CAF40 subunit of the CCR4–NOT complex (CAF40-binding motif [CBM]). In addition, nonconserved sequences flanking the CBM also contact other subunits of the complex. Crystal structures of the CBM–CAF40 complex reveal a mutually exclusive binding surface for NOT4 and Roquin or Bag of marbles mRNA regulatory proteins. Furthermore, CAF40 depletion or structure-guided mutagenesis to disrupt the NOT4–CAF40 interaction impairs the ability of NOT4 to elicit decay of tethered reporter mRNAs in cells. Together with additional sequence analyses, our results reveal the molecular basis for the association of metazoan NOT4 with the CCR4–NOT complex and show that it deviates substantially from yeast. They mark the NOT4 ubiquitin ligase as an ancient but nonconstitutive cofactor of the CCR4–NOT deadenylase with potential recruitment and/or effector functions.

[Keywords: deadenylation; mRNA decay; translational repression; ubiquitination]

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The CCR4–NOT complex plays a central role in the post-transcriptional regulation of gene expression by catalyzing the removal of the mRNA poly(A) tail, thereby repressing translation and promoting mRNA degradation (Wahl and Winkler 2013; Collart 2016). In addition, the CCR4–NOT complex has the ability to repress translation independently of deadenylation (Cooke et al. 2010; Chekulaeva et al. 2011; Bawankar et al. 2013).

The CCR4–NOT complex (Fig. 1A) is a multisubunit complex (Chen et al. 2001; Lau et al. 2009; Temme et al. 2010) that assembles on the NOT1 scaffold protein, which consists of several α-helical domains that serve to dock the other subunits of the complex (Bawankar et al. 2013). Deadenylation is carried out by two interacting deadenylases; namely, CCR4 and CAF1. They dock onto a central α-helical domain in NOT1 (labeled “MIF4G”), forming the “catalytic module” of the complex (Basquin et al. 2012; Petit et al. 2012). The C-terminal end of NOT1 contains the NOT1 superfamily homology domain (SHD), which is another α-helical domain that interacts with the NOT2–NOT3 heterodimer to form the “NOT module” of the complex [Bhaskar et al. 2013; Boland et al. 2013]. The catalytic module and the NOT module are connected by the CAF40-binding domain of NOT1 (labeled “CN9BD”) and a connector domain (labeled “MIF4G-C”) of unknown function [Chen et al. 2014; Mathys et al. 2014; Raisch et al. 2018]. Both the NOT module and the CAF40 subunit of the CCR4–NOT complex have been reported as important peptide-docking sites for the recruitment of the complex by mRNA-associated proteins. The NOT module provides binding sites for Bicaudal-C (Chicione et al. 2007), Nanos (Bhandari et al. 2014; Raisch et al. 2016), and Roquin [Sgromo et al. 2017]. CAF40 is known to be contacted by Roquin [Sgromo et al. 2017], Bag of marbles [Ban] [Sgromo et al. 2018], and TTP [Bulbrook et al. 2018] as well as the GW182/TNRC6 family of proteins that mediates microRNA-mediated mRNA repression and decay (Chen et al. 2014; Mathys et al. 2014).
The N-terminal portion of NOT1 is less well conserved than its C-terminal portion [Basquin et al. 2012] and serves to dock NOT10 and NOT11 as additional subunits of the complex in metazoan species [Bawankar et al. 2013; Mauxion et al. 2013]. The CCR4–NOT complexes of *Saccharomyces cerevisiae* (*Sc*) and *Schizosaccharomyces pombe* lack NOT10 and NOT11 proteins. Furthermore, these CCR4–NOT complexes are special because they contain NOT4 as an integral component [Bai et al. 1999; Chen et al. 2001; Nasertorabi et al. 2011; Stowell et al. 2016; Ukleja et al. 2016].

NOT4 (Fig. 1B) is an evolutionarily conserved E3 ubiquitin ligase that contains a RING domain, a linker region with coiled-coil propensity (CC), an RNA recognition motif (RRM) domain, and a C3H1-type zinc finger domain (ZNF). Together, they define the conserved N-terminal region of NOT4 (NOT4-N) (Fig. 1B). The C-terminal region of NOT4 (NOT4-C) (Fig. 1B) is predicted to be unstructured, and its sequence and length are not conserved among NOT4 proteins [The UniProt Consortium 2018]. NOT4 has been implicated in cotranslational mRNA quality control and translational repression in the context of stalled ribosomes, such as in the "No-Go"
mRNA decay pathway (Dimitrova et al. 2009, Matsuda et al. 2014; Panasenko 2014; Preissler et al. 2015; Wu et al. 2018).

A crystal structure demonstrated how Sc Not4 interacts with the SHD of Sc Not1 via an elongated polypeptide from the C-terminal region of Sc Not4 (Bhaskar et al. 2015). Using yeast two-hybrid assays, the human NOT4 and NOT1 proteins [Homo sapiens [Hs] NOT4 and Hs NOT1] were also shown to interact via the C-terminal portion of Hs NOT1 (Albert et al. 2002). However, the Sc Not1-binding sequence of Sc Not4 is only partially conserved, at best (Bhaskar et al. 2015), and NOT4 was not detected in mass spectrometric analyses of the native human and Drosophila melanogaster (Dm) CCR4–NOT complexes (Lau et al. 2009; Temme et al. 2010). This raised the question of whether NOT4 should be regarded as a component or cofactor of the CCR4–NOT complex in metazoans and how it would be recruited to the complex in species other than S. cerevisiae.

We therefore performed pull-down experiments from human cell extracts and with purified bacterially expressed proteins to identify and map interactions of human NOT4 with the CCR4–NOT complex. Assisted by alignments of metazoan NOT4 proteins, we uncovered a 23-amino-acid peptide motif in NOT4-C that binds to the CAF40 subunit of the CCR4–NOT complex and hence was termed the NOT4 CAF40-binding motif [CBM]. Crystal structures of the CBM–CAF40 complex identified critical contacts required in human and Dm S2 cells for an efficient recruitment of NOT4 to the CCR4–NOT complex and for NOT4-mediated mRNA deadenylation and decay via the CCR4–NOT complex. Consequently, NOT4 emerges as a nonconstitutive cofactor of the CCR4–NOT complex in metazoans with a conserved mode of interaction via the CAF40 subunit.

