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CSNAP is a stoichiometric subunit of the COP9 signalosome

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Summary

The highly conserved COP9 signalosome (CSN) complex is a key regulator of all cullin-RING-ubiquitin ligases (CRLs), the largest family of E3 ubiquitin ligases. Until now it was accepted that the CSN is comprised of eight canonical components. Here, we report the discovery of an additional integral and stoichiometric subunit that thus far evaded detection, and named it CSNAP (CSN Acidic Protein). We show that CSNAP binds CSN3, CSN5 and CSN6 and its incorporation into the CSN complex is mediated through the C-terminal region involving conserved aromatic residues. Moreover, depletion of this small protein influences the cell phenotype, leading to reduced proliferation, and a flattened and enlarged morphology. Finally, based on sequence and structural properties shared by both CSNAP and DSS1, a component of the related 19S lid proteasome complex, we propose that CSNAP, the ninth CSN subunit, is the missing paralogous subunit of DSS1.

Introduction

The COP9 signalosome (CSN) complex is an evolutionarily conserved protein complex that exists in all eukaryotes (for reviews, see: (Schwechheimer, 2004; Wei et al., 2008)). It contains eight canonical subunits that are termed CSN1 through CSN8, according to the descending order of molecular weights. The complex was originally identified as an essential factor that regulates light-induced development in *Arabidopsis thaliana* (At) (Chamovitz et al., 1996; Wei et al., 1994); since then, it has been shown to play a critical role in diverse cellular processes including early development, DNA repair, cytokine signaling, regulation of nuclear transport, cell cycle progression, angiogenesis, and antigen-

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Author Contributions

S.R., G.B.N., M.G.F.L. and M.S. designed the experiments and analyzed the data. S.R., G.B.N., M.G.F.L. and L.M. performed the experiments. A.G., and Y.L. performed the absolute protein quantification. M.E. and S.B.D did the structural and bioinformatics analyses, respectively. M.S. and G.B.N. wrote the manuscript.

induced responses (Schwechheimer, 2004; Wei et al., 2008). The involvement of the CSN in multiple cellular pathways is tied to its biochemical function as a regulator of the ubiquitin proteasome degradation pathway (Adler et al., 2006). Specifically, CSN coordinates the activity of cullin-RING ligases (CRL) (Adler et al., 2006; Deshaies and Joazeiro, 2009).

The CRLs are a family of ubiquitin E3 enzymes that conjugate ubiquitin onto target proteins, thereby exerting a huge impact on cellular regulation (Deshaies and Joazeiro, 2009). Ubiquitination frequently leads to degradation of the target protein; indeed, ~20% of proteasome-dependent degradation is mediated by CRL ubiquitination (Soucy et al., 2009). Yet in some cases, CRL-dependent ubiquitination acts as a switch to activate, repress or relocalize target proteins. The CSN deactivates CRL function in two ways: i) by deconjugation of the ubiquitin-like protein Nedd8 from the cullin subunit (deneddylation), an enzymatic process carried out by CSN5 (Cope et al., 2002); or ii) by physically binding to deneddylated CRLs, precluding interactions with E2 enzymes and ubiquitination substrates (Emberley et al., 2012; Enchev et al., 2012; Fischer et al., 2011). In mammals, the CRL family comprises eight cullin members (Cul1 to Cul7 and PARC), and hundreds of substrate receptor modules that enable specific ubiquitination of multiple proteins involved in diverse cellular processes (Lydeard et al., 2013). Thus, vigorous control of CRLs by the CSN is critical for an organism's normal development and survival. It is therefore not surprising that impairment of CSN function is linked with multiple cancers (reviewed in (Lee et al., 2011; Richardson and Zundel, 2005; Zhang et al., 2013)).

The recently determined crystal structure of the recombinant CSN (Lingaraju et al., 2014) exposed the dynamic and fairly extended conformation of this complex. In particular, its architecture is governed by two organizational centers (Lingaraju et al., 2014); An open horseshoe-shaped structure formed by the 'winged helix' subdomains of the six PCI subunits (proteasome, COP9, and initiation factor 3) (Hofmann and Bucher, 1998), CSN1-CSN4, CSN7, and CSN8, and an elaborate bundle comprising the carboxy-terminal ends of each subunit. Sitting atop this platform is the heterodimer formed by the MPN (Mpr1p and Pad1p N-terminal) (Aravind and Ponting, 1998; Glickman et al., 1998; Hofmann and Bucher, 1998) subunits, CSN5 and CSN6. Binding of neddylated CRLs to CSN triggers substantial remodeling of the complex, activating the isopeptidase activity of CSN5. Given that the CSN is a key regulator of all CRLs, a high degree of flexibility is essential to facilitates its binding to this structurally diverse family of more than 200 distinct complexes, including higher-order structures (Errington et al., 2012; Zhuang et al., 2009).

Notably, the CSN complex shares sequence similarities with two multi-subunit protein complexes: the lid component of the 19S proteasome, and the eukaryotic translation initiation factor 3 (eIF3) (Glickman et al., 1998; Scheel and Hofmann, 2005; Seeger et al., 1998). While the eIF3 complex is more distinct, and contains a larger number of subunits, the lid and the CSN exhibit a remarkable one-to-one correspondence between their two MPN and six PCI subunits. Beside this sequence homology, recent studies revealed that the two complexes also display similar architectures (da Fonseca et al., 2012; Enchev et al., 2012; Lander et al., 2012; Lasker et al., 2012; Lingaraju et al., 2014; Rockel et al., 2014). However, the 19S lid complex contains an additional small, non-PCI or MPN subunit,

known as DSS1 (Sem1 in yeast), that has not yet been identified in the CSN assembly (Sone et al., 2004).

Here, we provide several lines of evidence indicating that a small, intrinsically disordered protein, which we named CSNAP (previously named Myeov2), is an integral subunit of the CSN complex. Our findings support results indicating that CSNAP is pulled down together with CSN subunits (Dunham et al., 2011; Ebina et al., 2013; Sowa et al., 2009). Moreover, we demonstrate that CSNAP, which is present at unit stoichiometry, tethers together the two distinct structural elements of the complex by mutually binding the MPN subunits, CSN5 and CSN6, and the PCI subunit CSN3. Furthermore, the C-terminal end of CSNAP, which is enriched with phenylalanine and aspartic acid residues, is crucial for its integration into the CSN. The lack of CSNAP yields a cellular phenotype characterized by reduced cell proliferation, and a flattened and enlarged morphology. Finally, we suggest that CSNAP is the missing homologous subunit of DSS1, the only 19S lid subunit missing a counterpart in the CSN.

Results

The CSN complex associates with CSNAP

Once the endogenous CSN complex was isolated from human erythrocytes, we examined its composition using a liquid chromatography mass spectrometry (LC-MS)-based approach that we recently developed (Rozen et al., 2013). This method, which couples bottom up and top down MS analysis, enables characterizing the protein complexes' subunit composition. Initially, the constituent subunits are separated on a column under denaturing conditions. Following subunit elution from the column, the flow is split into two fractions. One fraction is directed straight into the mass spectrometer for intact protein mass measurements, while the rest of the flow is fractionated into a 96-well plate, for subsequent peptide sequencing and subunit identification. The heterogeneity of subunit composition is then determined by correlating the subunit mass with its sequence identity.

By applying the LC-MS approach, all eight subunits of the CSN complex could be separated, including the two isoforms of CSN7 (Fig. 1, Table S1). Data analysis demonstrated that all CSN subunits except for CSN1, CSN2 and CSN8 lack the first methionine (Met) residue, and carry an N-terminal acetylation. In addition, the molecular mass of CSN2 suggests that the protein is a product of an alternative translation site at Met9. Similarly, two alternatively translated forms of CSN8 were identified: the mass of the heavier CSN8 variant corresponds to the removal of the first methionine, and the lighter form is a product of an alternative translation initiation site at Met6. The relative intensity of the two forms indicated that only ~30% of the CSN8 integrated within the CSN complex corresponds to the shorter form of the subunit, while the majority of CSN8 corresponds to the full-length protein.

Beyond the inherent diversity of CSN subunits, we noted the repetitive co-elution (retention time: 8 min) of an additional component, along with CSN subunits (Fig. 1, A and B). According on-line MS measurements, we determined that this is a small protein, with a molecular weight of 6.2 kDa. Proteomic analysis identified the protein as Myeloma

overexpressed gene 2 (*Myeov2*), which from now on we refer to as CSNAP, for CSN Acidic Protein. As will become apparent, CSNAP is the ninth subunit of the CSN complex. Though this subunit could have been named CSN9, this already designates the CSN7-like subunit of *Saccharomyces cerevisiae* (Maytal-Kivity et al., 2003).

