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Structure of an Rrp6-RNA exosome complex bound to polyA RNA

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Abstract

The eukaryotic RNA exosome processes and degrades RNA by directing substrates to the distributive or processive 3' to 5' exoribonuclease activities of Rrp6 or Rrp44, respectively. The non-catalytic nine-subunit exosome core (Exo9) features a prominent central channel. While RNA can pass through the channel to engage Rrp44, it is not clear how RNA is directed to Rrp6 or if Rrp6 utilizes the central channel. Here, we report a 3.3 Å crystal structure of a ten-subunit RNA exosome complex from *Saccharomyces cerevisiae* composed of the Exo9 core and Rrp6 bound to single-stranded polyA RNA. The Rrp6 catalytic domain rests atop the Exo9 S1/KH ring above the central channel, the RNA 3' end is anchored in the Rrp6 active site, and the remaining RNA traverses the S1/KH ring in an opposite orientation to that observed in a structure of a Rrp44-containing exosome complex. Solution studies with human and yeast RNA exosome complexes suggest that the RNA path to Rrp6 is conserved and dependent on the integrity of the S1/KH ring. While selection to Rrp6 or Rrp44 is stochastic *in vitro*, the fate of a particular RNA may be determined *in vivo* by the manner in which co-factors present RNA to the RNA exosome.

The eukaryotic RNA exosome core is an essential nine-subunit complex formed by two stacked rings of six RNase PH-like proteins on the bottom and three S1/KH domain “cap” proteins on the top. Exo9 associates with the endo- and 3' to 5' processive exoribonuclease Rrp44, and the 3' to 5' distributive exoribonuclease Rrp6 (ref. 1). In *Saccharomyces cerevisiae*, the cytoplasmic exosome includes Exo9 and Rrp44 (Exo10^{Rrp44}), the nuclear exosome contains Exo9, Rrp44 and Rrp6 (Exo11^{Rrp44/Rrp6})², and a nucleolar exosome with

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Author Contributions E.V.W. and C.D.L. designed and E.V.W. performed experiments for the *S. cerevisiae* exosomes. E.V.W. and C.D.L. determined the structure. K.J. and C.D.L. designed and K.J. performed the experiments for the *H. sapiens* exosome. E.V.W. and C.D.L. wrote the manuscript.

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Exo9 and Rrp6 (Exo10^{Rrp6}) has been posited in human³. Each exosome complex may be uniquely equipped to target different RNA substrates, in conjunction with protein cofactors, to catalyze RNA turnover, quality control, or processing in the context of their respective cellular compartments⁴.

Rrp6 is a RNase D family member⁵, and is proposed to hydrolyze RNA via two metal ion catalysis^{6,7}. It includes an N-terminal PMC2NT domain that associates with a cofactor Rrp47 (ref. 8), an exoribonuclease domain (EXO), an HRDC domain, and a C-terminal domain (CTD) that associates with the Exo9 core⁹. Structural studies revealed that the EXO and HRDC domains constitute the catalytic module^{10,11} while a Exo10^{Rrp44+Rrp6Cterm} structure revealed how Rrp6 CTD residues interact with Exo9 (ref. 12). No structures yet exist for Rrp6 in complex with Exo9 or RNA.

We proposed that Rrp44 and Rrp6 utilize an overlapping channel within the S1/KH ring to engage RNA, that Rrp6 activities are modulated by Exo9, and that Rrp6 stimulates Rrp44 in binding and degradation of ssRNA in the context of Exo11^{Rrp44/Rrp6} (ref. 13). To provide a structural basis for these observations, we crystallized Rrp6 with Exo9 and a 24-nt single-stranded polyA RNA (polyA₂₄ RNA).

Global architecture of Exo10^{Rrp6}-polyA RNA

The 3.3 Å structure of Exo10^{Rrp6} was obtained in the presence of a 24-nucleotide (nt) single-stranded polyA RNA using Rrp6 (128-685) that lacked exoribonuclease activity (D238N)^{13,14}, the PMC2NT domain (1-127)¹³ and its last 48 C-terminal residues (Extended Data Fig. 1; Extended Data Table 1). Rrp6 is positioned atop the Exo9 S1/KH ring with Exo9 subunits resembling those of human Exo9 (ref. 15) and yeast Exo9 (ref. 12) in Exo10^{Rrp44+Rrp6Cterm} (Fig. 1a, b). PolyA₂₄ RNA is coordinated within the S1/KH ring with Rrp6 active site residues contacting the RNA 3' end (Fig. 1c). The Rrp6 EXO domain contacts two of the three S1/KH ring proteins, Rrp4 and Rrp40, and the HRDC domain is proximal to Rrp4, although no direct contacts are apparent. Consistent with previous results¹², the Rrp6 CTD wraps around the N-terminal domain (NTD) of Csl4 and PH-like subunit, Rrp43, before emerging at the top of the Exo9 core (Fig. 1a; Extended Data Fig. 2a,b). No electron density was observed for CTD residues 629-684 and for residues (517-524) that link the HRDC domain and CTD. The location of the Rrp6 N-terminus (Met128) places the Rrp6 PMC2NT domain over the Exo9 central channel in an ideal position to interact with Rrp47 to facilitate substrate recruitment (Fig. 1a). Alignment of exosome-associated Rrp6 to the yeast Rrp6 catalytic module (EXO-HRDC) reveals few differences (Extended Data Fig. 2c).

The NTD of Rrp4 and S1 domains of Rrp4 and Rrp40 interact with the Rrp6 EXO domain burying 2230 Å² of surface area in the complex. Rrp6 interactions with the Rrp4 NTD (Region 2) and S1 domain (Region 3) are more extensive (1750 Å²) while interactions with the Rrp40 S1 domain and Rrp4 S1 domain encompass a region proximal to the RNA binding site (Region 1). These surfaces are highly conserved (Fig. 2a,b) in comparison to the Rrp6 CTD (Extended Data Fig. 2a), which is important for Rrp6 interaction with the exosome

core as illustrated by analytical gel filtration studies showing the CTDs of both yeast and human Rrp6 are required for association with the exosome (Extended Data Fig. 3).