Results

Hs NOT4-C shows a stable interaction with the CCR4–NOT complex

To investigate whether and how Hs NOT4 interacts with the CCR4–NOT complex in HEK293T cells, we expressed Hs NOT4 with a V5-SBP-MBP [V5-streptavidin-binding peptide-maltose-binding protein] tag in HEK293T cells and performed SBP pull-down assays. In agreement with previous reports [Lau et al. 2009; Temme et al. 2010], full-length Hs NOT4 failed to pull down the endogenous CCR4–NOT complex efficiently [Fig. 1C, lane 6]. Hs NOT4-C, however, showed a stable interaction with the CCR4–NOT complex, as indicated by the detection of endogenous NOT1, NOT2, and NOT3 subunits in the pull-down fraction [Fig. 1C, lane 8]. This is consistent with previous yeast two-hybrid experiments [Albert et al. 2002]. In contrast, Hs NOT4-N did not interact with the CCR4–NOT complex [Fig. 1C, lane 7]. The lack of an efficient interaction with the full-length protein remains unexplained but hints at a possible regulation of NOT4-C binding by NOT4-N. Additional SBP pull-down experiments showed that it is the presence of the positively charged CC linker and of the RRM domain in NOT4-N that prevents NOT4-C from interacting with the CCR4–NOT complex [Supplemental Fig. S1A,B].

Tethered Hs NOT4 causes deadenylation-dependent mRNA decay

To address the relevance of an interaction between NOT4 and the CCR4–NOT complex with a functional assay, we investigated the molecular consequences resulting from the presence of NOT4 in the context of an mRNA. Usually, the recruitment of the CCR4–NOT complex to an mRNA target promotes its deadenylation-dependent decay [Wahle and Winkler 2013]. It is unknown, however, whether NOT4 can bind to an mRNA despite the presence of putative and conserved RNA-binding domains in NOT4-N [CC-RRM-ZNF] [Fig. 1B]. In the absence of known mRNA targets, we therefore used a tethering assay to direct NOT4 toward the 3′ untranslated region [UTR] of defined reporter mRNAs and tested them for NOT4-dependent deadenylation and decay.

In a first series of experiments, we tethered MS2-tagged Hs NOT4 to a β-globin mRNA reporter containing six MS2-binding sites in the 3′ UTR [β-globin-6xMS2bs] [Lyykke-Andersen et al. 2000]. Even though full-length Hs NOT4 did not associate with the CCR4–NOT complex in SBP pull-down assays [Fig. 1C, lane 6], we found that tethered MS2-HA–Hs NOT4 caused a substantial reduction of the β-globin-6xMS2bs mRNA level compared with the negative control MS2-HA [Fig. 2A,B]. Tethering Hs NOT4-C also reduced mRNA levels, whereas tethering Hs NOT4-N had no effect [Fig. 2A,B]. All Hs NOT4 fragments were expressed at comparable levels [Fig. 2C], and none of them affected the expression of the control β-globin mRNA lacking MS2-binding sites [Fig. 2B]. Regarding full-length Hs NOT4, the discrepancy with the SBP pull-down assay [Fig. 1C] might be rationalized by a higher sensitivity of the tethering assay for weak and possibly transient interactions or, alternatively, by conformational changes of Hs NOT4 in the presence of RNA that promote the availability of NOT4-C to the deadenylase complex.

In a second series of experiments, we also tethered MS2-tagged Hs NOT4 to another reporter mRNA that encoded Renilla luciferase [R-Luc-6xMS2bs] instead of β-globin, allowing for the quantification of protein abundance in addition to mRNA levels. In agreement with the β-globin mRNA reporter, we observed a clear reduction of both Renilla luciferase mRNA and Renilla luciferase protein levels in the case of tethered Hs NOT4 or Hs NOT4-C [Supplemental Fig. S2A,B]. By comparison, a Renilla luciferase mRNA lacking MS2-binding sites was not affected by the expression of MS2-tagged full-length Hs NOT4 or its fragments [Supplemental Fig. S2C,D].

To verify whether the reduction of reporter mRNA levels upon tethering Hs NOT4 or Hs NOT4-C resulted from deadenylation-dependent decay, we overexpressed a GFP-tagged catalytically inactive mutant of the human mRNA decapping enzyme DCP2 (GFP–DCP2 mut, E148Q). The presence of this mutant is known to impair mRNA decapping in a dominant-negative manner and hence
5′-to-3′ mRNA decay by XRN1 (Loh et al. 2013; Bhandari et al. 2014; Chang et al. 2014; Kuzuoğlu-Öztürk et al. 2016; Sgromo et al. 2017). Indeed, we observed the accumulation of a shorter deadenylated decay intermediate of the β-globin-6xMS2bs reporter mRNA upon tethering Hs NOT4 or upon tethering the Hs Nanos2 mRNA-binding protein (Bhandari et al. 2014), which served as a positive control (Supplemental Fig. S3A–C). We therefore attributed reporter mRNA decay to the recruitment of the CCR4–NOT complex, although we could not formally exclude contributions from other deadenylases at this stage.

**The capacity of NOT4-C to mediate tethered mRNA decay is conserved in metazoans**

The C-terminal region of NOT4 is not conserved in sequence and length (Fig. 1B). We therefore investigated the functionality of Dm NOT4 as an example from an in-vertebrate species and to allow more general conclusions on NOT4 recruitment to the CCR4–NOT complex in metazoans. For Dm NOT4, we used a N-terminal tethering assay in Dm S2 cells with an F-Luc-5BoxB reporter mRNA (Behm-Ansmant et al. 2006). Similar to Hs NOT4 and despite highly divergent sequences of NOT4-C, tethered Dm NOT4 and Dm NOT4-C efficiently mediated reporter mRNA decay (Fig. 2D,E). Dm NOT4 and its fragments were expressed at equal levels (Fig. 2F), and none of them affected the expression of an F-Luc reporter lacking the BoxB sites (Supplemental Fig. S2E,F).

Again, reporter mRNA decay was deadenylation-dependent. Deadenylated F-Luc-5BoxB reporter mRNA was stabilized in the presence of tethered Dm NOT4, when a GFP-tagged Dm DCP1 mutant (GFP-DCP1 mut; R70G, L71S, N72S, and T73G) known to prevent mRNA decapping in a dominant-negative manner (Chang et al. 2014; Kuzuoğlu-Öztürk et al. 2016) was overexpressed (Supplemental Fig. S3D–F). Tethered GW182 protein was used
as a positive control (Kuzuoğlu-Öztürk et al. 2016). Together, our results indicate that the interaction of NOT4-C with the CCR4–NOT complex is conserved between humans and flies.