To determine whether CSNAP specifically interacts with the CSN complex, rather than being a contaminate protein, we purified the complex from a different source: HEK293T cells stably expressing a FLAG-tagged CSN2 subunit. Since entirely different purification strategies were applied to isolate the CSN complex in each system – multi-column biochemical purification for erythrocytes, and a single-step FLAG protocol for HEK293T cells – as well as the great difference between these two cell types, it is unlikely that a similar contaminant protein would co-elute with the CSN in both systems. The subunit elution profile of the CSN complex isolated from HEK293T cells was very similar to that obtained for the CSN isolated from erythrocytes (Fig. 1C).

Apart from the CSN subunits, we identified additional proteins common to both erythrocytes and HEK293 cells that were co-purified with the CSN (Table S2). These include expected members of the ubiquitin-proteasome system, the proteasome inhibitor PI31, SKP1, a member of the SCF (Skp1–Cullin–F-box protein) complex, as well as chaperones and cytoskeleton-related proteins. Included among these proteins was also CSNAP (Fig. 1C), suggesting that it is not a contaminant protein, but rather a specific binding partner of the CSN complex.

CSNAP physically binds the CSN complex

To determine whether CSNAP physically interacts with the CSN complex, we applied the native mass spectrometry (MS) approach under conditions that maintain non-covalent interactions between protein subunits (Sharon, 2013). In this experiment, the intact CSN complex appeared as a charge state series at around ~9,500 m/z (Fig. 1D). However, due to the diversity of CSN subunits (Table S1), and CSNAP's small size, it was impossible to determine unambiguously whether it binds to the CSN. Therefore, tandem MS (MS/MS) experiments were performed, in which a single peak corresponding to CSN ions was isolated. The ions were then subjected to collisional activation, and the individual subunits stripped from the complex were identified (Sharon, 2013). This process not only yielded the dissociation of canonical CSN subunits as CSN8, CSN7b and CSN6, but also to the ejection of CSNAP (Fig. 1E). By extrapolation, we can conclude that prior to the MS/MS analysis, CSNAP was bound to the CSN complex.

The short isoform of CSNAP is a component of the CSN complex

The *Myeov2* gene comprises two alternatively spliced isoforms (Fig. 2A). The first encodes a protein containing 57 amino acids (molecular weight: 6.2 kDa), which we termed CSNAP. The second transcript generates a 252 amino acid-long protein, with a calculated molecular weight of 27.7 kDa, known as *Myeov2-L*. Even though the sequence of the shorter variant, CSNAP, is embedded within *Myeov2-L*, we can unambiguously distinguish between the two proteins via proteomic analysis. This is because *Myeov2-L* contains a short sequential insert between residues 43 to 52, and CSNAP includes a unique C-terminal glutamine residue

(Gln57). Considering that our LC-MS analysis (Fig. 1) involved both molecular weight measurements and proteomic analysis, we could clearly demonstrate that CSNAP is the variant associated with the CSN. Nevertheless, we wished to determine whether Myeov2-L can also interact with the complex.

We began by separating the endogenous CSN complex on a 6% native PAGE (Fig. 2B). Only a single band was detected in the gel, indicating not only the high purity of the sample, but also the stability of the complex. We then extracted the proteins from the band and performed a proteomic LC-MS/MS analysis, using a 30% sequence identity coverage cutoff. Together with the eight canonical CSN subunits, CSNAP, but not Myeov2-L, was also identified (Fig. 2B, right panel), suggesting that unlike CSNAP, Myeov2-L is not a component of the CSN complex.

To further strengthen our results, we generated a polyclonal antibody against a synthetic peptide containing the entire CSNAP sequence, which recognized both the CSNAP and Myeov2-L isoforms. We then performed a gel filtration analysis of HEK293 cell extract, to monitor whether these proteins co-elute with the CSN complex. Western blot analysis indicated that CSNAP displays an elution profile similar to those of CSN1, CSN3, and CSN8, while Myeov2-L elutes at lower molecular weight fractions (Fig. 2C), supporting our premise that Myeov2-L is not part of the CSN complex.

Next, reciprocal co-immunoprecipitation experiments were performed, using both HeLa and HEK293 cell extracts (Fig. 2D). The results obtained for the two cell lines were virtually the same: When cells transiently expressing CSNAP-myc-FLAG were co-immunoprecipitated using an anti-FLAG resin, CSN1, CSN3, CSN5, and CSN8 were pulled down. When anti-CSN3 was used for pulldown in HeLa cells, the antibody directed toward CSNAP gave rise to two bands, an intense band at CSNAP's molecular weight, and a second, faint band corresponding in size to Myeov2-L. Notably, the intensity of the Myeov2-L band was similar to the non-specific signal that appeared when non-conjugated beads were used as a negative control, suggesting that this faint band is a product of a non-specific interaction. When we used HEK293T cells stably expressing FLAG-CSN2 and an anti-FLAG resin, CSNAP was pulled down, but not Myeov2-L. Similar results were also obtained when reciprocal co-immunoprecipitation experiments were performed using A31N-ts20 BALB/c mouse embryo fibroblast cells (Fig. S1). Moreover, pulldown assays using anti-FLAG resin or an anti-CSN3 antibody depleted both CSNAP and FLAG-CSN2 from the unbound fraction, unlike Myeov2-L or GAPDH, respectively (Fig. 2E). This result suggests that cellular CSNAP is present in the majority of CSN complexes. Altogether, the data imply that only CSNAP, and not Myeov2-L, is a bona fide member of the CSN complex.

CSNAP is a stoichiometric subunit of the CSN complex

To determine whether CSNAP is a stoichiometric component of the CSN, we subjected the anti-CSN3 immunoprecipitated complex to targeted proteomic analysis by selective reaction monitoring (SRM) MS for absolute quantification (based on the AQUA approach) (Menetret et al., 2007). Custom stable isotope-labeled peptides of CSN2, CSN4 and CSNAP were synthesized and quantified by amino acid analysis. Those were spiked into the complex samples, enabling absolute quantification of the CSN and CSNAP subunits by converting

light/heavy peak area ratios into absolute protein amounts. Values were then normalized against a representative peptide (Fig. 2F). We found that the measured stoichiometry of CSN2, CSN4 and CSNAP was ~1:1:1, indicating that CSNAP, like the other CSN subunits, is present in equimolar amounts.

The C-terminal F/D-rich region of CSNAP mediates its association with CSN, with Phe44 and Phe51 playing direct roles in the interaction

To clarify which domain within CSNAP mediates its interaction with the CSN complex, we transiently expressed several versions of CSNAP, including deletion mutants of either the N- or C-terminal regions, fused to the fluorescent protein Cerulean (Cer) in HEK293 cells (Fig. 3A). Initially we examined the cellular localization of CSNAP. Full-length CSNAP constructs were seen throughout the cytoplasm and nucleus, but were excluded from nucleoli (Fig. S2), as previously observed for other CSN subunits (Fuzesi-Levi et al., 2014). We then performed reciprocal co-immunoprecipitation analyses, using antibodies against CSN3 and GFP (Fig. 3B). The results indicated that fusing Cerulean to either terminus of CSNAP did not hinder its ability to interact with CSN. Likewise, deletion of the N-terminal region of CSNAP did not affect the CSNAP/CSN association. However, CSNAP was unable to bind CSN in the absence of the C-terminal fragment, suggesting that this region is essential for interaction with the complex.

Analysis of the amino acid sequence of CSNAP indicates that it is enriched with charged and polar amino acids, characteristic of intrinsically disordered proteins (Dyson and Wright, 2005). Such proteins, however, tend to adopt distinct structures in their bound state. When we examined CSNAP's C-terminal sequence, the region critical for its incorporation into the CSN, we noticed that it is not only rich in phenylalanine and aspartic residues (F/D rich domain), but that the two types of residues are alternately spaced three and four residues apart. This observation suggests that this C-terminal region might adopt a binding configuration of an amphipathic α -helix, containing both aromatic and negatively charged sides (Fig. 3C). It is therefore reasonable to assume that the aromatic face is buried within the complex forming the CSNAP/CSN interface.

To test this hypothesis, single and double mutants were introduced into N-CSNAP-Cer at positions 44 and 51 (F44A, F51A, F44A-F51A). The mutant proteins were transiently expressed in HEK293 cells, and subjected to reciprocal co-immunoprecipitation with antibodies against CSN3 and GFP (Fig. 3D). Though the F44A mutation did not interfere with the CSNAP/CSN interaction, and binding of the F51A CSNAP mutational variant to CSN was noticeably reduced, it was the F44A-F51A double mutant that displayed the most striking phenotype, completely abolishing the ability of CSNAP to interact with the CSN. Therefore, within the F/D rich C-terminal region of CSNAP, F44 and F51 are necessary mediators of its interaction with the CSN.