The structure of Exo10^{Rrp6} bound to polyA₂₄ RNA reveals how an RNase D family member interacts with RNA (Fig. 2c). The Rrp6 catalytic domain and active site bind A₂₁A₂₂A₂₃A₂₄ with the 2'-OH and 3'-OH of the terminal A₂₄ coordinated via two main chain hydrogen bonds to the backbone amide and carbonyl oxygen of His241 and Glu240 side chain carboxylate, similar to that observed for AMP in the Rrp6 catalytic domain¹⁰. An additional contact is observed to A₂₄ between the N7 adenine atom and Gln345 side chain amide. The scissile phosphate of A₂₄ and A₂₃ is coordinated by a single magnesium ion that bridges the phosphate and the side chain carbonyl of Asn238 (Asp in wild-type enzyme). The scissile phosphate is within hydrogen bond distance of the invariant side chain hydroxyl of Tyr361 (Extended Data Fig. 4a,c). Consistent with the proposed reaction mechanism⁷, two metal ions coordinated by conserved side chain carboxylates in the yeast Rrp6-AMP complex are ideally positioned to contact the scissile phosphate (Extended Data Fig. 4c,d). Several contacts between metal ions and RNA are missing in our structure due to the D238N substitution.

The remaining contacts to polyA₂₄ RNA reveal several non-specific interactions, consistent with the ability of Rrp6 to degrade RNA of any sequence. A₂₃ 2'-OH is within hydrogen bonding distance to the side chain carboxylate of Asp296 while the A₂₃ phosphate is within hydrogen bonding distance of the Leu328 backbone amide. The A₂₃ base is situated over the A₂₄ base with van der Waals contacts provided by side chains of Trp299, Met295 and Phe294. Phe294 is positioned between the A₂₃ and A₂₂ bases, presumably disrupting base stacking interactions. The A₂₂ base and ribose are cradled by van der Waals contacts to Phe294, Gly292, His291 and Tyr315 while its phosphate is within hydrogen bonding distance with the backbone amide of His291. Lys319, His326 and Tyr315 contribute the remaining contacts to the A₂₁ base. Importantly, none of the aforementioned contacts appear to interrogate the identity of the base. Sixteen of eighteen residues observed in direct contact to the RNA or metal cofactors are conserved in human Rrp6 (Extended Data Fig. 4a,b)^{10,11}.

RNA passes through the S1/KH ring

Four of six nucleotides of polyA₂₄ RNA observed in electron density (5'-A₁₉-A₂₄-3') are coordinated by Rrp6 placing the 5' end (A₁₉) proximal to conserved, basic side chains from Rrp4 (Lys122 and Arg123), while Rrp40 Lys108 points toward A₂₄ and Arg110 is proximal to the phosphate of A₂₁ and ribose of A₂₀ (Fig. 3a). Additional densities with spacing consistent with the RNA phosphate backbone, modeled as PO_{4a} and PO_{4b}, are observed within the S1/KH ring nearest to Arg150 and proximal to Arg145 and Arg202 of Csl4 (Extended Data Fig. 1a-c). The aforementioned Csl4, Rrp4 and Rrp40 residues are highly conserved¹⁵. No additional density consistent with RNA is observed within the PH-like ring.

Rrp6 and Rrp44 utilize a shared portion of the central channel to engage RNA substrates¹³, and alignment of Exo10^{Rrp6} and Exo10^{Rrp44+Rrp6Cterm} reveals overlapping paths for RNA within the S1/KH central channel (Fig 3a,b), albeit in opposing directions. A subset of side chains from the S1/KH cap proteins contact RNA in both structures, but none make contacts

that would enforce directionality. To determine the importance of the S1/KH ring for RNA degradation, yeast Exo10^{Rrp6} and Exo11^{Rrp44/Rrp6} were reconstituted with Csl4 and Rrp40 containing amino acid substitutions for Arg145, Arg150, Arg202; and Lys107, Lys108, and Arg110, respectively. These mutations severely diminish Rrp6 activity in Exo10^{Rrp6} and Exo11^{Rrp44/Rrp6} as well as Rrp44 activity in Exo11^{Rrp44/Rrp6} (Fig. 3c; Extended Data Fig. 5a). In contrast, Rrp6 activity is not inhibited by an insertion in Rrp45 that occludes the PH-like ring channel below the S1/KH ring¹³ although this mutation inhibits Rrp44 activity. Reconstituted human Exo10^{Rrp6} (Extended Data Fig. 3b) reveals similar dependencies as a channel occluding insertion in Rrp41 near the S1/KH ring impaired Rrp6 activity while a channel occluding insertion in Rrp45 well below the S1/KH ring failed to inhibit Rrp6 (Fig. 3d; Extended Data Fig. 5b). Combining insertions inhibits Rrp6 to greater degree, likely because the Rrp45 insertion restricts movement of the Rrp41 insertion. These data suggest that the RNA path to Rrp6 depends on the integrity of the S1/KH ring but does not require the full extent of the PH-like ring central channel. In contrast, at least for yeast Exo11^{Rrp44/Rrp6}, the RNA path to Rrp44 relies on the integrity of central channel throughout both S1/KH and PH-like rings. These data are consistent with contacts observed to RNA in structures of Exo10^{Rrp6} and Exo10^{Rrp44+Rrp6Cterm}.

To map RNA interactions through Exo9 to Rrp6 and Rrp44, catalytically dead yeast exosome complexes were reconstituted and subjected to long wavelength UV crosslinking using 36-nt polyA or AU-rich RNA substrates bearing single 4-thioU substitutions 6, 21, or 29 nt from the 3' end (Fig. 4a, Extended Data Fig. 6a-c). As the 4-thioU probe is moved 3' to 5', AU-rich RNA crosslinks to Rrp44, followed by the PH-like ring, and finally to the S1/KH cap in Exo10^{Rrp44} and Exo11^{Rrp44/Rrp6}, in agreement with previous studies^{12,13,16,17}. In contrast, crosslinks are only observed to Rrp44 in Exo10^{Rrp44} with polyA₃₆ RNA; however, addition of Rrp6 in Exo11^{Rrp44/Rrp6} results in crosslinks that progress from Rrp44 through the PH-like ring to the S1/KH ring and Rrp6 as the 4-thioU probe is moved 3' to 5'. The results with polyA₃₆ RNA are in line with previous biochemistry showing that Rrp6 stimulates Rrp44 binding to polyA RNA in Exo11^{Rrp44/Rrp6} (ref 13).