**Hs NOT4 directly interacts with the NOT1 and CAF40 subunits of the CCR4–NOT complex**

To test whether the interaction between Hs NOT4 and the CCR4–NOT complex is direct, we expressed MBP-tagged Hs NOT4, Hs NOT4-N, and Hs NOT4-C in Escherichia coli and performed pull-down experiments with a reconstituted and purified subcomplex of human CCR4–NOT components that we had described previously (Sgromo et al. 2017) and that we here call the “pentameric” complex (Figs. 1A, 3A). This subcomplex comprises the C-terminal portion of NOT1 together with the CAF1 and CAF40 subunits and the C-terminal fragments of NOT2 and NOT3. Indeed, we observed a direct interaction with the pentameric complex. In contrast to the result from the SBP pull-down experiment in HEK293T cells (Fig. 1C, lane 6), the interaction occurred even with the full-length Hs NOT4 (Fig. 3A, lane 7). Furthermore, Hs NOT4-C interacted with the pentameric complex as efficiently as the recombinant full-length protein (Fig. 3A, lane 9), whereas recombinant Hs NOT4-N did not interact (Fig. 3A, lane 8).

To map the interactions to individual components of the CCR4–NOT complex, we also used previously described smaller subassemblies of the complex (Petit et al. 2012; Boland et al. 2013; Chen et al. 2014; Sgromo et al. 2017) in MBP pull-down experiments (Figs. 1A, 3B). These were the NOT1–NOT10–NOT11 module (comprising the N-terminal region of NOT1 and NOT10 and the C-terminal half of NOT11), CAF1 bound to the CAF40-binding domain of NOT1 (labeled “CN9BD”), the NOT1 connector domain (labeled “MIF4G-C”), and the NOT module (comprising the C-terminal regions of NOT1, NOT2, and NOT3). We detected interactions of Hs NOT4 with both the CAF40-containing subcomplex and the NOT module (Figs. 3B, lanes 13, 17), pointing to multiple NOT4-binding sites within the CCR4–NOT complex. Finally, we narrowed down these interactions even further, primarily to the NOT1-SHD (Fig. 3C, lane 7) and CAF40 alone (Fig. 3C, lane 11), with only minor contributions from the NOT2–NOT3 heterodimer (Fig. 3C, lane 9).

**The central C2 region of Hs NOT4-C is sufficient to contact CCR4–NOT and trigger decay of tethered mRNA**

It is not uncommon for the long unstructured portions of regulatory mRNA-associated proteins (such as in GW182/...
To identify a potential sequence motif that could be responsible for the interaction of the C2 region with the NOT module and CAF40, we generated and analyzed separate alignments of NOT4 proteins from metazoans, plants, and yeasts. These alignments revealed locally conserved sequences at different positions in the C-terminal regions of NOT4 [Fig. 1B; Supplemental Figs. S4, S5A; Supplemental Alignment Files SF1–SF3].

Most striking is the conservation of a 23-amino-acid motif with α-helical propensity in the C2 region of HS NOT4, which is present throughout all metazoans [Fig. 1B; Supplemental Fig. S4]. A similar sequence also exists in plants but is not found in fungi. Conversely, the NOT1-binding motif of Sc Not4 [Bhaskar et al. 2015] is conserved in yeast but cannot be identified in plants and metazoans [Supplemental Fig. S5A; Supplemental Alignment Files SF1–SF3].

Beside the 23-amino-acid motif, the alignments also uncovered the presence of a proline-rich PPPGΦ motif (Φ = F, L, or I) at least once in each of the NOT4 sequences from metazoans, plants, and yeasts. The position of the PPPGΦ motif within NOT4-C varies with the phylogeny and can be found before or after the 23-amino-acid motif with the potential for misaligning distantly related sequences [Supplemental Figs. S4, S5A; Supplemental Alignment Files SF1–SF3].PPPΦ motifs are known to be recognized by proteins containing proline-binding GYF domains [Koller and Freund 2006].

The 23-amino-acid motif [residues E400–E422] occupies the second half of the C2 region [Supplemental Fig. S5B]. We therefore again performed MBP pull-down experiments with the NOT module and CAF40, where either the first half (C2a) or the second half (C2b) of the C2 region or the entire C2 region was deleted from HS NOT4-C. These deletions did not detectably affect the interaction with the NOT module, indicating that the C1 and C3 regions of HS NOT4-C together are still sufficient to pull down the NOT module [Supplemental Fig. S5C, lanes 9–12]. However, the interaction with CAF40 was clearly diminished by deleting the second half of the C2 region [Supplemental Fig. S5C, lanes 15,17]. This observation suggests that the 23-amino-acid motif acts as a conserved CBM. This assumption was confirmed in the following by X-ray crystallography. The motif is hence called the NOT4 CBM.

Crystal structure of the CBM of Dm NOT4 in complex with HS CAF40

To understand in molecular detail how NOT4 interacts with the CCR4–NOT complex, we used peptides corresponding to the putative CBM of HS NOT4 or Dm NOT4 to set up cocrystallization trials with HS CAF40. We obtained crystals—but only of a heterologous complex consisting of HS CAF40 and the CBM of Dm NOT4. We obtained two distinct crystal forms, each with two crystallographically independent complexes per asymmetric unit and diffracting X-rays to a maximum resolution of 2.1 Å (Table 1). CBM binding is highly similar among the four CAF40–CBM complexes, and therefore only one of them [polypeptide chains A and B from space group P2₁2₁2₁] is described [Fig. 4A–C; Supplemental Figs. S5D, E, S6].

From Dm NOT4 L816 to E835 [see Fig. 4D for human sequence numbers], the Dm CBM adopts a common conformation in all of the available structures, with Dm NOT4 P820 to E835 folding into four turns of an amphipathic α helix that is bent between turns two and three toward the surface of HS CAF40. Only the very N-terminal and very C-terminal residues of the crystallized peptide show differing orientations in the four available complexes, probably due to crystal-packing interactions and indicating structural flexibility [Supplemental Fig. S5E]. HS CAF40 adopts its rigid and previously described crescent-like shape, made from six armadillo repeats. Despite its highly negative overall charge (pI = 3.6), the Dm CBM interacts with HS CAF40 primarily via hydrophobic contacts. It engages the concave surface of HS CAF40, contacting residues from three parallel α helices (α5, α8, and α11) and burying a surface on HS CAF40 of 842 Å² [Fig. 4A–C].