Cellular analyses indicate that CSNAP is an integral subunit of the CSN complex

We recently demonstrated that following induction of UV damage, the CSN complex is transiently recruited to the nucleoplasmic and chromatin fractions (Fuzesi-Levi et al., 2014). We therefore wished to explore whether CSNAP acts as an integral CSN component, and

displays a similar relocalization pattern. To this end, HeLa cells were exposed to UV irradiation, and cells were then fractionated (Fig. 4A). Our findings indicated that the level of cytosolic CSNAP did not significantly change in response to UV irradiation, though a clear increase in CSNAP intensity was observed in the nucleoplasmic and chromatin-associated fractions immediately after UV irradiation, as observed for CSN subunits. However, no change in the intensity levels of Myeov2-L was detected. This finding confirms the analogous cellular response of CSNAP and the CSN complex.

To study the diffusion kinetics of CSNAP in live cells, we performed fluorescence recovery after photobleaching (FRAP) measurements, in both the cytosol and nuclear compartments of HEK293 cells stably expressing different fluorescently tagged CSNAP constructs (Luijsterburg et al., 2007). Previously, we had demonstrated that the recovery curves of CSN2, CSN3, CSN6 and CSN7 subunits are very similar, indicating that they commonly reside within the holoCSN complex (Fuzesi-Levi et al., 2014). Thus, if CSNAP is an integral subunit of the complex, it is expected to display comparable mobility. For comparison, we conducted similar measurements on cells stably expressing free Cerulean and Cer-CSN3, representing the dynamics of a fully mobile, monomeric protein (Dross et al., 2009) and the CSN complex, respectively. As an additional control, we used cells stably expressing fluorescently tagged DDB2, a nuclear DNA-binding protein previously shown to interact with the CSN, though not an integral subunit of the complex (Groisman et al., 2003; Olma et al., 2009).

Examination of the resulting data indicated that Cer-CSNAP and Cer-CSN3 recovery curves were remarkably similar, displaying significantly slower mobility compared to that of free Cerulean, and suggesting that Cer-CSNAP is part of the CSN complex (Figs. 4B and S3). In contrast, the mobility kinetics of the CSNAP construct lacking the C-terminal F/D region was comparable to that of free Cerulean, further demonstrating that this region is critical to CSNAP incorporation into the CSN. The absence of the N-terminal region did not affect the recovery rate and which was similar to that of Cer-CSNAP and Cer-CSN3, suggesting that the F/D region alone is sufficient for CSNAP/CSN assembly. In the nucleus, DDB2 exhibited a clearly slower recovery compared to that of Cer-CSN3 and Cer-CSNAP, reflecting some degree of transient immobilization. Taken together, the similar mobility kinetics of CSNAP and CSN3 strongly suggests that CSNAP is an integral component of the holoCSN complex.

CSNAP interacts with CSN3, CSN5, and CSN6

To determine which of the CSN subunits binds CSNAP, we performed chemical cross-linking reactions using BS³, a homobifunctional amine-reactive compound that reacts predominantly with the primary amines in lysine side-chains, and the N termini of polypeptide chains (Kalkhof and Sinz, 2008). We then monitored the change in the protein's band pattern by running denaturing gel electrophoresis (SDS-PAGE) before and after cross-linking. Immunoblotting with anti-CSNAP and various anti-CSN antibodies revealed the appearance of similarly sized, higher molecular mass species for CSNAP and CSN3, CSN5 and CSN6 (Fig. 5A). The CSNAP-CSN3, CSNAP-CSN5, and CSNAP-CSN6 bands were sensitive to the presence of a denaturing agent: when SDS was added and the sample was

boiled prior to the cross-linking reaction, the cross-linked species were eliminated, implying the specificity of the interaction (Fig. 5A). In contrast, cross-links with CSNAP were not identified for CSN1, CSN2 and CSN8.

Despite numerous attempts, we were unable to map the specific sites of association between CSNAP and CSN3, CSN5 and CSN6 using proteomic MS analysis. This is likely due to CSNAP's relatively small size and the few peptides it produces, reducing the odds of identifying explicit CSNAP cross-linked species within the complex mixture of CSN peptides. Nonetheless, when we examined the recently solved crystal structure of the recombinant CSN complex (Lingaraju et al., 2014), the only region in which CSN3, CSN5 and CSN6 are in proximity, and their surfaces not obscured by the other subunits, is near the C-terminal helices of CSN3 and CSN6, and the loop 284-295 of CSN5, which connects its two C-terminal helices (Fig 5B). Near this region, we noted a highly positive patch on the CSN3 surface that contains several exposed lysine residues capable of forming a cross-link with CSNAP. Lysine residues of CSN5 and CSN6, which could act as additional cross-link targets, are located near the CSN3 positive patch (Fig. 5C). This patch, which corresponds to the PCI domain of CSN3, includes several hydrophobic pockets, which are preferred phenylalanine binding sites as determined by ANCHORSmap (Ben-Shimon and Eisenstein, 2010). From the crosslinking results, and the structural features of CSNAP and the CSN subunits, we suggest that the positive patch of CSN3 binds the CSNAP F/D-rich motif by forming ionic interactions with the negative charges, and hydrophobic interactions with CSNAP Phe44, Phe51, and perhaps also Phe47. Other parts of CSNAP bind near the helical bundle region, thus tethering the MPN dimer of CSN5 and CSN6 to the PCI proteins, through CSN3.

CSNAP and DSS1 share sequential and structural properties

At this point we referred to the related 19S lid complex, which shares one-to-one subunit correspondence with the CSN, except for DSS1 that is missing a counterpart. We noticed, however, that CSNAP and DSS1 not only share sequential similarity (Fig. S4A), but they also belong to the intrinsically unstructured protein family (Fig. S4B). Thus, CSNAP could potentially be the subunit homologous to DSS1 and as such, may occupy a similar position within the CSN architecture.

The molecular architecture of the 26S holocomplex was recently determined using cryo-EM single-particle analyses; however, the position of DSS1/Sem1 within the lid was not resolved, due to its small size (Beck et al., 2012; Lander et al., 2012; Lasker et al., 2012). Nevertheless, cross-linking analysis and biochemical characterization indicated that DSS1/Sem1 forms a subcomplex with RPN3 and RPN7, enforcing their joint incorporation into the lid (Fukunaga et al., 2010; Sharon et al., 2006; Tomko and Hochstrasser, 2014). Comparative cryo-electron microscopy maps and cross-linking restraints position DSS1/Sem1 in the cleft between the PCI domains of Rpn7 and Rpn3, while its central part lies near the N-terminal region of Rpn3. This mode of binding, near the 19S lid helical bundle, fastens the lid together (Bohn et al., 2013; Kao et al., 2012; Tomko and Hochstrasser, 2014). Based on the cross-linking results it seems that CSNAP occupies an analogous binding mode, in which it tethers the MPN subunits CSN5 and CSN6, to the PCI CSN3 subunit at the helical bundle

region. More evidence of this binding mode is provided by superpositioning the PCI subunit of PCID2 in the PCID2/DSS1 complex (Ellisdon et al., 2012) onto the PCI domain of CSN3, showing that the DSS1 binding surface of PCID2 corresponds to the CSN3 positive patch, proposed here to be the CSNAP binding surface (Fig. 5D). Small differences in the actual binding modes support the observation that CSNAP and DSS1/Sem1 are intrinsically unstructured proteins; therefore, their interaction partners co-determine their structure, as previously shown for DSS1 (Ellisdon et al., 2012; Yang et al., 2002). Overall, it suggests that like DSS1/Sem1, CSNAP connects subunits within the complex.

CSNAP-depleted cells display a distinct phenotype

To test whether CSNAP confers functional significance, HAP-1 cell lines lacking CSNAP (CSNAP cells) were generated using the CRISPR system. Initially, pull-down assays using an anti-CSN3 antibody confirmed that CSNAP is absent from the CSN complex in the CSNAP cell line (Fig. 6A). Next, cell extracts of wild type and CSNAP cells were passed through a column containing CSNAP-conjugated beads. After washing, beads were boiled in SDS-PAGE sample buffer, and the bound proteins analyzed by Western blots. We found that only CSN from the CSNAP cells was able to bind the CSNAP-column, whereas that from the wild-type cells did not bind (Fig. 6B). This result not only validated the lack of CSNAP in the engineered cells, it also confirmed the stoichiometric CSN/CSNAP association in wild type cells.

We then examined whether the enzymatic activity of the CSNAP and wild-type CSN complexes differ. No significant difference in the deneddylation capacity was observed between the wild-type or CSNAP cells (Fig. 6C). This observation is in accordance with a previous study which compared the rate of deneddylation of endogenous CSN prepared from HEK293 cells, with that of recombinant CSN expressed in insect cells (Emberley et al., 2012). Although the recombinant CSN might contain the insect CSNAP protein, it is unlikely to be stoichiometric, due to its very high overexpression levels. Thus, CSNAP might not have a significant effect on the enzymatic activity of the CSN.