Crosslinking to the Exo10^{Rrp6} complex reveals RNA contacts to Rrp6 and Rrp4 when 4-thioU is 6 nt from the 3' end. Although no structural impediment exists to prevent RNA from entering the PH-like ring, polyA₃₆ and AU-rich RNA crosslinks are only observed to Rrp4, Csl4 and Rrp40 even when the 4-thioU is positioned 21 or 29 nt from the 3' end (Fig. 4c; Extended Data Fig. 6). This pattern contrasts with that observed for Exo10^{Rrp44} and Exo11^{Rrp44/Rrp6}, where crosslinks are observed to PH-like ring subunits when 4-thioU is placed 21 nt from the 3' end. Crosslinking patterns to human Exo10^{Rrp6} utilized a 36-nt AU-rich substrate as reported previously¹³. Similar to results obtained for yeast, UV-induced crosslinks are observed to the three S1/KH ring proteins and RRP45, but to none of the other five PH-like proteins (Fig. 4b). UV crosslinking to complexes with loop insertions mirror results observed in decay assays (Fig. 3d) with diminished crosslinking when a loop insertion is placed proximal to the S1/KH ring (Extended Data Fig. 7). These data suggest that the integrity of the S1/KH ring central channel is important for Rrp6 activity in both human and yeast systems.

The results are consistent with a model in which distinct but overlapping paths guide RNA to Rrp6 or to Rrp44 (Fig. 4c). So how is a path selected? Path selection appears stochastic *in vitro* because degradation products of Rrp6 and Rrp44 are observed under conditions of limiting enzyme (Fig. 3c) or limiting substrate (Fig. 4d). The distributive mechanism underlying Rrp6 activity suggests repeated substrate binding and release while the processive mechanism utilized by Rrp44 suggests that it binds and holds onto substrates until completely degraded. Thus at steady state, binding and UV crosslinking likely reflect the stable interaction with Rrp44 even when Rrp6 is present¹³ (Extended Data Fig. 6a). Additional evidence for stochastic sampling of the two paths is evident by UV crosslinking under conditions of slight enzyme excess (Fig. 4d). As predicted based on the distributive and processive mechanisms of Rrp6 and Rrp44, respectively, crosslinked products are observed to Rrp44, Rrp6 and the S1/KH ring proteins at the earliest times, and this pattern is lost once most of the RNA finds its way to the Rrp44 active site.

Structural analysis of the Exo10^{Rrp6} polyA complex suggests at least four potential paths past the S1/KH ring to Rrp6 although paths 1 and 2 appear most likely with respect to electrostatics and conservation (Fig 4e; Extended Data Fig. 8a). Importantly, these paths are available in Exo11^{Rrp44/Rrp6} as they do not involve surfaces from the PH-like ring or Rrp44. Modeling Rrp6 onto the Exo10^{Rrp44} RNA complex shows that the central channel is still accessible and that RNA paths to Rrp6 and Rrp44 are available in Exo11^{Rrp44/Rrp6} (Fig 4e; Extended Data Fig. 8b).

A wider channel in Exo10^{Rrp6}

Rrp6 can stimulate Rrp44 binding and decay activities¹³, a phenomenon readily apparent in crosslinking to polyA RNA (Fig. 4a; Extended Data Fig. 6a). These results suggest that Rrp6 enhances RNA access to the PH-like ring and central channel. Structures of Exo10^{Rrp6} and Exo10^{Rrp44+Rrp6Cterm} were compared to query differences that might account for this activity. While the global architecture of the PH-like ring does not differ (Fig. 5d), the Exo9 channel widens in Exo10^{Rrp6} through movement of Rrp4, Rrp40 and Csl4 away from the central channel, widening the gap between Rrp4 and Rrp40 S1 domains by ~4 Å (Fig. 5a-c). Although Exo10^{Rrp6} and Exo10^{Rrp44+Rrp6Cterm} structures are bound to RNA, the increase in channel width in Exo10^{Rrp6} might account for Rrp6-mediated stimulation of Rrp44, especially since Rrp44 and Rrp6 do not physically interact. It remains unclear how Rrp6 exerts this change or if an Exo10^{Rrp44} apo structure differs from its RNA-bound configuration, but it is clear that Rrp6 EXO and CTD are both required for this activity as addition of either element alone is not sufficient to stimulate Exo10^{Rrp44} when added *in trans* (Fig. 5e). These data suggest that the Rrp6 CTD is required to bring the catalytic module in proximity to the S1/KH ring, perhaps eliciting channel widening through EXO domain interaction with Rrp4 and Rrp40. Further data will be required to determine if channel widening is a regulated feature of Exo11^{Rrp44/Rrp6} or if additional cytoplasmic factors elicit channel widening of Exo10^{Rrp44}.

Conclusions

The structure of Exo10^{Rrp6} shows Rrp6 positioned above the Exo9 S1/KH ring while the Exo10^{Rrp44+Rrp6Cterm} structure shows Rrp44 below the PH-like ring. It is notable that Rrp6 activity is altered and becomes dependent on the S1/KH ring when associated with Exo9 (ref. 13). Although shorter RNAs may be directed to Rrp44 via a channel-independent “direct access” route¹⁸, Rrp44 remains highly dependent on the integrity of the central channel throughout both S1/KH and PH-like rings. The dependency on the Exo9 core is remarkable given that both Rrp44 and Rrp6 active sites are exposed to solvent in Exo10^{Rrp6} and Exo10^{Rrp44+Rrp6Cterm} complexes.

That a similar segment of the S1/KH ring is used to engage RNA in opposing directions suggests that overlapping paths to Rrp6 and Rrp44 may serve to commit the exosome to distributive or processive degradation depending on how a particular RNA substrate is delivered to the exosome. Because the paths overlap, the exosome would be unable to interact with another substrate until completing the task at hand. While path choice appears stochastic *in vitro* (Fig. 4d), the nuclear cofactors Mpp6 (ref. 19), Rrp47 (ref. 8), and TRAMP²⁰ may bias selection of a particular path to facilitate transitions between editing, processing or degradation activities of the exosome.

Rrp6 can stimulate Rrp44 activities when associated with the Exo9 core, perhaps through widening of the S1/KH ring. Channel gating mechanisms as a point of regulation have been described in many systems including, but not limited to, the proteasome²¹⁻²⁵. While many degradation complexes place their catalytic activities in a protected compartment that is accessed by gating a substrate responsive channel, the RNA exosome appears inside out, with its catalytic subunits located on the periphery of the exosome core. While the functional relevance of this architecture is not yet clear, it is evident that the exosome core can modulate the activities of its catalytic subunits by requiring RNA to pass through distinct elements of the central channel before being processed or degraded.