Importantly, the very same surface of CAF40 was described previously to be engaged also by the CBMs of Roquin and Bam [Fig. 4E,F; Sgromo et al. 2017, 2018]. These CBMs also fold into amphipathic helices, covering surface areas of 841 Å² and 748 Å², respectively, and excluding any simultaneous associations of multiple CBMs with CAF40. Strikingly, however, whereas the CBMs of Roquin and Bam run in parallel to the α helices α5, α8, and α11, the CBM of NOT4 runs in an antiparallel fashion and hence is structurally and evolutionarily unrelated to the other two CBMs.
Details of the interaction between the CBM of Dm NOT4 and Hs CAF40

The amphipathic α helix of the Dm NOT4 CBM is preceded by an “LGFDP” sequence motif that is invariant in our alignment of metazoan species (Fig. 4D). This sequence motif by itself forms a characteristic structure that helps to pin down the N-terminal half of the α helix (Fig. 4G, H). Probably due to the backbone flexibility provided by Dm NOT4 G817, the side chains of Dm NOT4 L816, F818, and P820 can join to form a small hydrophobic cluster, which is centrally contacted and completed by V181 from the α helix α11 of Hs CAF40. Furthermore, V181 is assisted by F184 to fix Dm NOT4 L816 and by L177 to fix Dm NOT4 F818, allowing Dm NOT4 L816 and F818 to intercalate between the side chains on the α helix α11 of Hs CAF40. Dm NOT4 P820 also initiates the α helix of NOT4 and is assisted by Dm NOT4 D819, which caps the helix and compensates for the positive charge of the helix dipole. Moreover, P820 is positioned straight over G141 in the α helix α8 of Hs CAF40 and would be spatially incompatible with any other residue at this position apart from a glycine.

Following P820, the hydrophobic surface of the NOT4 α helix probes the groove between α helices α8 and α5 of Hs CAF40 [Fig. 4D,H,J] using side chains of Dm NOT4 F821, T824, L828, and L831. These side chains are lined by residues from α helices α8 (Hs CAF40 T138, L137, Y134, and P131) and α5 (Hs CAF40 A95, N92, C91, N88, S87, H85, and A84), all of which are within van der Waals distance and frequently contribute to the interactions with the aliphatic portions of their side chains. At the C-terminal end of the CBM [Fig. 4D,H,J], the invariant Dm NOT4 E835 pins down the C-terminal half of the NOT4 α helix, using two hydrogen bonds to coordinate the free main chain nitrogens of Hs CAF40 A84 and H85 at the beginning of α helix α5 and compensating for the positive charge of the helix dipole. As a consequence of this interaction, the side chain of A84 gets completely surrounded by residues from NOT4 (Dm NOT4 L828, L831, M832, and E835), tolerating no side chains at this position that are larger than alanine. Additional specificity arises from an H bond between the side chains of Dm NOT4 T824 and Hs CAF40 T138 on α helix α8, which is deeply buried in the interface [Fig. 4D,H]. Finally, it is important for Hs CAF40 Y134 on α helix α8 to rotate away from its preferred

| Table 1. Data collection and refinement statistics for the Hs CAF40–Dm NOT4 CBM complex |
|----------------------------------|----------------------------------|
| Space group                      | P 2 1 2 1 2                     |
| Unit cell                        | I 2 1 2 1 2                     |
| Dimensions $a$, $b$, $c$          | 83.9 Å, 109.6 Å, 69.7 Å        |
| Angles $\alpha$, $\beta$, $\gamma$ | 90.0°, 90.0°, 90.0°            |
| Data Collection$^a$              |                                 |
| Wavelength                       | 1.0000 Å                       |
| Resolution range                 | 50 Å–2.1 Å (2.14 Å–2.10 Å)     |
| $R_{\text{sym}}$                 | 7.0% (91.4%)                   |
| Completeness                     | 99.6% (99.8%)                  |
| Mean $I/\sigma(I)$               | 12.2 [1.5]                     |
| CC 1/2                           | 99.0 [55.6]                    |
| Unique reflections               | 38,143 [2780]                  |
| Multiplicity                     | 4.0 [4.2]                      |
| Refinement                       |                                 |
| $R_{\text{work}}$                | 19.1%                          |
| $R_{\text{free}}$                | 22.2%                          |
| Number of atoms                  |                                 |
| All atoms                        | 4971                           |
| Protein                          | 4760                           |
| Ligands                          | 34                             |
| Water                            | 177                            |
| Average B factor                 |                                 |
| All atoms                        | 52.7 Å$^2$                     |
| Protein                          | 52.3 Å$^2$                     |
| Ligands                          | 95.5 Å$^2$                     |
| Water                            | 53.7 Å$^2$                     |
| Ramachandran plot                |                                 |
| Favored regions                  | 99.2%                          |
| Disallowed regions               | 0.0%                           |
| RMSD from ideal geometry         |                                 |
| Bond lengths                     | 0.002 Å                        |
| Bond angles                      | 0.426$^a$                      |

$^a$Values in parentheses are for the highest-resolution shell. 
(RMSD) Root mean square deviation.
rotamer position that is observed in free CAF40 (Garces et al. 2007; Chen et al. 2014; Mathys et al. 2014) and in the complexes with Roquin (Sgromo et al. 2017) and Bam (Sgromo et al. 2018). Hence, the orientation of Hs CAF40 with the CBMs of Dm Roquin [E; Protein Data Bank [PDB] ID 5lsw] (Sgromo et al. 2017) and Dm Bam [F; PDB ID 5onb] (Sgromo et al. 2018), shown in the same style and orientation as in C and excluding structurally variable flanks of the CBMs. Note the inverted orientation of the CBMs. (C–I) Close-up views of the interface between Dm NOT4 and Hs CAF40. Selected side chains of NOT4 and CAF40 are shown as blue and orange sticks, respectively, with nitrogens in dark blue and oxygens in red. Hydrogen bonds are indicated by red dashed lines. Residues mutated in this study are underlined. (J) Rotamers of CAF40 Y134 as found in the complexes of CAF40 with the CBMs of Dm Roquin (yellow) and Dm Bam (lime).

Validation of the binding interface

To validate the specificity of the interface observed in the crystal structure, we generated mutations in the CBM of Dm NOT4 and in Hs CAF40 and tested them in MBP pull-down assays. First, we demonstrated that the CBM of Dm NOT4 indeed also interacts with Hs CAF40 in solution [Fig. 5A, lane 14], confirming that it is a bona fide interaction. To validate the specificity of the interface observed in the crystal structure, we generated mutations in the CBM of Dm NOT4 and in Hs CAF40 and tested them in MBP pull-down assays. First, we demonstrated that the CBM of Dm NOT4 indeed also interacts with Hs CAF40 in solution [Fig. 5A, lane 14], confirming that it is a bona fide interaction. We disrupted the interaction from the side of CAF40 using either a single point mutation targeted at the “LGFDP” motif [Hs CAF40 V181E] [Fig. 4G] or a double point mutation targeted at the α-helical part of the CBM [Hs CAF40 2x mut; Y134D and G141W] [Fig. 4G,I].
case of the Dm Roquin CBM [Sgromo et al. 2018], we could not determine a precise dissociation constant for the Dm NOT4 CBM because it aggregated at concentrations needed to perform isothermal titration calorimetry and microscale thermophoresis experiments.