Next, we examined whether the lack of CSNAP gives rise to a detectable cellular phenotype. To this end, we performed cell proliferation assays, and compared the growth potential of wild-type and CSNAP HAP-1 cells. Our data (Fig. 6D) show that there is a clear reduction in cell proliferation in the CSNAP clone, compared to the wild-type cells. Furthermore, microscopy analysis of both cell types demonstrated that CSNAP cells display a flattened and enlarged cell phenotype, compared to wild-type cells (Fig. 6E-G). These results indicate that the lack of CSNAP generates a distinct phenotype, indicating the functional significance of this small subunit. Nevertheless, further research is required in order to determine the specific mechanistic function of CSNAP.

Discussion

Here, we show that CSNAP, a 6.2 kDa protein, is a stoichiometric subunit of the CSN complex. The protein's second, longer isoform, Myeov2-L (27.7 kDa), which contains the entire sequence of CSNAP except for the C-terminal amino acid, is not, however, part of the complex. Moreover, we demonstrate that the C-terminal F/D-rich region of CSNAP is

necessary and sufficient for its incorporation into the CSN and, in particular, conserved Phe44 and Phe51 take part in this interaction. Our data also demonstrate that CSNAP binds to CSN3, CSN5 and CSN6, possibly linking the MPN and PCI substructures. Finally, we found that the absence of CSNAP had a marked effect on the cellular morphology and proliferation rate. Although this result implies biological relevance, further investigation is required to determine the specific functional role of CSNAP. It is tempting to suggest that it contributes to the complex stability or flexibility, enabling the CSN to associate and dissociate from the variable CRL complexes.

CSNAP was first identified as an overexpressed gene in ARH-77 cells (Tang et al., 2003), which were considered to be cultures of human myeloma cells, giving the protein its original name, Myeov2 (myeloma-overexpressed gene 2). Notably, however, these cells were shown to be a false malignant hematopoietic cell line, essentially representing EBV-transformed B lymphoblastoid cells (Drexler et al., 1999), a fact that supports our suggestion to rename the protein. Still, this study stimulated us to examine whether overexpression of CSNAP in ARH-77 cells is coordinated with the other CSN subunits, as expected of constituents of the same protein complex (Eisen et al., 1998; Jansen et al., 2002). Interestingly, analysis of microarray data using the GeneVestigator database not only revealed that the expression levels of the entire CSN complex, including CSNAP, are highly comparable in ARH-77 and various other cell lines (Fig. S5A), but also that similar expression coherence is obtained across different tissues and organs (Fig. S5B). Likewise, when we examined the expression patterns in a range of cancers (e.g., colon, leukemia and lung) reported to induce overexpression of CSN5 and/or CSN6 (Adler et al., 2008; Chen et al., 2014), the entire CSN complex, including CSNAP, displayed similar expression levels (Fig. S5C). Together, these results indicate that all CSN-encoding genes, including CSNAP, show similar expression patterns, as expected from subunits of the same protein complex (Jansen et al., 2002).

The first documentation concerning the existence of CSNAP at the protein level came from a global proteomic analysis aimed at identifying deubiquitinating enzyme-associated proteins (Sowa et al., 2009). The study indicated that CSNAP is pulled down, together with CSN6. Similarly, a methodological study focused on the effectiveness of multidimensional separation techniques for MS analysis indicated that Myeov2 is repeatedly identified in FLAG-CSN5 purifications (Dunham et al., 2011). A later study, focusing on the subnuclear localization of the ribosomal L11 protein, indicated that overexpression of a FLAG-tagged construct containing the N-terminal region of Myeov2-L, comprising the F/D-rich motif, is co-immunoprecipitated with CSN5, Cul1 and Cul3 (Ebina et al., 2013). In contrast, a construct containing the long, Myeov2-L C-terminal tail could not interact with the CSN complex. Moreover, a genetic screen searching for *Arabidopsis thaliana* mutants that display resistance to the auxin inhibitor p-chlorophenoxyisobutyric acid identified the Small Acidic Protein 1 (SMAP1) (Rahman et al., 2006), the plant ortholog of CSNAP. A subsequent study indicated that SMAP1 physically binds the CSN complex via the F/D-rich region (Nakasone et al., 2012). Taken together, these results support our view that CSNAP is an integral component of the CSN complex.

The CSN complex is conserved throughout eukaryote evolution, from fungi to humans (Schwechheimer, 2004; Wei et al., 2008). Therefore, it would be expected that as a CSN

component, CSNAP would be similarly conserved. To examine this possibility, we performed bioinformatics searches, the results of which indicated that CSNAP is highly conserved in all chordates, and also displays a high degree of similarity in lower eukaryotes (Fig. S6). In plants, conservation is mainly maintained in the C-terminal F/D-rich region, the domain essential for CSN integration. We could not, however, identify a yeast ortholog of CSNAP, perhaps because the protein diverged to such an extent that it is difficult to identify it by homology database searches; rather, experimental approaches are required. More conservational support for the CSN/CSNAP link comes from the fact that CSN7b, CSN8 and CSNAP are all located in the same chromosomal region (2q37), and that they are found together in syntenic regions in all mammalian species (e.g., mouse, rat, horse and pig) whose DNA sequences are available so far. In marsupials, there is no CSN7b gene; however, CSN8 and CSNAP are co-localized on the same chromosome. The shared synteny in mammals may reflect selection due to a functional relationship (Moreno-Hagelsieb et al., 2001), strengthening the link between CSNAP and the CSN.

Unlike CSNAP, there is very little transcriptional evidence that Myeov2-L is expressed in humans; thus far, no such evidence has been found in other species at all (Fig. S7). The lack of conservation in all non-primates makes it highly unlikely that it exists in those species. Even in primates, where the exon sequences are mostly present, the splice junctions, particularly in the fourth exon, are defective in all species except chimp and human. Therefore, this isoform is most likely human- or human/chimp-specific, and unlikely to play a role in a complex as highly conserved as the CSN.

A striking one-to-one correspondence exists between the CSN subunits, and those of the 19S lid subcomplex of the 26S proteasome, suggesting that they diversified from a common ancestor (Glickman et al., 1998; Seeger et al., 1998). The two signature structural motifs entail six subunits with a PCI and two with an MPN domain, all found within each of the complexes. The lid subunit DSS1 (Sem1 in budding yeast) is the only exception: DSS1 does not contain a PCI or MPN fold; moreover, it is missing a counterpart within the CSN. Considering that i) like CSNAP, DSS1 links 19S proteasome lid subunits (Paraskevopoulos et al., 2014; Sone et al., 2004; Tomko and Hochstrasser, 2014), though an intact complex can be formed in its absence (Paraskevopoulos et al., 2014); ii) CSNAP and DSS1 are remarkably small proteins that share 18% sequential identity and 26% similarity, including a C-terminal region enriched with aromatic and acidic residues (Fig. S4A); iii) both proteins belong to the intrinsically unstructured protein family (Fig. S4B), we suggest that DSS1 and CSNAP are paralogous subunits, filling the missing gap of homology between the 19S lid and CSN complexes (Fig. 7).

Two decades have passed since the discovery of the CSN complex (Chamovitz et al., 1996; Wei et al., 1994); thus, a key question arises: How has the CSNAP/CSN association escaped previous detection? The answer probably lies within the sequence of CSNAP. First, the low molecular weight of the protein is likely to cause its rapid migration outside polyacrylamide electrophoresis gels, preventing its detection. Second, even if CSNAP is retained within the gel, the fact that it contains only two basic amino acids – i.e., two lysine residues – gives rise to poor Coomassie dye staining (Syrový and Hodny, 1991). Finally, proteomic identification of CSNAP might also be challenging, considering that tryptic peptides are typically

analyzed (Mann et al., 2001). This procedure not only generates just two CSNAP peptides, but their long length, which is in the upper mass scale for typical analysis, also challenges identification. Hence, the methodology we described herein, which couples bottom up and top down MS analysis, set the stage for the discovery of CSNAP, and paves the way toward identification of additional small proteins that act as integral subunits of large protein complexes.

Experimental Procedures

Purification of the CSN complex

CSN was isolated from human erythrocytes, according to a previously published protocol (Hetfeld et al., 2005). FLAG purification of CSN is described in the Supplemental Experimental Procedures.

Generation of a polyclonal antibody against CSNAP

The full-length CSNAP peptide was conjugated to KLH (Sigma Aldrich) using the BS³ crosslinker (Pierce, Thermo Scientific, USA), and used for immunization in rabbits. After 3 boosts, serum was obtained, and anti-CSNAP antibodies were affinity-purified on CSNAP peptide coupled to Aminolink beads (Pierce, Thermo Scientific, USA) and stored in 50% glycerol at -20°C. Specificity was validated in comparison to pre-immune serum, against synthetic CSNAP and cell lysates.