Methods

Yeast exosome reconstitution and purification

Expression, purification, and reconstitution of recombinant exosome subunits and complexes have been previously described in detail^{13,26}. In brief, all subunits were cloned as N-terminal 6× histidine Smt3 fusions in pRSFDuet-1 expression vectors (Novagen), and expressed in the *E. coli* expression strain BL21 (DE3) RIL (Stratagene) either as heterodimers (Smt3-Rrp41/Rrp45, Smt3-Rrp42/Mtr3, Smt3-Rrp46/Rrp43) or as distinct subunits (Smt3-Csl4, Smt3-Rrp4, Smt3-Rrp40, Smt3-Rrp6, Smt3-Rrp44). Generation of the Rrp40 and Csl4 point mutants was performed by PCR-mediated site-directed mutagenesis. For the mutant subunits, cells were grown in shaker flasks at 37°C in Superbroth (Teknova), induced by cold shock on ice, addition of ethanol to a final concentration of 2% and 0.05 mM isopropyl-β-D-thiogalactoside (IPTG), and grown at 18°C overnight. For all other proteins, cells were grown in shaker flasks at 37°C in Superbroth and induced by cold shock on ice and addition of 0.4 mM IPTG, and then subjected to shaking at 18°C overnight. Cells were harvested and lysed as described previously. After high speed centrifugation to pellet

the insoluble material, the supernatant was loaded on a nickel-NTA column (Qiagen) and allowed to flow by gravity. After washing with 350 mM NaCl, 20 mM Tris pH 8.0, 1 mM BME, and 20 mM imidazole, the column was further washed with 350 mM NaCl, 50 mM KCl, 20 mM Tris pH 8.0, 1 mM BME, 2 mM ATP, 10 mM MgSO₄ to displace chaperone impurities. Protein was eluted with 350 mM NaCl, 20 mM Tris pH 8.0, 1 mM BME, and 250 mM imidazole and then purified by size exclusion chromatography. All subunits were purified on the Superdex 200 (GE) with the exception of Smt3-Csl4 and Smt3-Rrp40, which were purified on a Superdex 75 (GE). Wild-type and mutant subunits eluted at the same volumes on gel filtration. Only Smt3-Rrp6, Smt3-Rrp44, and Smt3-Rrp42/Mtr3 were subjected to overnight cleavage by the SUMO protease Ulp1 (ref. 29) with another step of purification on the Superdex 200 to remove the 6x histidine Smt3 tags; Smt3-Rrp42/Mtr3 was cleaved by Ulp1 prior to formation of Exo9 as inclusion of this tag interferes with reconstitution. At this stage, subunits were concentrated to 6 to 12 mg/mL and stored at -80°C. For reconstitution, Exo9 subunits were mixed together as Smt3 fusions, incubated on ice for 30 minutes, followed by addition of Ulp1 for another 30 minutes prior to overnight dialysis. After purification, exosomes were concentrated to 12-14 mg/mL and stored at -80°C in a final buffer of 100 mM NaCl, 20 mM Tris pH 8.0, 0.1 mM MgCl₂, 1 mM TCEP.

Human exosome reconstitution and purification

For human, expression, purification and reconstitution strategies of the Exo9 core were identical except Smt3-RRP43/MTR3/RRP42 was expressed as a trimer, Smt3-RRP46 was expressed alone (as described previously^{15,26}), and RRP45¹⁻³⁰²/RRP41 was purified with a non-cleavable N-terminal hexa-histidine tag on RRP45. RRP6 (EXOSC10, PM/SCL-100) residues 180 to 804 were PCR-amplified from a previously generated full-length expression construct¹¹, cloned into pET28a-Smt3 and transformed into BL21 (DE3) RIL. For expression, cells were grown in shaker flasks to ~0.6 OD₆₀₀, and induced overnight at 18°C with 0.25 mM IPTG in the presence of 2% ethanol. Subsequent purification of human Smt3-RRP6¹⁸⁰⁻⁸⁰⁴ and reconstitution of HsExo10^{Rrp6} is identical to that described for the yeast system. The purification method for the catalytic region, RRP6¹⁸⁰⁻⁶⁰⁶, used for core interaction studies has been described previously¹¹. The channel occluding RRP41 and RRP45 mutants were engineered based on those described for *S. cerevisiae*¹³, and were generated by PCR to encode electrostatic and steric loop insertions with the primary sequence GTGESEGESES between amino residues Gly93 and Arg94 of RRP45 and between Arg62 and Ala63 of RRP41.

Crystallization and structure determination

Exosome samples were mixed with polyA₂₄ RNA (Invitrogen) in a 1:1.1 molar ratio and incubated on ice for 1 hour prior to crystallization. Crystals grew at 18°C by vapor diffusion in either sitting (Greiner Bio One, Crystalquick) or hanging (Hampton Research, VDX) drop formats, in 7-11% PEG3350, 100 mM MES pH 6.7, 4-15% MPD, typically taking 3 to 5 days to appear. Crystals were harvested within 2 weeks. Partial degradation of RNA (20-24 nt) was observed in washed crystals (Extended Data Fig. 1d). For cryoprotection, well solution was replaced with crystallization buffer augmented with 25-30% MPD and crystals were incubated for three days before harvesting; one day before harvest, trays were transferred to 4°C. A light polarizer was used to exclude multiple crystals and to identify