Finally, we also tested the interaction of the Hs NOT4 CBM with Hs CAF40. Similar to the results obtained with the Dm NOT4 CBM, the single Hs CAF40 V181E mutation [Fig. 4G] was sufficient to abolish the interaction with the Hs NOT4 CBM [Fig. 5C, lane 15 vs. 14]. The double point mutation [Hs CAF40 2x mut] [Fig. 4G,I] also reduced the interaction but was not sufficient to abolish it, indicating species-specific adaptations in the molecular details of the coevolved interface of the human proteins [Fig. 5C, lane 16 vs. 14]. Together, the results confirm that the interactions observed in the crystal structure also occur in solution.

The CBM is essential for the interaction of NOT4 with the human CCR4–NOT complex

To investigate the significance of the CBM–CAF40 interaction in cells and in the context of the entire CCR4–NOT complex, we repeated SBP pull-down assays with Hs NOT4-C where either the CBM [ΔC2b], the C2a region [ΔC2a], or the entire C2 region [ΔC2] were deleted from the construct (see Supplemental Fig. S5B for boundaries). We found that the presence of the CBM is required to pull down the CCR4–NOT complex [Fig. 6A, lane 7], as detected by the absence of NOT1, NOT2, NOT3, and CAF40 in the pull-down fraction when the C2b region was deleted [Fig. 6A, lane 10]. Consistently, the deletion of the entire C2 region as well disrupted the interaction [Fig. 6A, lane 8]. Surprisingly however, the presence of the C2a region was also required for an efficient pull-down of the CCR4–NOT complex [Fig. 6A, lane 9]. The C2a region could simply act as a spacer to allow for a proper interaction of the CBM with CAF40. More likely, however, its requirement reflects and underlines the importance of auxiliary and possibly species-specific interactions of NOT4-C with other parts of the CCR4–NOT complex, such as the NOT module. In line with these results, the deletion of the entire C2 region strongly diminished the NOT4-mediated degradation of the β-globin-6xMS2bs reporter mRNA in HEK293T cells [Fig. 6B,C]. The separate deletion of the C2a region [ΔC2a] or the CBM [ΔC2b] did so as well but to a lesser degree. All tethered proteins were expressed at a similar level [Fig. 6D].

To probe more directly for the interface observed in the crystal structure, we engineered a quadruple point mutant of the Hs NOT4 CBM (4x mut; F405D, F408A, T411E, and L415E) [Fig. 4D] affecting both the LGFDP motif and the hydrophobic surface of the α helix. This mutation strongly reduced the interaction with the CCR4–NOT complex in the SBP pull-down assay [NOT4-C 4x mut] [Fig. 6E], whereas individual point mutations were not as effective [Supplemental Fig. S7A]. Also in the tethering assay, the quadruple point mutation [NOT4 4x mut] impaired the ability of NOT4 to degrade the β-globin-6xMS2bs reporter
mRNA to a degree that is comparable with the deletion of the entire CBM [Fig. 6, F,G vs. B,C]. All of the tethered proteins were equally expressed [Fig. 6H]. These findings demonstrate the importance of the CBM for the function of NOT4 and indicate that this is a major site of interaction with the CCR4–NOT complex.

**CAF40 plays a crucial and conserved role for the recruitment of NOT4 to the CCR4–NOT complex in metazoans**

The present crystal structures show how the CBM interacts with CAF40, but our previous experiments did not formally exclude that the CBM makes similarly important contacts with other subunits of the CCR4–NOT complex. Therefore, we took advantage of HEK293T cells in which CAF40 had been knocked out by CRISPR–Cas9 genome editing [CAF40 knockout cells] [Fig. 7A–D; Supplemental Fig. 7B–D; Sgromo et al. 2018]. In this cell line, the levels of endogenous NOT1, NOT2, and NOT3 proteins were not altered [Fig. 7B], but the NOT4-mediated decay of the tethered R-Luc-6xMS2bs mRNA reporter [Fig. 7A,D] or of the tethered β-globin-6xMS2bs mRNA reporter [Supplemental Fig. 7B,C] was impaired. This observation underlines the importance of CAF40 for the efficient recruitment of the CCR4–NOT complex.

To test whether the remaining mRNA repression in CAF40 knockout cells [Fig. 7A; Supplemental Fig. 7B,C]...
CAF40 is a central mediator for the interaction of NOT4 with the CCR4–NOT complex in HEK293T and Dm S2 cells. (A) Tethering assay with Hs NOT4 and a luciferase reporter in HEK293T cells lacking CAF40 (CAF40 knockout [KO]) (Sgromo et al. 2018). Renilla luciferase mRNA served as a reporter and contained six binding sites for the MS2 protein [R-Luc-6xMS2bs]. Firefly luciferase mRNA served as a reference and transfection control [F-Luc]. R-Luc activity was normalized to the reference and plotted with respect to the values obtained from the expression of MS2-HA alone (set to 100). The shRNA-mediated depletion of the CCR4–NOT complex is indicated by white bars [sh-NOT1 RNA and NOT1 knockdown [KD]] as compared with black bars [sh-scr RNA, control]. Error bars correspond to standard deviations. n = 3. (B) Efficiency of CRISPR–Cas9-mediated gene editing of CAF40. The Western blot shows CAF40 knockout cells in comparison with a dilution series of wild-type HEK293T cells using tubulin as a loading control. (C) Efficiency of shRNA-mediated depletion of NOT1. The Western blot shows HEK293T cells expressing sh-NOT1 RNA in comparison with a dilution series of HEK293T cells expressing sh-scr RNA using PABP as a loading control. (D) Tethering assay with the CBM mutation variant of Hs NOT4 (4x mut) and the luciferase mRNA reporter in CAF40 knockout cells. Error bars correspond to standard deviations. n = 3. (E–G) Tethering assay with Dm NOT4 in Dm S2 cells overexpressing Dm CAF40 with a mutated CBM-binding surface. Experiments were done with AN-HA-tagged Dm NOT4 or Dm NOT4-C and analyzed as in Figure 2D, but cells were additionally overexpressing either wild-type Dm CAF40 or mutant Dm CAF40 (GFP-CAF40 V186E, corresponding to Hs CAF40 V181E). GFP-V5 was overexpressed as a negative control. (E) Relative protein and mRNA levels, with error bars corresponding to standard deviations. n = 3. (F) Northern blot. (G) Western blot. The asterisk denotes the additional detection of GFP-V5 on the anti-V5 blot.
was due to the CCR4–NOT complex, we additionally disrupted and depleted the remainder of the complex by a shRNA-mediated knockdown of NOT1 [Fig. 7C; Supplemental Fig. S7D; Boland et al. 2013]. Under these conditions, tethering of NOT4 left reporter mRNA expression almost unaffected [CAF40 knockout + NOT1 knockdown] [Fig. 7A; Supplemental Fig. S7B,C]. Consequently, the remaining subunits of the complex in CAF40 knockout cells still interact with tethered NOT4, which seems to act exclusively via the CCR4–NOT complex.