Subcellular fractionation

HeLa cells were fractionated to cytoplasmic, nuclear soluble, and chromatin-bound fractions, with or without exposure to 20 J/cm² UV-C, as described (Fuzesi-Levi et al., 2014). UV-C exposed samples were collected ten minutes after induction of UV damage.

Activity assay

Deneddylation activity of HAP1 cell lines was measured as in (Fuzesi-Levi et al., 2014). At each time point, deneddylation activity was calculated as the measured intensity of the deneddylated band divided by the total intensities of the neddylated and deneddylated bands, and normalized to maximal activity (10 minutes). Average activity levels and standard errors were calculated for each time point from three independent experiments.

LC-MS approach

The monolithic-LC-MS approach was performed as previously described (Rozen et al., 2013).

Native MS analysis

Nanoelectrospray ionization mass spectrometry (MS) and tandem MS (MS/MS) experiments were performed on a Synapt G2 instrument (Waters, USA), as described in (Kirshenbaum et al., 2010).

Confocal fluorescence microscopy

Confocal fluorescence imaging and FRAP (fluorescence recovery after photobleaching) measurements were performed as previously described (Fuzesi-Levi et al., 2014).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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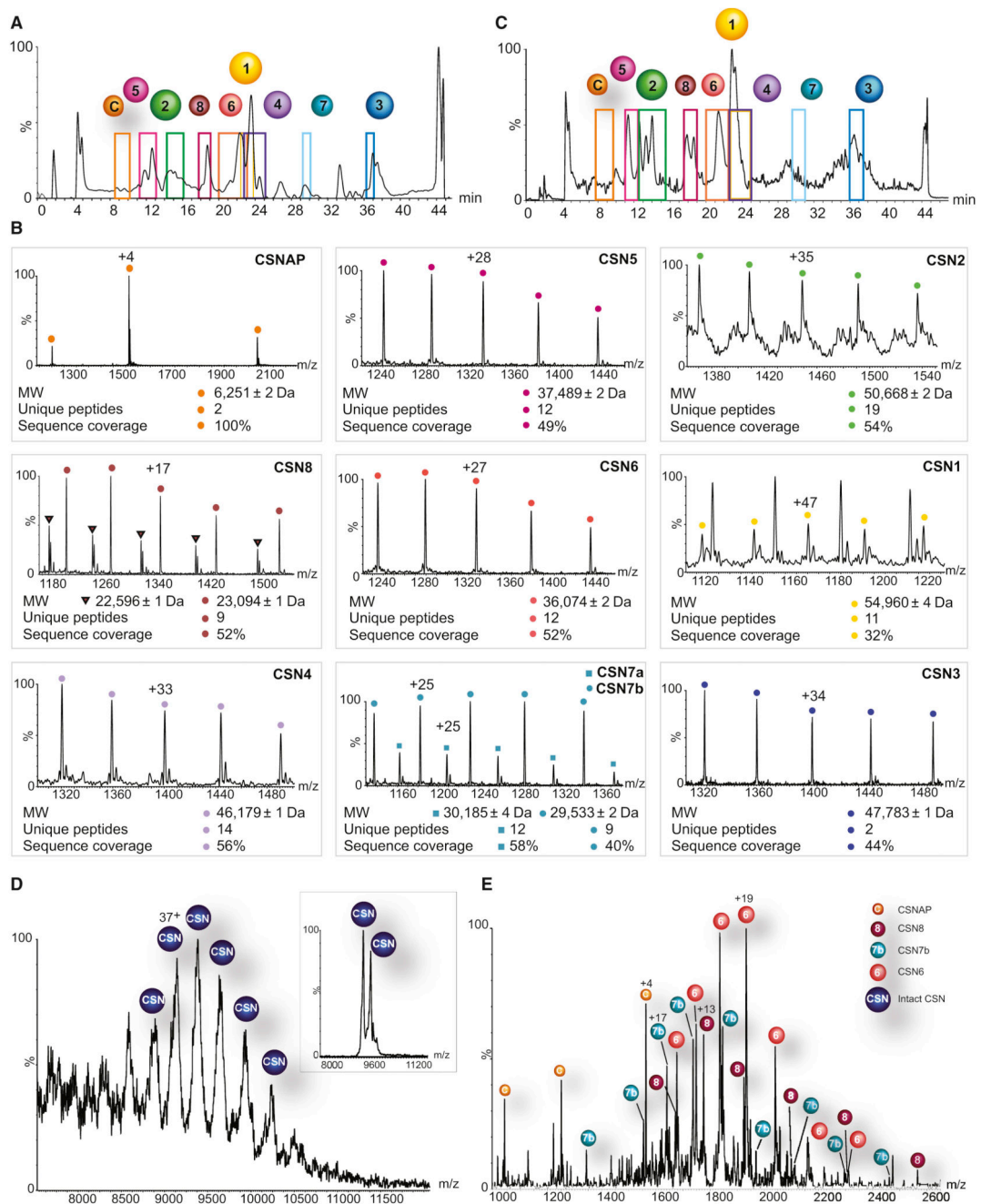


Figure 1. CSNAP physically associates with the CSN complex.

The endogenous CSN complex isolated from human erythrocytes (A) and HEK293 cells (C) was separated into its component subunits, using a monolithic column under denaturing conditions. A colored frame highlights the retention time of each eluted protein. CSNAP (labeled as C) was persistently detected alongside the eluted CSN subunits, implying its association with the CSN complex. (B) The resulting ESI-QToF mass spectra are shown. These spectra made it possible to determine the mass of the eluted protein and its associated variants; proteomics analysis was performed for sequence identification. CSNAP repeatedly

co-eluted with the CSN subunits, suggesting that it associates with the complex. Indicated masses are an average of biological and technical measurements. Subunit variants are differentiated by labeling with circles and squares. (D) Nano-electrospray mass spectrum recorded under native conditions of the human CSN complex isolated from HEK293T cells. The intact CSN complex is observed between 8,500 and 10,500 m/z. The 36⁺ and 35⁺ charge states (inset) were selected for tandem MS analysis. (E) MS/MS spectrum showing the individual subunits stripped from the CSN complex. In addition to the dissociation of CSN6, CSN7b, and CSN8, we could also assign peaks corresponding in mass to CSNAP, indicating that it interacts with the CSN complex. The different species are denoted with labeled circles.