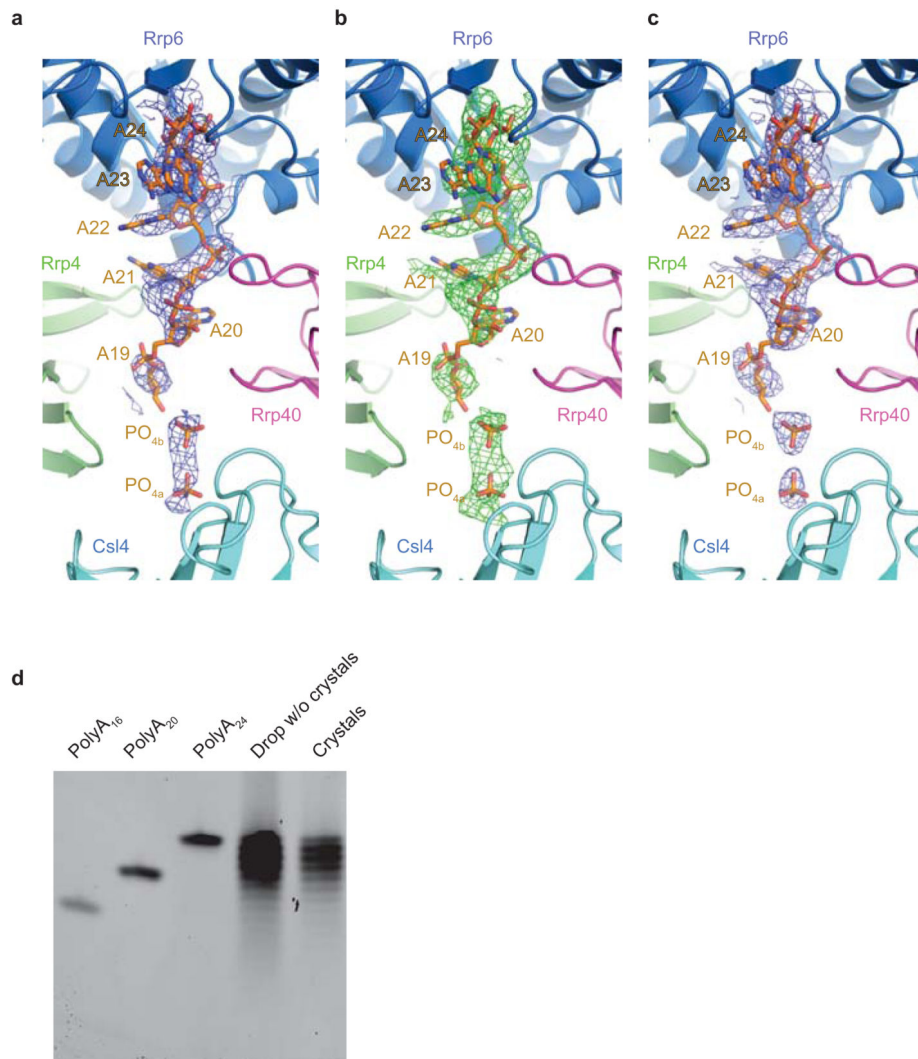
crystals with single regions, which were broken off and flash frozen in liquid nitrogen for data collection. X-ray diffraction data were collected at the Advanced Photon Source 24-ID-E and 24-ID-C beam lines, and the National Synchrotron Light Source X29 beam line. Data was obtained from a single crystal diffracted at NSLS X29 at a wavelength of 1.075 Å at 100 K. Data were processed using HKL2000³⁰ and statistics reported in Extended Data Table 1 were obtained using Phenix³¹ including $CC_{1/2}$ (%) and CC^* (%) values of 99.9 (33.8) and 100.0 (71.1) for data between 50-3.3 (3.42-3.3) Å. The structure solved by molecular replacement using Phaser³² and coordinates of human apo Exo9 (PDB: 2NN6) and yeast Rrp6^{CAT} (PDB: 2HBL) as search models followed by docking yeast Exo10^{Rrp44+6Cterm} (PDB: 4IFD) (Extended Data Table 1). The structure is refined to R/R_{free} values of 0.227/0.265. The final model includes six nucleotides of the polyA 24mer, and two phosphate ions that likely represent the RNA backbone (Extended Data Fig. 1a-c), and 2751 of the 3156 amino acids present in the crystal (Extended Data Table 2). The asymmetric unit contains one complex. Iterative rounds of refinement were accomplished using Phenix³¹. RNA and side chains were manually built using O³³ and Coot³⁴. The model was initially refined using secondary structure restraints in conjunction with positional refinement followed by individual B-factor refinement. Secondary structure restraints were released and a final round of positional and B-factor refinement was performed followed by refinement of TLS parameters that resulted in a further decrease in R and R_{free} values. Figures depicting the structure were prepared with Pymol²⁷. Surface conservation was calculated using ConSurf³⁵. Structure quality was assessed using MolProbity³⁶ indicating the model has excellent geometry with 95.7% in favored and 100% in allowed regions of Ramachandran space. The structure also scored in the 100th percentile for the Clash and MolProbity scores.

RNA biochemistry

For yeast exosomes, unless otherwise noted, exoribonuclease assays were performed under multiple turnover conditions as described previously¹³. In brief, 10 nM synthetic 49-nt RNAs (Invitrogen) bearing 5' fluorescein labels were incubated at 30°C with 1 nM RNA exosome for various time points, and RNA degradation was monitored by resolving reaction intermediates by denaturing TBE-PAGE (Invitrogen), and detected using a Fuji FLA-5000 scanner (FITC filter). A similar protocol was followed for human exosome complexes, the exception being that these assays were performed at 37°C, with 5 nM of the 49-nt RNA and 50 nM HsExo10^{Rrp6}. For mix-in experiments (Fig. 5e), 2-fold molar excess of Rrp6 protein was incubated with Exo10^{Rrp44} on ice for 1 hour prior to initiating RNA decay by addition of RNA. For steady state UV-RNA crosslinking to 4-thioU-bearing RNAs, 500 nM of exosome complexes with mutations in the exoribonuclease active sites (D238N for Rrp6, D551N for Rrp44) were incubated for 20 minutes on ice with 50 nM 36-nt 5' fluorescein RNAs with a single internal 4-thioU (Thermo Scientific) in a 70 µL reaction volume. Binding buffer includes 50 mM KCl, 20 mM Tris pH 8.0, 10 mM DTT, 0.5 mM MgCl₂. Crosslinking was performed by subjecting the RNA-exosome mixture to long-range UV (365 nm) for 15 minutes in the dark using a 4W handheld lamp. For the time course described in Fig. 4d, 350 nM of exosome was incubated with 150 nM of RNA, and the binding reactions allowed to proceed for indicated times before initiating crosslinking for 10 minutes. 15 µL was quenched with LDS loading buffer, and the crosslinked products were separated by SDS-PAGE (Invitrogen) and visualized with a Fuji FLA-5000 scanner (FITC