Finally, there was only a small difference between tethering the quadruple point mutation of NOT4 [NOT4 4x mut] and tethering wild-type NOT4 in CAF40 knockout cells, whereas this difference was considerable in wild-type cells [Fig. 7D]. This result confirms CAF40 as the primary interaction partner of the NOT4 CBM in human cells.

The conservation of the CBM suggests that the interaction of NOT4 with CAF40 is preserved in metazoans, albeit modulated by additional contacts, such as with the NOT1-SHD [Fig. 3C]. Likely due to such taxon-specific or species-specific modulation, it was therefore possible in Dm S2 cells to obtain a dominant-negative effect on CCR4–NOT recruitment by overexpressing a V186E mutant of GFP-tagged Dm CAF40 [Fig. 7E–G]. This mutant (corresponding to Hs CAF40 V181E) was also shown previously to impair CCR4–NOT recruitment by the CBM of Bam [Sgromo et al. 2018]. Apparently, in this case, the overexpression of GFP-tagged CAF40 can functionally replace the endogenous protein, and a single point mutation is then sufficient to disrupt the interaction with NOT4. Again, these observations demonstrate the central and conserved role of the CBM for the recruitment of NOT4 to the CCR4–NOT complex.

Discussion

The present work demonstrates that metazoan NOT4 contains a conserved CBM in its variable C-terminal tail and elucidates the molecular details of the CBM–CAF40 complex. The CBM is required in cells for an efficient recruitment of NOT4 to the CCR4–NOT complex or a recruitment of the complex to NOT4-mediated cellular processes. The interaction of the CBM is assisted by auxiliary flanking sequences in NOT4-C that vary between metazoan species. These sequences also contact other subunits of the CCR4–NOT complex, such as the SHD of NOT1. From an evolutionary point of view, the CBM therefore appears to represent the ancestral mode of coupling the NOT4-dependent E3 ubiquitin ligase activity with the CCR4- and CAF1-dependent deadenylation activity of the CCR4–NOT complex. In yeast, however, the CBM seems to have become dispensable, possibly because the contacts with NOT1 are sufficient to maintain the interaction [Bhaskar et al. 2015]. The conservation of the CBM marks NOT4 as a ubiquitous but apparently facultative cofactor of the metazoan CCR4–NOT complex that likely has important functions in a subset of CCR4–NOT-dependent cellular processes.

Mutual corecruitment of NOT4 and the CCR4–NOT complex

The widespread conservation of the CBM in NOT4 proteins reveals NOT4 as an ancient cofactor of the CCR4–NOT complex. Furthermore, we show that NOT4 is able to cause CCR4–NOT-mediated mRNA decay if tethered to an mRNA target. However, it remains unclear from these experiments whether metazoan NOT4 is needed primarily as an upstream recruitment factor that directs the CCR4–NOT complex to selective mRNA targets or rather as a downstream effector that recruits additional proteins to the CCR4–NOT complex and/or ubiquitinates nearby protein targets, e.g., to mark them for proteasomal degradation. In contrast to selective mRNA-binding proteins such as TTP, Nanos, or Roquin [Newman et al. 2016], there are currently no known RNA targets for the coiled-coil linker, RRM, or ZNF domains of NOT4, i.e., for its putative RNA-binding domains. This argues against an RNA-specific recruitment function.

In the context of cotranslational mRNA quality control, however, NOT4 could act as both an upstream recruitment factor of the CCR4–NOT complex to mRNAs with stalled ribosomes and a downstream effector for the ubiquitination and degradation of protein targets [Panasenko
Additional protein-binding partners may modulate or stabilize the interactions in this case.

Finally, it is worthwhile to follow up also on the PPPGΦ motifs that we found to be highly conserved in the NOT4 proteins from metazoans, plants, and yeasts. PPPGΦ motifs tend to interact with GYF domain proteins [Koller and Freund 2006], such as the GIGYF1/2 translational repressors [Krysaszke et al. 2016; Peter et al. 2017; Amaya Ramirez et al. 2018] that were described to bind CAF40 in human cancer cells [Ajiro et al. 2009]. Quite likely, therefore, the CCR4–NOT complex and NOT4 frequently support each other in a mutual corecruitment that is difficult to disentangle experimentally.

Competition of mRNA-associated proteins for the CBM-binding site of CAF40

In most of the known cases where CCR4–NOT gets recruited to an mRNA target, the CBM-binding surface on CAF40 appears to remain available for a simultaneous recruitment of NOT4. This is, for example, the case for the TNRC6/GW182-mediated microRNA-dependent mRNA regulation, where tryptophans of TNRC6/GW182 bind to the convex side of CAF40 [Chen et al. 2014; Mathys et al. 2014]. Similarly, mRNA-specific CCR4–NOT recruitment proteins such as TTP or Nanos apparently do not structurally interfere with NOT4 binding to the concave surface of CAF40 [Fabian et al. 2013; Bhandari et al. 2014; Bulbrook et al. 2018], allowing for a combinatorial mRNA regulation.

In contrast, the CBMs of Roquin and Bam were shown to target the exact same binding surface on CAF40 as the CBM of NOT4 [Sgromo et al. 2017, 2018], making their binding mutually exclusive. It is therefore possible that Roquin proteins have evolved to displace NOT4 in a context-specific manner, since they bring along their own E3 ubiquitin ligase domain. Conversely, in the case of Bam, the CBM might serve to prevent NOT4-mediated and ubiquitination-dependent processes downstream from CCR4–NOT recruitment in the germline of D. melanogaster. Future work will show whether such competition indeed occurs in vivo and whether there are additional CBM-containing mRNA-binding proteins in fungi, plants, or metazoans that operate in a similar manner.

Clearly, however, the present identification of a conserved CBM in the NOT4 protein underlines the role of CAF40 as a hub for peptide-mediated interactions and adds to the ever more complex regulation of mRNA expression in eukaryotic cells.