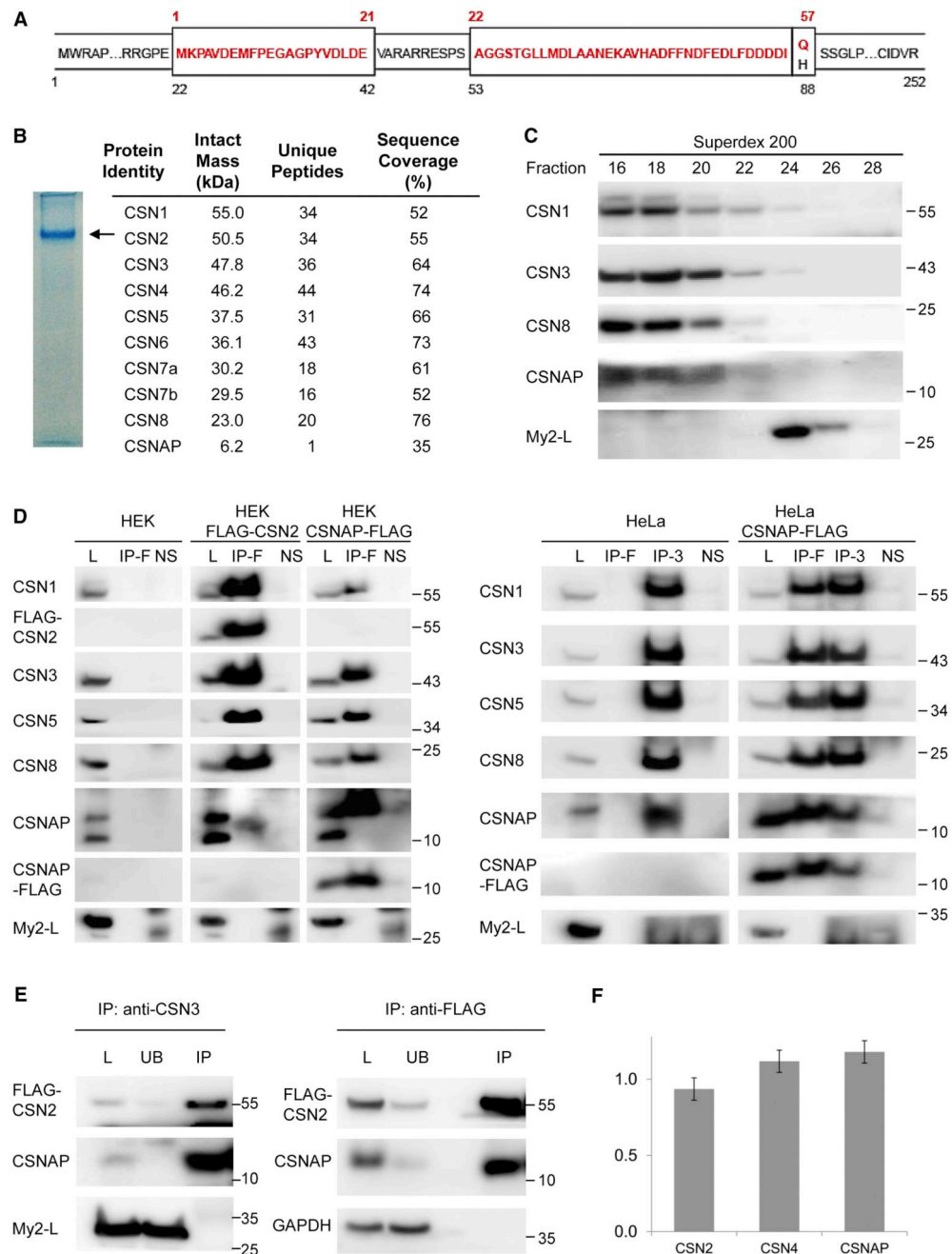


Figure 2. The short isoform of the Myeov2 gene, CSNAP, interacts with the CSN complex. (A) Schematic representation of the two alternatively spliced products of the Myeov2 gene. Amino acid residues colored in red represent the CSNAP sequence (57 amino acids). The long Myeov2 protein (252 amino acids, My2-L) contains the entire sequence of the short transcript except for glutamine 57, as well as additional sequence stretches in internal and end regions (shown in black). (B) Native PAGE separation (6%) of the purified CSN complex. The position of the complex is denoted by an arrow; the absence of additional bands in the gel indicates the high integrity of the complex. Proteins extracted from the

labeled band were subjected to proteomic LC-MS/MS analysis. Identified proteins with sequence coverage $\geq 30\%$ are listed. (C) HEK293T FLAG-CSN2 cell extract was subjected to Superdex 200 gel-filtration chromatography. Fractions were collected and analyzed by SDS-PAGE and immunoblotting. CSNAP co-fractionated with the CSN complex, unlike Myeov2-L, that eluted in lower molecular weight fractions. (D) Whole-cell lysates (L) from HEK293T cells stably expressing FLAG-CSN2, or HEK293 cells transiently expressing CSNAP-FLAG (left panel), as well as lysates from HeLa cells transiently expressing CSNAP-FLAG (right panel), were subjected to immunoprecipitation (IP) using FLAG affinity gel (IP-F) or anti-CSN3 antibody (IP-3). Pulldowns were analyzed by Western blots, using antibodies against CSN subunits, CSNAP, and FLAG. The results show that CSNAP, but not the long version of Myeov2, co-immunoprecipitate with the CSN complex. As controls, WT HEK293 and HeLa lysates were used. To rule out non-specific interactions, lysates were also incubated with Protein G Sepharose, without the addition of the primary antibodies (NS). (E) CSN was co-immunoprecipitated from HEK293 stably expressing FLAG-CSN2, using an anti-CSN3 antibody or anti-FLAG resin. The whole-cell lysates (L) bound (IP) and unbound (UB) fractions were analyzed by Western blots using antibodies against CSNAP and FLAG. GAPDH and My2-L were used as negative controls. The depletion of CSN2 and CSNAP from the unbound fraction not only suggests that the majority of CSN complexes are bound to CSNAP, but also that this small protein is not part of another protein complex. In all Western blots, molecular weights are indicated in kDa units. All experiments were repeated at least 3 times. (F) CSN was subjected to targeted proteomic analysis by selective reaction monitoring (SRM) mass spectrometry and stable, isotopically labeled peptide standards. Absolute quantification was done by referencing the native peptide intensities to the heavy labeled standards, and then normalizing against a representative peptide. Results indicate that CSN subunits and CSNAP are present in equimolar amounts. Data shown are the result of three biological replicates, with two technical replicates each. Error bars indicate the standard deviations of all six measurements.

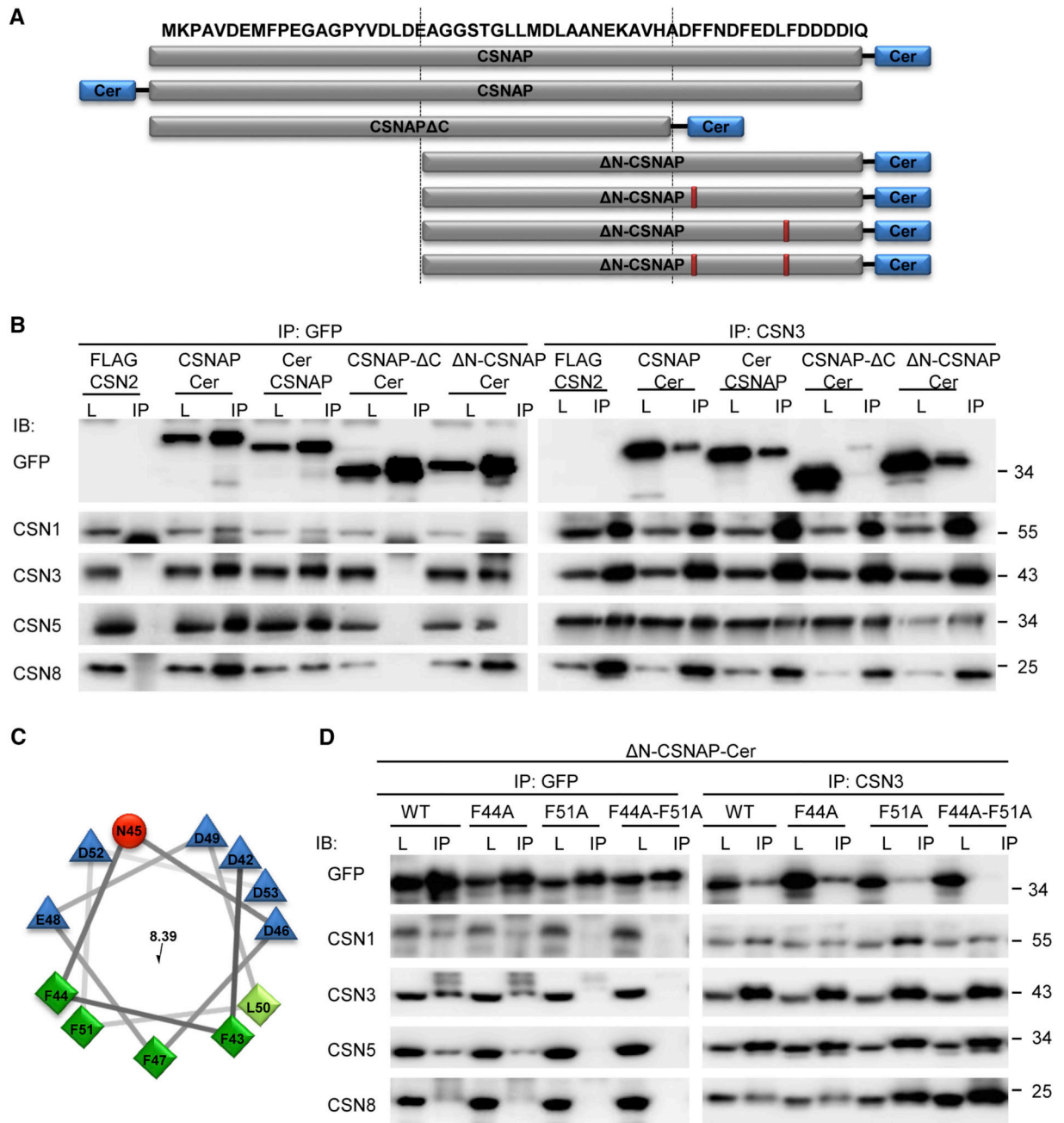


Figure 3. The F/D-rich C-terminal domain of CSNAP, specifically Phe44 and Phe51, are involved in its interaction with the CSN complex.

(A) Schematic representation of the different CSNAP constructs used in this experiment. (B) Cellular proteins extracted from the different fluorescently tagged HEK293 cell lines were immunoprecipitated, using anti-GFP and anti-CSN3 antibodies. As control, lysate from HEK293 cells stably expressing FLAG-CSN2 was used. Lysates (L) were run side-by-side with their corresponding immunoprecipitated proteins (IP), and visualized using various antibodies, as indicated (IB). Results show that CSNAP- C-Cer did not interact with the CSN, indicating that the C-terminal domain is responsible for its interaction with the

complex. (C) Helical wheel representation of the CSNAP C-terminal region. Hydrophilic and hydrophobic residues are colored blue and green, respectively. Distribution of the residues on either side of the helix suggests amphipathic properties for this structure. (D) Lysates (L) from HEK293 cells expressing N-CSNAP-Cer (WT) and its mutational variants, consisting of single (F44A and F51A) and double (F44A-F51A) amino acid substitutions, were immunoprecipitated by either anti-GFP or anti-CSN3 (IP). Pulldowns were analyzed by Western blot, using antibodies against GFP and CSN subunits. Findings show that while Phe44 displacement yielded results similar to those of the WT construct, the F51A mutant extensively weakened the interaction of N-CSNAP with the CSN. The most pronounced effect was observed for the F44A-F51A double mutant, which entirely abolished the N-CSNAP/CSN interaction.