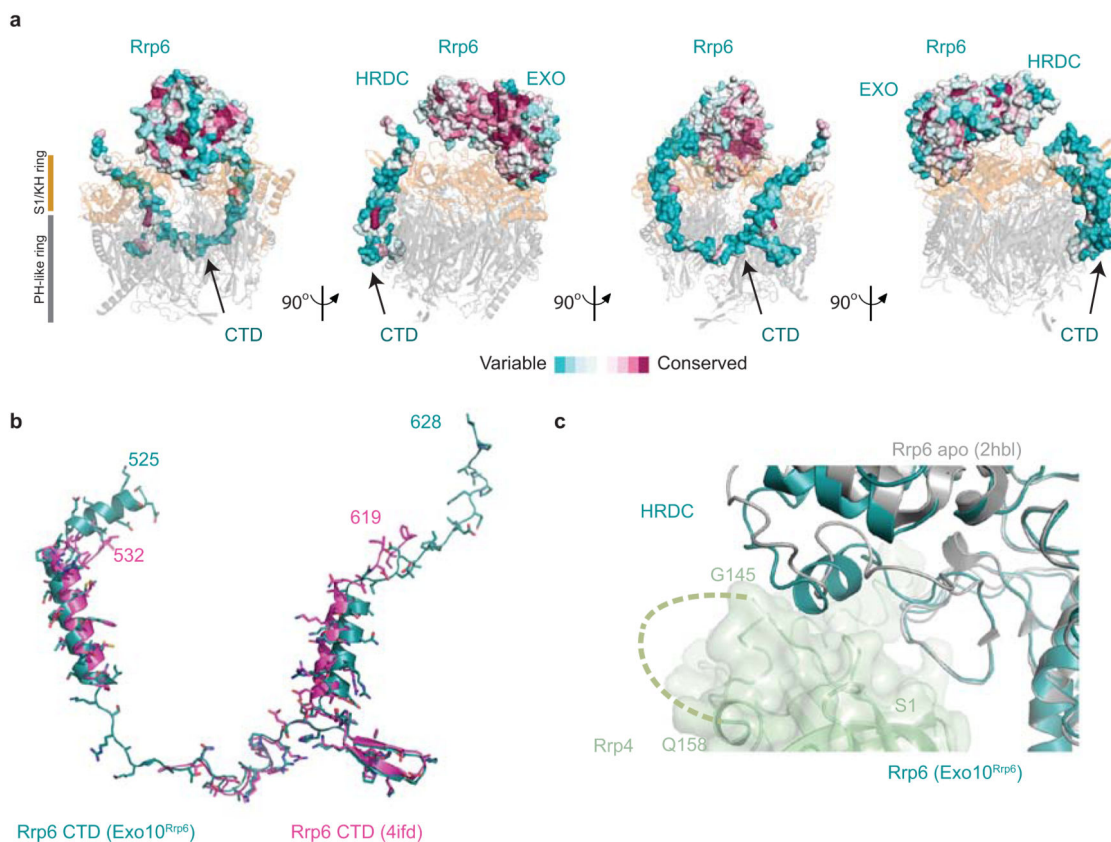
filter). 4-thioU RNA included the following sequences, each of which contained a 5'-Fluorescein: 5'-Fl-AAU UAU 4thioUUA UUA UUU AUU UAU UAU UUA UUU AUU UAA; 5'-Fl-AAU UAU UUA UUA UU4thioU AUU UAU UAU UUA UUU AUU UAA; 5'-Fl-AAU UAU UUA UUA UUU AUU UAU UAU UUA UU4thioU AUU UAA; 5'-Fl-AAA AAA 4thioUAA AAA AAA AAA AAA AAA AAA AAA AAA AAA; 5'-Fl-AAA AAA AAA AAA AA4thioU AAA AAA AAA AAA AAA AAA AAA; 5'-Fl-AAA AAA AAA AAA AAA AAA AAA AAA AA4thioU AAA AAA. For UV crosslinking to human exosome complexes, samples were incubated with 250 nM of the AU-rich RNA and protein in binding buffer for 60 minutes on ice, and then placed in a UV-Stratalinker (Stragene) and subjected to 300,000 μJ of short-wave (254 nm) UV radiation. Samples were quenched with LDS loading buffer, and crosslinked products were separated and visualized as described above.

Extended Data



Extended Data Figure 1. View highlighting electron densities covering RNA in the Exo10^{Rrp6}-polyA structure

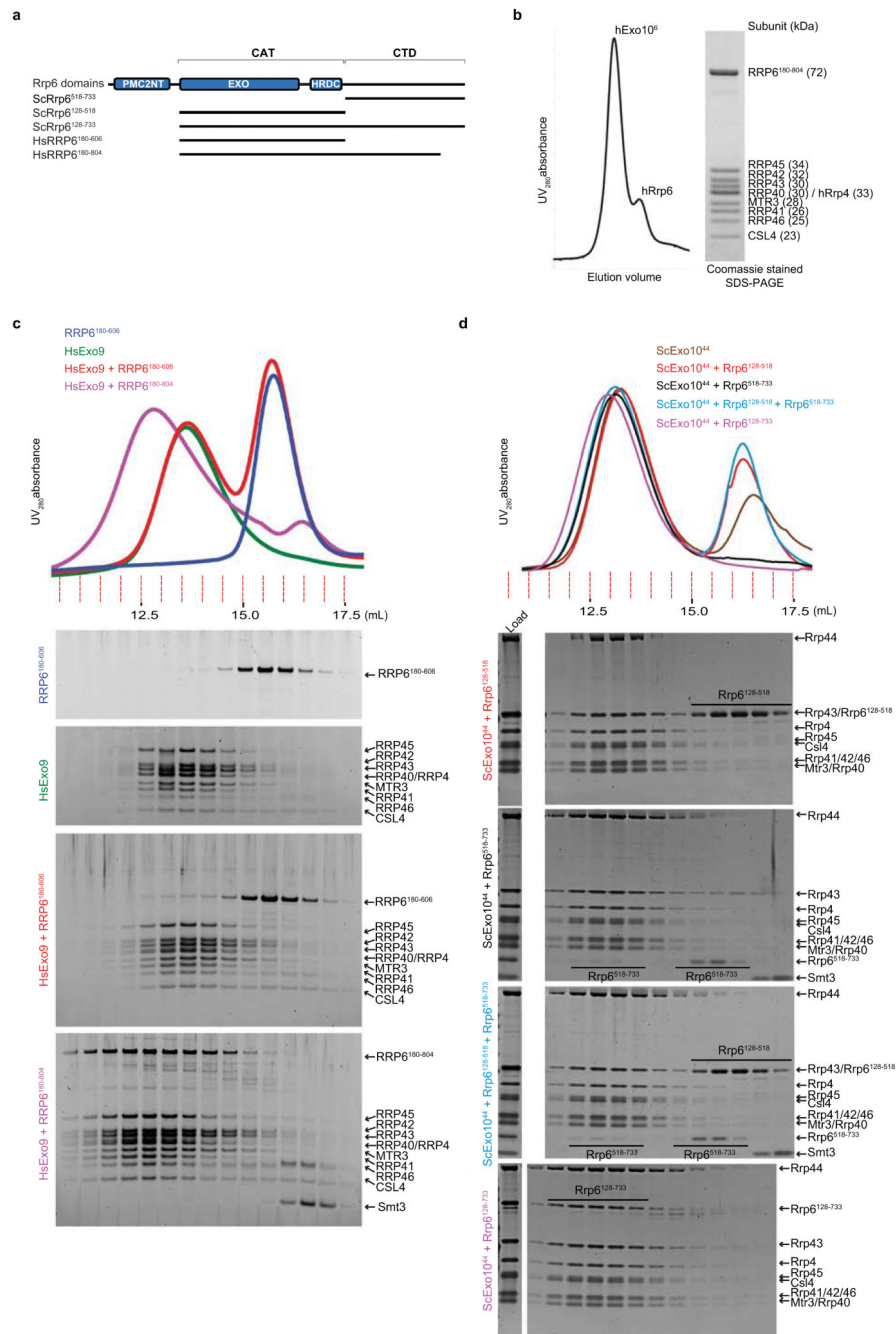
a, Simulated annealing $2F_o-F_c$ omit map contoured at 1.0σ . **b**, Simulated annealing F_o-F_c omit map contoured at 2.0σ . **c**, Final $2F_o-F_c$ omit map contoured at 1.0σ . RNA and phosphate oxygen, nitrogen, carbon and phosphate atoms are colored red, blue, orange and deep orange, respectively. Electron density maps shown as wire mesh colored blue for $2F_o-F_c$ maps and green for the F_o-F_c map. Rrp6, Rrp40, Rrp4 and Csl4 are labeled and shown in cartoon representation in teal, magenta, green and light blue, respectively. **d**, 20 to 24 nucleotides of polyA RNA are present in crystals of Exo10^{Rrp6}. Crystals of Exo10^{Rrp6} bound to polyA₂₄ RNA were first washed by two rounds of transfer to 1 μ L well solution, then dissolved in water and TBE-urea sample buffer, and analyzed by 15% TBE-urea PAGE. A drop without crystals was run as a control. RNA was stained by Sybr Gold.