Materials and methods

DNA plasmid constructs

For bacterial expression of recombinant Hs NOT4 constructs in E. coli, cDNA sequences were inserted between the Xhol and Nhel restriction sites of the pNEA-pM plasmid, resulting in fusion proteins carrying N-terminal MBP tags cleavable by the human rhinovirus 3C [HRV3C] protease and, in addition, C-terminal GB1 and hexahistidine tags. For bacterial expression of recombinant Dm NOT4 constructs, cDNA sequences were inserted between the Xhol and BamHI restriction sites of the pνYC-vM plasmid, resulting in tobacco etch virus (TEV) protease-cleavable MBP fusion proteins. For the expression of Hs NOT4 constructs in human [HEK293T] cells, cDNA sequences were inserted into the pCIneo-V5-SBP-MBP plasmid or the pcDNA3.1-MS2-HA plasmid using the NotI and BstBI restriction sites. For the expression of Dm NOT4 constructs in Dm S2 cells, cDNA sequences were inserted into the pAc5.1-BN-AN-HA plasmid using the NotI and BamHI restriction sites. All of the plasmid constructs generated in this study, including backbone references, are listed in Supplemental Table S1.

MBP pull-down assays with bacterially expressed proteins

For initial pull-down assays with full-length Hs NOT4 and its fragments, the proteins were expressed in E. coli BL21 [DE3] Star cells [Invitrogen] overnight in LB medium at 20°C. Cells were homogenized in lysis buffer (50 mM Na/HEPES at pH 7.5, 300 mM NaCl, 5 µg/mL DNaseI, 1 mg/mL lysozyme, Roche “Complete” EDTA-free protease inhibitors) supplemented with 20 mM imidazole and 2 mM β-mercaptoethanol. The proteins were immobilized and isolated from the lysate on Ni-NTA resin (Qiagen) followed by elution in lysis buffer supplemented with 500 mM imidazole. They were then immobilized on 50 µL of amyllose resin and incubated with an excess of purified CCR4–NOT proteins for 1 h in 500 µL of binding buffer (50 mM Na/HEPES at pH 7.5, 300 mM NaCl, 2 mM β-mercaptoethanol). Finally, the amyllose beads were washed five times with binding buffer, and the proteins were eluted in 50 µL of binding buffer supplemented with 25 mM D(+) -maltose.

For pull-down assays with Hs and Dm NOT4 CBM constructs, proteins were purified from cells homogenized in lysis buffer supplemented with 2 mM DTT. Proteins were immobilized and isolated from the lysate on an amyllose resin [New England Biolabs] followed by anion exchange chromatography over a HiTrap Q column (GE Healthcare) and size exclusion chromatography over a Superdex 200 26/600 column (GE Healthcare) in a buffer containing 10 mM Na/HEPES (pH 7.5), 200 mM NaCl, and 2 mM DTT. Forty micrograms of purified MBP-tagged NOT4 fragments or 20 µg of MBP were then incubated with approximately equimolar amounts of the respective purified CCR4–NOT proteins and 50 µL of amyllose resin in 500 µL of binding buffer (50 mM Na/HEPES at pH 7.5, 200 mM NaCl, 2 mM DTT). After the incubation and washing steps, the proteins were eluted in 200 µL of binding buffer supplemented with 25 mM D(+) -maltose and precipitated with trichloroacetic acid.

The purifications of other human CCR4–NOT proteins, including Hs CAF40 for crystallization, were described previously [Petit et al. 2012; Boland et al. 2013; Bhandari et al. 2014; Chen et al. 2014; Raisch et al. 2016; Sgromo et al. 2017, 2018]. The protein samples were resolved and analyzed by SDS-PAGE.

Crystallization

Hs CAF40 [GPHMLE-R19–E285] [Chen et al. 2014] was mixed with a twofold molar excess of the Dm NOT4 CBM peptide [D813–Q838, chemically synthesized and purchased from EMC Microcollections] in 10 mM Na/HEPES [pH 7.5], 200 mM NaCl, and 2 mM DTT. Initial screens were carried out in sitting drops at 22°C by mixing 200 nL of sample solution [6 mg/mL CAF40, 1.2 mg/mL NOT4] with 200 nL of reservoir solution. Crystals appeared within 1 d in many conditions. Crystals of crystal form 1 appeared in the initial screen over a reservoir
containing 0.2 M ammonium sulfate, 0.1 M Bis-Tris/Cl (pH 5.5), and 25% (w/v) PEG 3350. The crystals were cryoprotected in reservoir solution supplemented with 25% glycerol and flash-cooled in liquid nitrogen. Optimized crystals of crystal form 2 grew at 18°C in hanging drops mixing 1 µL of sample solution and 1 µL of reservoir liquid nitrogen. Optimized crystals of crystal form 2 grew at 18°C in hanging drops mixing 1 µL of sample solution and 1 µL of reservoir solution containing 0.9 M K2HPO4 and 0.3 M NaH2PO4. Crystals were cryoprotected in 4.0 M sodium formate and flash-cooled in liquid nitrogen.

**Data collection and structure determination**

X-ray diffraction data were collected at a wavelength of 1.000 Å on a Pilatus 6M detector (Dectris) at the PXII beamline of the Swiss Light Source (Villigen) and processed using XDS and XSCALE (Kabsch 2010). Crystal form 1 (space group P212121) diffracted X-rays to a resolution of 2.1 Å, whereas crystal form 2 (space group I212121) diffracted X-rays to a comparable resolution of 2.2 Å but with an increased Bweight of 56.7 Å² vs. 39.2 Å². For each crystal form, we identified two copies of two sets of 435¢sa per asymmetric unit by molecular replacement using Phaser (McCoy et al. 2007) from the CCP4 package (Winn et al. 2011) and using chain A of Protein Data Bank (PDB) ID 2vf2 (Garces et al. 2007) as a search model. Initial models were improved and completed by iterative cycles of model building in COOT (Emsley et al. 2010) and refinement using PHENIX (Afonine et al. 2012). The NOT4 CBM peptides were then built manually into the remaining electron density and improved by additional building and refinement cycles. For crystal form 1, final refinement rounds were done using PHENIX with one TLS group per polypeptide chain and including small molecule ligands (one molecule each of Tris and glycerol plus four sulfate ions) in addition to 177 water molecules. This resulted in an Rwork of 19.0% and an Rfree of 21.8%. For crystal form 2, final refinement rounds were done using BUSTER (https://www.globalphasing.com/buster), also with one TLS group per polypeptide chain but in addition to small molecule ligands (one sodium and two chloride ions) and 64 water molecules, also autorrefining NCS restraints. This resulted in an Rwork of 19.3% and an Rfree of 23.0% (Table 1). Illustrations were prepared in PyMOL (http://www.pymol.org).