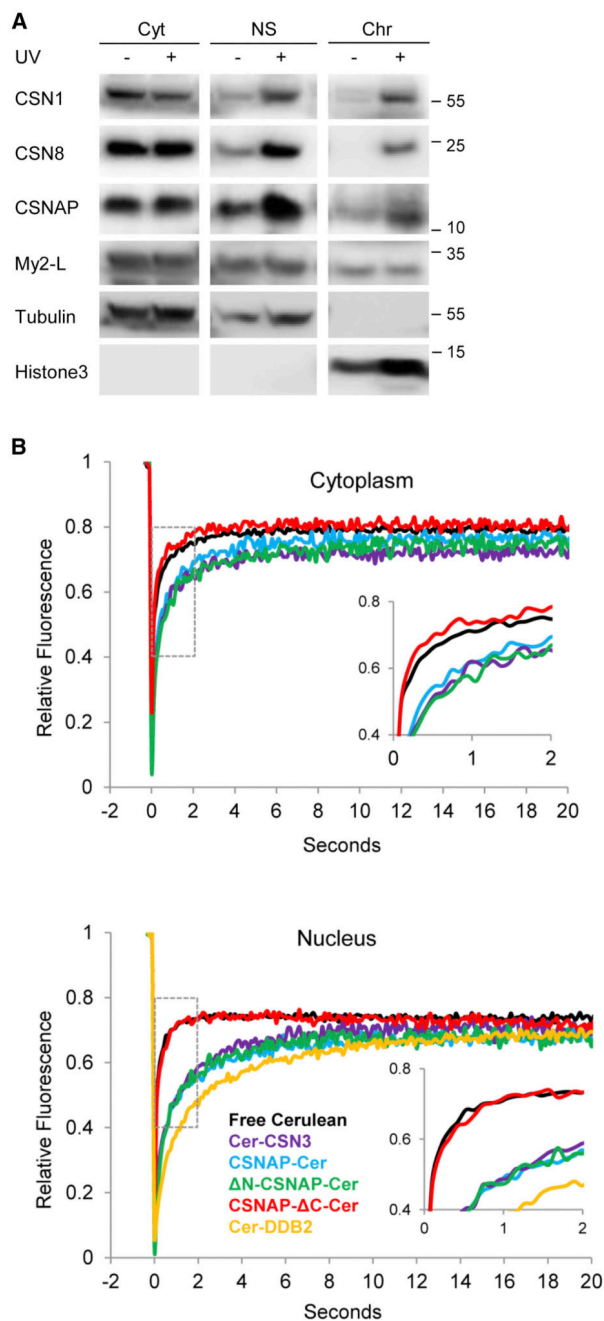


Figure 4. Live cellular analyses indicate that CSNAP is an integral CSN subunit.

(A) HeLa cells were fractionated into cytoplasmic, nuclear soluble, and chromatin-associated fractions, with or without prior exposure to UV light. Fractions were separated on tricine-SDS gels, and blots were probed with anti-CSN1, CSN8 and CSNAP antibodies. As controls, anti-tubulin and anti-histone 3 antibodies were used. Like CSN subunits, CSNAP, but not Myeov2-L (My2-L), is recruited to the nucleoplasmic and chromatin fractions following DNA damage induction. (B) FRAP curves of full-length CSNAP-Cer, as well as its deletion mutants, N-CSNAP-Cer and CSNAP- C-Cer, were compared to those obtained

for free Cerulean and fluorescently labeled CSN3 and DDB2. Each plot constitutes an average of at least 40 cells, normalized to pre-bleach intensity. To better display the mobility differences between the measured cell lines, the regions within the dashed squares were enlarged (insets).

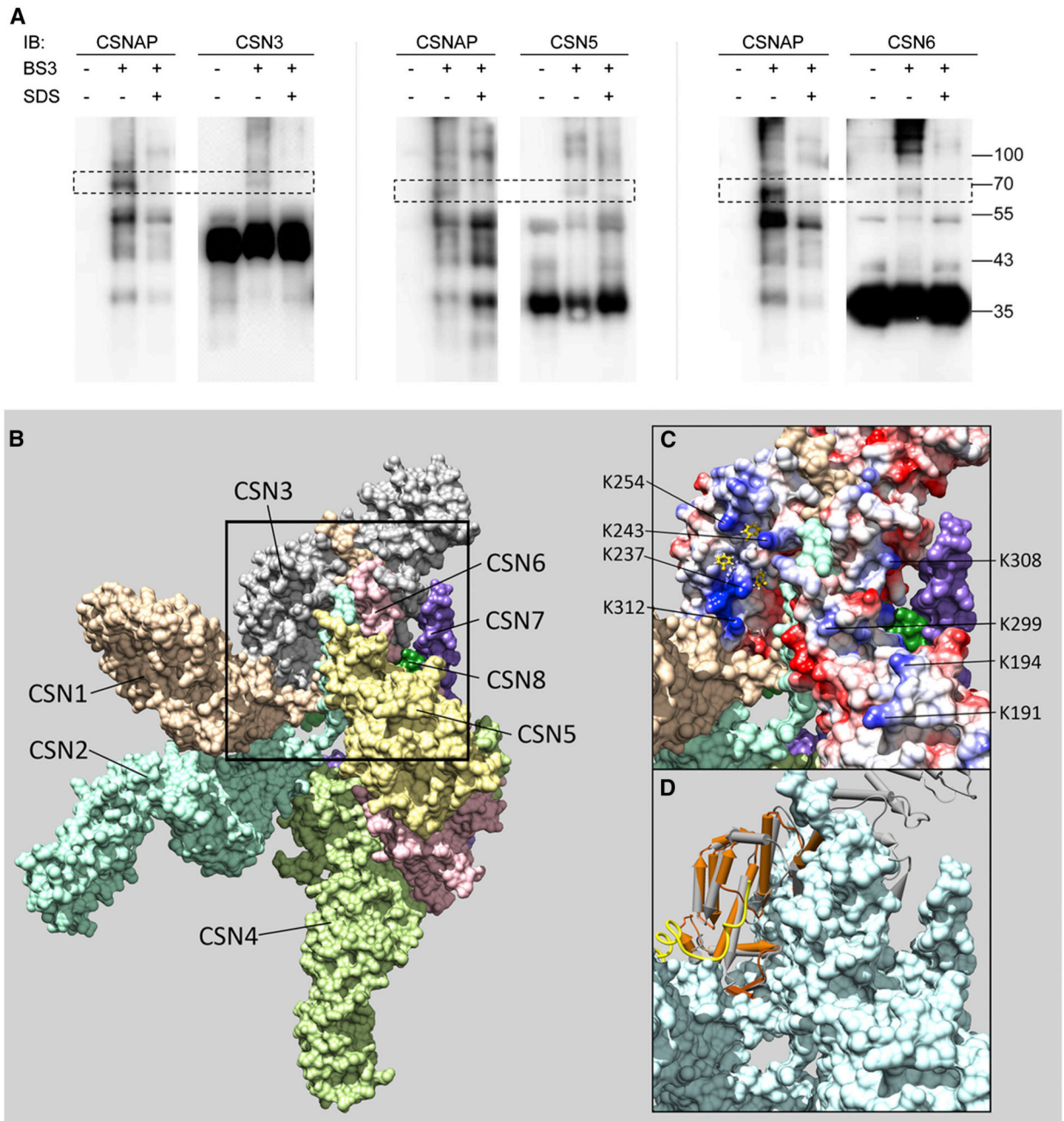


Figure 5. CSNAP interacts with CSN3, CSN5 and CSN6.