Extended Data Figure 2. Rrp6 features within Exo10^{Rrp6}

a, The primary sequence of the catalytic module of Rrp6 (EXO and HRDC) is highly conserved, whereas sequence conservation within the CTD varies. Rrp6 in Exo10^{Rrp6} is represented as a surface, and colored according to sequence conservation as calculated by ConSurf³⁵, colored from red (highly conserved) to blue (variable). The Exo9 core is depicted as a transparent cartoon, with the S1/KH ring in orange, and the PH-like ring in grey. **b**, Superposed structures of the Rrp6 CTD from Exo10^{Rrp6} (residues 525-628; teal) and Exo10^{Rrp44+Rrp6Cterm} (residues 532-557, 565-619; magenta; PDB 4IFD) reveal overall similarity for residues within the respective models. **c**, The conserved linker between the

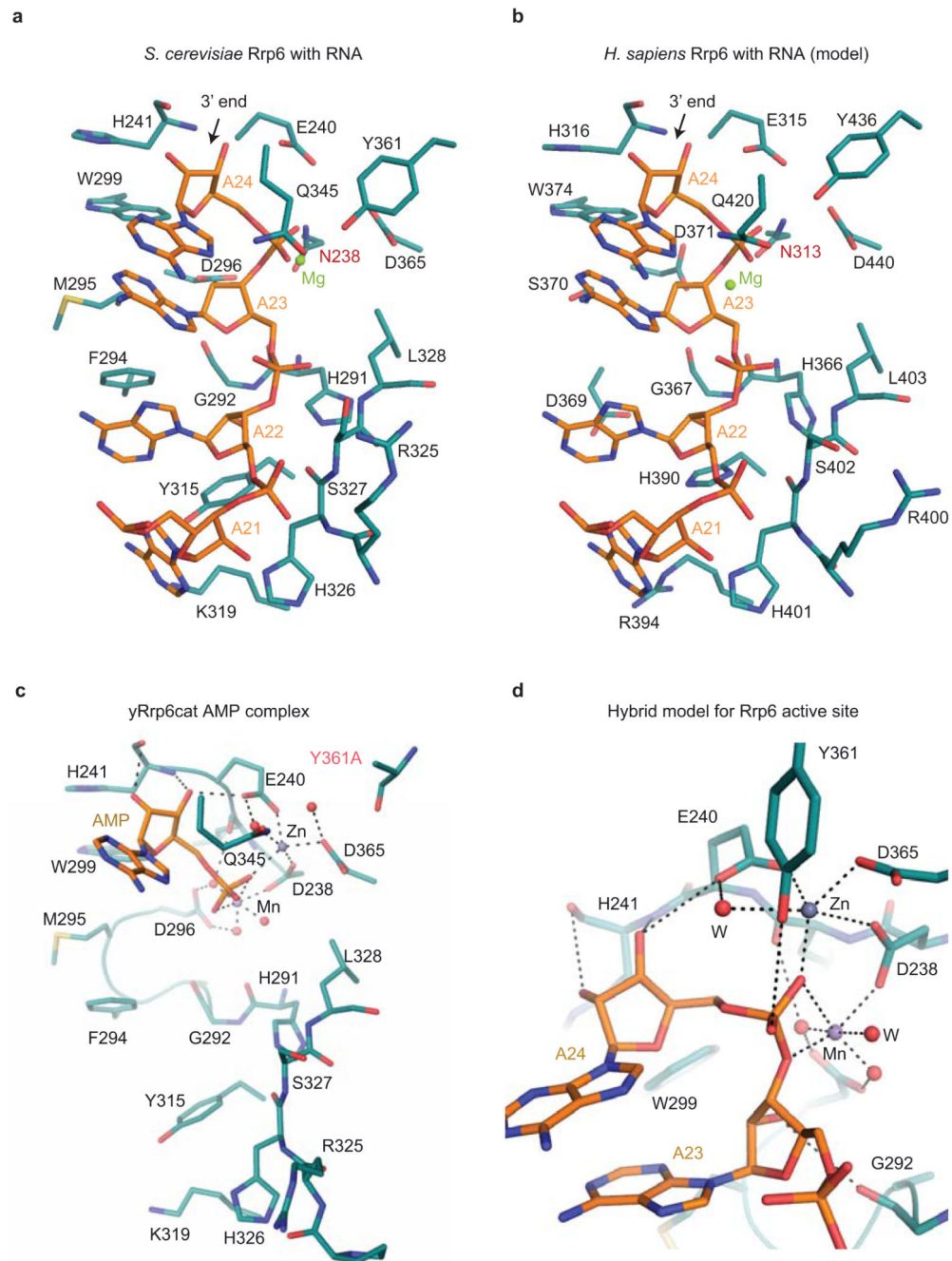
EXO and HRDC domain of Rrp6 adopts an α -helix when associated with the exosome core. Superposition of the Rrp6 catalytic domains from the structures of Exo10^{Rrp6} and Rrp6-AMP (PDB: 2HBL) shows residues Pro424 to Asn433 rearrange from a loop in the AMP-bound structure to a helix that is adjacent to a conserved albeit disordered loop in the S1 domain of Rrp4. These residues were previously thought to comprise a protein interaction module¹⁰. Rrp6 is represented as cartoons, with Rrp6 from the Exo10^{Rrp6} structure in teal, and the Rrp6-AMP structure in grey. Rrp4 is in light green and depicted as a cartoon with a transparent surface, with the S1 loop representing residues Gly145 to Gln158 shown as a dashed line.