**SBP pull-down assays from HEK293T cells**

HEK293T cells were seeded in 10-cm dishes (4 x 10^6 cells per plate and experiment) and transfected with pCIneo-V5-SBP-MBP-plasmids after 1 d using Turbofect (Thermo Scientific) according to the manufacturer’s protocol. Two days after transfection, the cells were lysed on ice in 1 mL of NET lysis buffer (50 mM Tris/Cl at pH 7.5, 150 mM NaCl, 0.1% Triton X-100, 1 mM EDTA, 10% glycerol) supplemented with protease inhibitors (Roche). After 15 min, lysates were centrifuged at 20,000 g for 15 min at 4°C. The cleared lysate was then treated with 200 µg/mL RNase A (Qiagen) for 30 min at 4°C and centrifuged again as before, resulting in the input fraction for the experiment (1 mL = 100%). The input fraction was then incubated for 1 h at 4°C with 50 µL of streptavidin sepharose resin (Genecart Healthcare). The beads were washed three times with NET buffer and finally resuspended in protein sample buffer, resulting in the pull-down fraction (100 µL = 100%). The samples were analyzed by Western blot (for antibodies, see Supplemental Table S2) using the ECL Western blotting detection system (Gene Healthcare).

**Tethering assays in HEK293T cells**

For MS2-dependent tethering assays with the β-globin mRNA reporter (Lykke-Andersen et al. 2000), HEK293T cells were seeded in six-well plates (0.7 x 10^6 cells per well) and transfected on the following day using Lipofectamine 2000 (Invitrogen). The transfection mixtures contained 0.5 µg of the β-globin reporter plasmid encoding six MS2-binding sites (β-globin-6xMS2bs), 0.5 µg of the β-globin reference and transfection control plasmid lacking MS2-binding sites and containing a partial sequence of GAPDH (control; β-globin-GAPDH) (Lykke-Andersen et al. 2000); and variable amounts (0.05-0.75 µg) of pcDNA3.1-MS2-HA plasmids (Supplemental Table S1) to achieve equivalent expression of MS2-HA fusion proteins. The cells were harvested 2 d after transfection. The total RNA was isolated using the RneGold TriFast reagent (Peqlab) and analyzed by Northern blot as described previously (Behm-Ansманt et al. 2006). Equivalent expression of MS2-HA-tagged proteins was tested in parallel by Western blot, expressing F-Luc-GFP (Kuzuoglu-Ozturk et al. 2016) as a transfection control.

For the experiment shown in Supplemental Figure S3, A-C, cells were additionally cotransfected with 0.5 µg of a plasmid expressing either wild-type Hs DCP2 (GFP-DCP2 wild-type) or the Hs DCP2 mutant (GFP-DCP2 mut; E148Q) (Loh et al. 2013). Equivalent expression of the GFP-tagged proteins was tested in parallel by Western blot, expressing V5-SBP-MBP as a transfection control.

For MS2-dependent tethering assays with the luciferase reporter system [Kuzuoglu-Ozturk et al. 2016], the transfection mixtures contained 0.2 µg of reporter plasmid containing or lacking six MS2-binding sites (R-Luc-6xMS2bs or R-Luc), 0.2 µg of reference and transfection control plasmid lacking six MS2-binding sites (F-Luc-GFP), and variable amounts of pcDNA3.1-MS2-HA plasmids (0.1–1.5 µg; Supplemental Table S1). The cells were harvested 2 d after transfection, mRNA levels were determined by Northern blot, and L-Ruc and F-Luc activities were measured using a “dual-luciferase reporter assay” system (Promega).

**Tethering assays in HEK293T cells with knockdown of NOT1**

The shRNA-mediated depletion of NOT1 has been described previously [Boland et al. 2013] using shRNA (Hs NOT1 target: ATTCAACATTCCTCCITTATA) and control shRNA (scr; control target: ATTTCTCCGAACGTGTCAGC). For tethering assays in cells depleted of NOT1, wild-type HEK293T cells or HEK293T CAF40 knockout cells (Sgromo et al. 2018) were transfected twice. For the first transfection, cells were seeded in six-well plates (0.7 x 10^6 cells per well) and transfected on the following day with mixtures containing 4 µg of plasmid expressing the respective shRNA. After 1 d, cells were selected for 24 h in DMEM supplemented with 1.5 µg/mL puromycin and subsequently seeded in six-well plates in medium without puromycin (0.7 x 10^6 cells per well). The following day, cells were transfected again with mixtures containing 2 µg of the respective shRNA plasmids but also containing the reporter and reference/transfection control plasmids (0.2 µg of R-Luc-6xMS2bs and 0.2 µg of F-Luc-GFP) and 0.125–0.25 µg of pcDNA3.1-MS2-HA plasmids (Supplemental Table S1). After 1 d, cells were selected for 48 h in DMEM supplemented with 1.5 µg/mL puromycin and analyzed as before.

**Tethering assays in Dm S2 cells**

For the N-terminally tagged tethering assay [Behm-Ansman et al. 2006] with the luciferase reporter system in Dm S2 cells, cells were seeded in six-well plates (2.5 x 10^6 cells per well) and transfected just thereafter using Effectene (Qiagen). The transfection mixtures contained 0.1 µg of the F-Luc-5BoxB reporter plasmid, 0.4 µg of an R-Luc reference and transfection control plasmid encoding a deadenylation-resistant mRNA lacking BoxB sequences...
For the experiments in Supplemental Figure S3, D–F, Dm S2 cells were additionally cotransfected with 1 μg of a plasmid expressing either wild-type Dm DCP1 (GFP-DCP1 wild-type) or the Dm DCP1 mutant (GFP-DCP1 mut; R70G, L71S, N72S, and T73G) (Kuzuoğlu-Öztürk et al. 2016), and, for the experiments in Figure 7, E–G, cells were cotransfected with 1.5 μg of a plasmid expressing either wild-type Dm CAF40 (GFP-CAF40 wild-type) or the Dm CAF40 mutant (GFP-CAF40 V186E) (Sgromo et al. 2018).

Accession numbers

Atomic coordinates and structure factors for the reported crystal structures have been deposited with the PDB under accession number 6hom for space group P212121 and 6hon for space group I212121.

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Author contributions: E.I. conceived the project. C.K., T.R., A.S., and D.B. performed experiments and analyzed data, together with C.I., D.B., O.W., and E.I. C.K., T.R., A.S., C.I., and D.B. wrote the initial manuscript, which was annotated by E.I. and finalized with help from O.W. All authors commented on and approved the initial manuscript, which was annotated by E.I. and finalized with help from O.W. All authors commented on and approved the final manuscript.

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