(A) FLAG-tagged CSNAP was purified from HEK293T cells and cross-linked with BS³. As a control for specific CSNAP association, FLAG-CSNAP was denatured with 1% SDS and boiled for 5 minutes, prior to addition of the cross-linker. After quenching the reaction, the complex was precipitated with acetone, followed by Western blot analysis with antibodies against CSNAP and CSN subunits. The data revealed that CSNAP forms a cross-link with CSN3, CSN5 and CSN6. (B) An overall view of the CSN crystal structure, showing the surfaces of the eight subunits (Lingaraju et al., 2014). The black frame delineates the region where

CSN3, CSN5, and CSN6 are found in close proximity. This region enlarged in (C) and (D), includes the C-terminal helices of CSN3 and CSN6, and the loop 284-295 of CSN5, which connects its two C-terminal helices. (C) The electrostatic potential on the surfaces of CSN3, CSN5 and CSN6: blue for positive, red for negative, and white for neutral. The highly positive patch of CSN3 is seen at the top left. Exposed lysine residues which may be involved in cross-linking are indicated: K237, K243, K254, and K312 of CSN3; K191, K194, and K299 of CSN5; and K306 of CSN6. The positive patch includes several hydrophobic pockets, which are preferred anchoring sites of Phe residues, as predicted by ANCHORSmap. Anchored Phe side chains with $\Delta G < -4$ Kcal/mol are shown in yellow. (D) Superposition of the PCI domain of PCID2 (brown) in the complex with DSS1 (Ellisdon et al., 2012), onto the PCI domain of CSN3 (gray). The F/D-rich region of DSS1 (golden coil) binds to the front face of PCID2, which corresponds to the positive patch of CSN3.

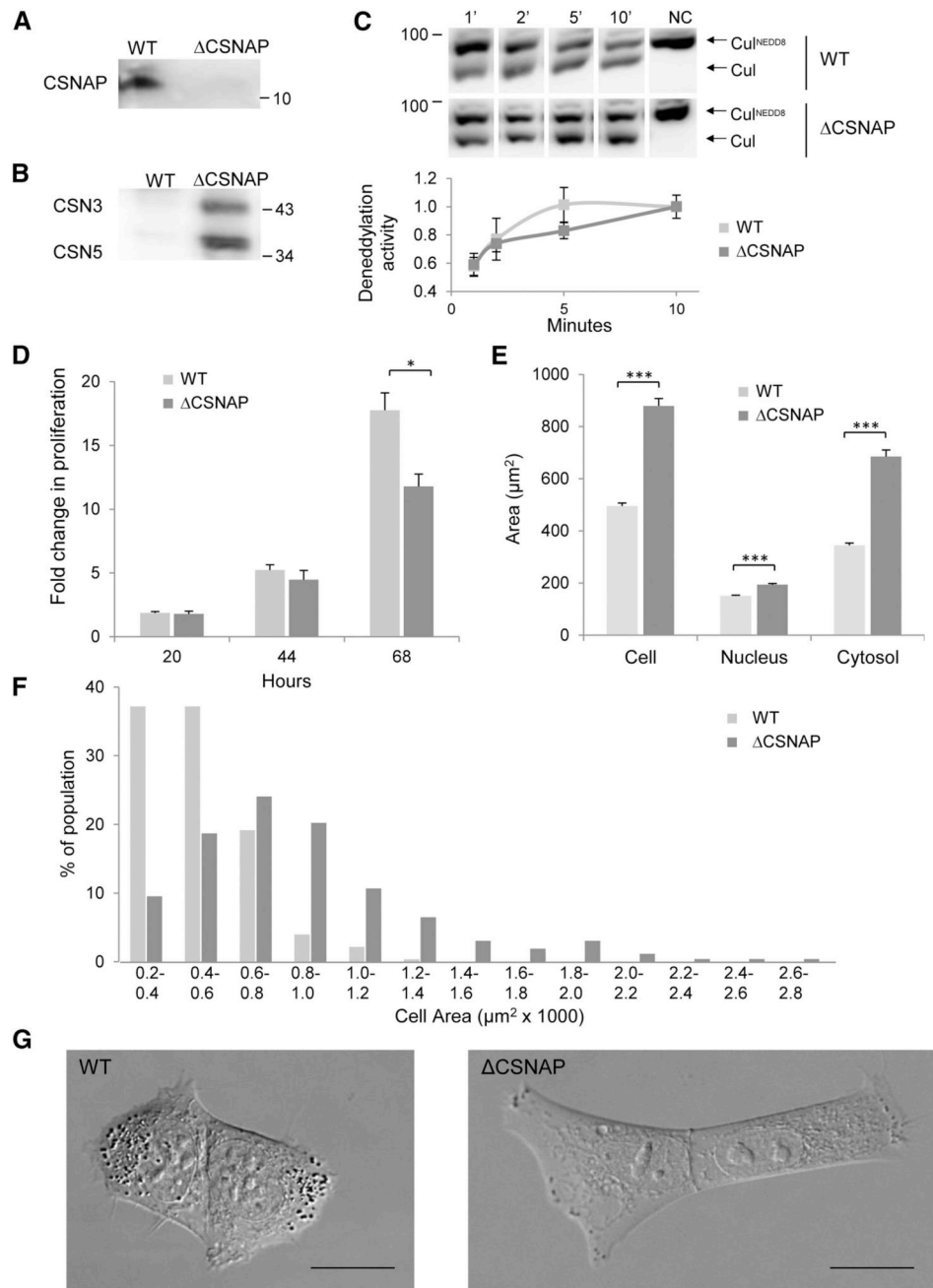


Figure 6. CSNAP-depleted cells display a distinctive phenotype.

(A) The CSN complex was FLAG-affinity purified from wild-type (WT) or from CSNAP knockout cells (Δ CSNAP), stably expressing FLAG-CSN1. CSNAP was detected by Western blot using an anti-CSNAP antibody only in the complex isolated from WT but not Δ CSNAP cells. (B) The WT CSN complex is saturated with endogenous CSNAP. Lysates from WT and Δ CSNAP cells stably expressing FLAG-CSN1 were passed through Aminolink beads coupled to a CSNAP-peptide. Only the CSN complex from the Δ CSNAP but not from the WT lysate, could bind to the beads. (C) WT and Δ CSNAP cells exhibit a

similar rate of deneddylation. Deneddylation was monitored at different time points. A representative deneddylation assay (top panel) and a plot demonstrating the average activity of three independent experiments (bottom panel). As a negative control (NC), lysates were denatured (boiled) prior to the assay. Bars represent standard errors. (D) CSNAP cells exhibit lower proliferation rates, as measured using the resazurin proliferation assay. Plot represents the average proliferation rate of three independent experiments. Measurements were subjected to T-test analysis; * indicates $p < 0.05$. (E) CSNAP cells are flatter and larger than WT cells. Cells plated at a low density were imaged using a confocal microscope. Partially dispersed cells displaying distinct cellular borders, were used to measure the cellular area of WT ($n=277$) and CSNAP cells ($n=263$). Bars represent standard errors. Measurements were subjected to T-test analysis; *** indicates $p < 10^{-15}$. (F) Size distribution of WT and CSNAP cell areas, represented as percent of the total population. (G) A representative confocal image of WT and CSNAP cells. Bar represents 20 μm .

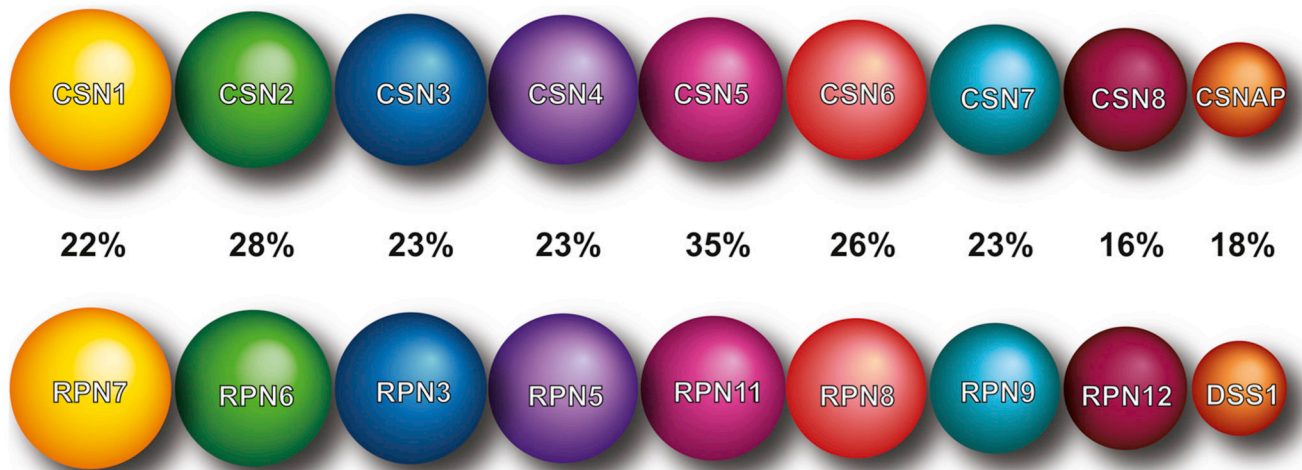


Figure 7. DSS1 and CSNAP are intrinsically disordered proteins that share sequential and functional homology.

Schematic illustration of the one-to-one sequence homology between the CSN and 19S proteasome lid complexes. CSNAP may represent the missing homologous partner of DSS1. The percentage of sequence identity between the subunits is labeled for each pair.