Extended Data Figure 3. The Rrp6 CTD is required for interaction with human and yeast RNA exosome cores

a, Schematic representation of Rrp6 including from N- to C-terminus the PMc2NT, EXO, HRDC and C-terminal (CTD) domains. EXO and HRDC comprise Rrp6^{CAT}. Below are lines representing individual elements used in gel filtration analysis in **c** and **d** along with amino acid numbering specific for *S. cerevisiae* (Sc) Rrp6 and *H. sapiens* (Hs) RRP6. **b**, Gel filtration profile for reconstituted human Exo10^{Rrp6} (left) with SDS-PAGE analysis of the peak fractions (right). Subunits labeled in capital letters corresponding to yeast nomenclature for clarity. Apparent molecular weights denoted in kilodaltons. **c**, Interaction

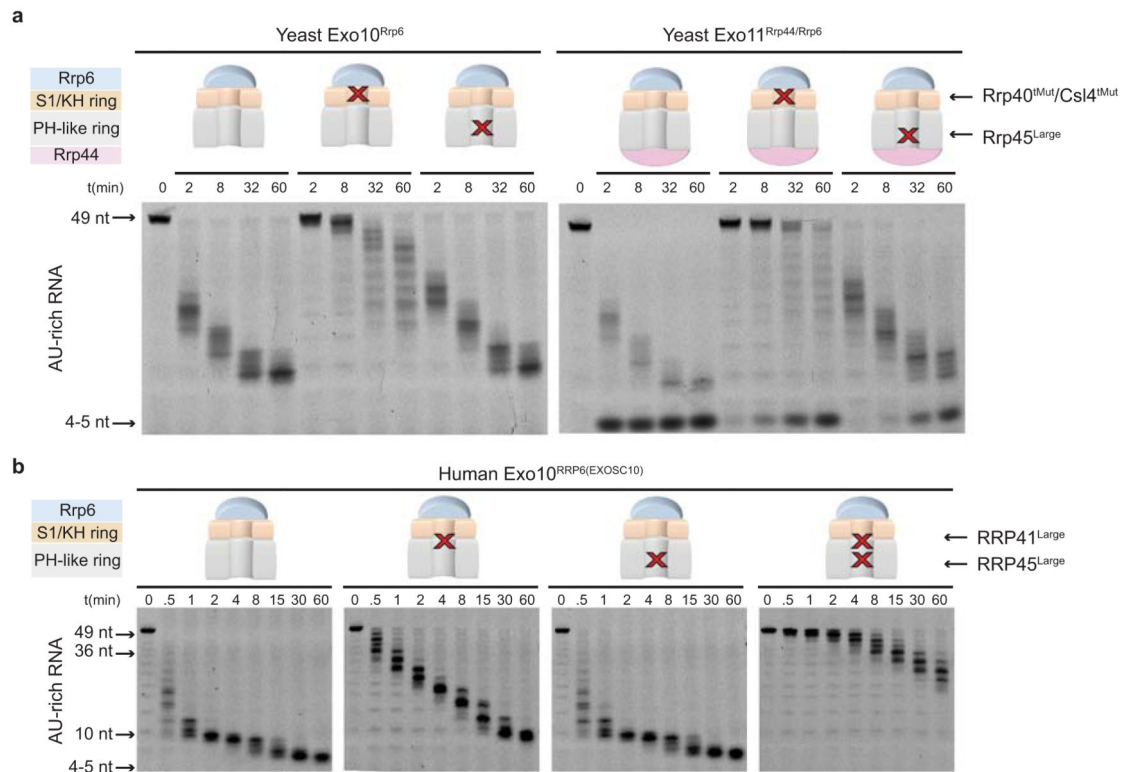
studies of human RRP6 with the HsExo9. The human Rrp6 catalytic domain (residues 180-606) does not interact well with human Exo9; however addition of Rrp6 180-804 suggests that CTD residues 607-804 are important for stable interaction with Exo9. Reconstitution experiments on mixtures of HsExo9, HsExo9 with 2-fold excess of RRP6¹⁸⁰⁻⁶⁰⁶ (the catalytic region), HsExo9 with 2-fold excess of RRP6¹⁸⁰⁻⁸⁰⁴ (the catalytic region with a portion of the carboxy terminus), and RRP6¹⁸⁰⁻⁶⁰⁶. Gel filtration analysis on RRP6¹⁸⁰⁻⁸⁰⁴ in isolation was not possible as this construct is unstable in the absence of the SMT3 tag. Mixtures (0.15 ml at ~2 mg/ml) were dialyzed overnight at 4°C against reconstitution buffer (50 mM NaCl, 20 mM Tris-HCl [pH 8.0], and 10 mM DTT). Mixtures were applied to a *Superose 6* 10/300 GL column (G.E. Health Sciences) equilibrated with reconstitution buffer with the resulting UV traces (upper panels) and analyzed fractions (lower panel). **d**, Interaction studies of budding yeast Rrp6 with ScExo10^{Rrp44}. Rrp6 (residues 128-733) and Rrp6^{CTD}, but not Rrp6^{CAT}, interact with Exo10^{Rrp44}. Rrp6 (residues 128-733), Rrp6^{CAT} (residues 128-518), the Rrp6^{CTD} (residues 518-733), or both were incubated with Exo10^{Rrp44} for one hour on ice before analysis by size exclusion chromatography (*Superose 6*) in reconstitution buffer. Rrp6 and Rrp6^{CAT} were added in 1.5-fold molar excess to Exo10^{Rrp44}, while Rrp6^{CTD} was added in 3-fold molar excess. For panel **c** and **d**, fractions were analyzed by SDS-PAGE and protein detected with Sypro Ruby.



Extended Data Figure 4. RNA interactions within the Rrp6 active site

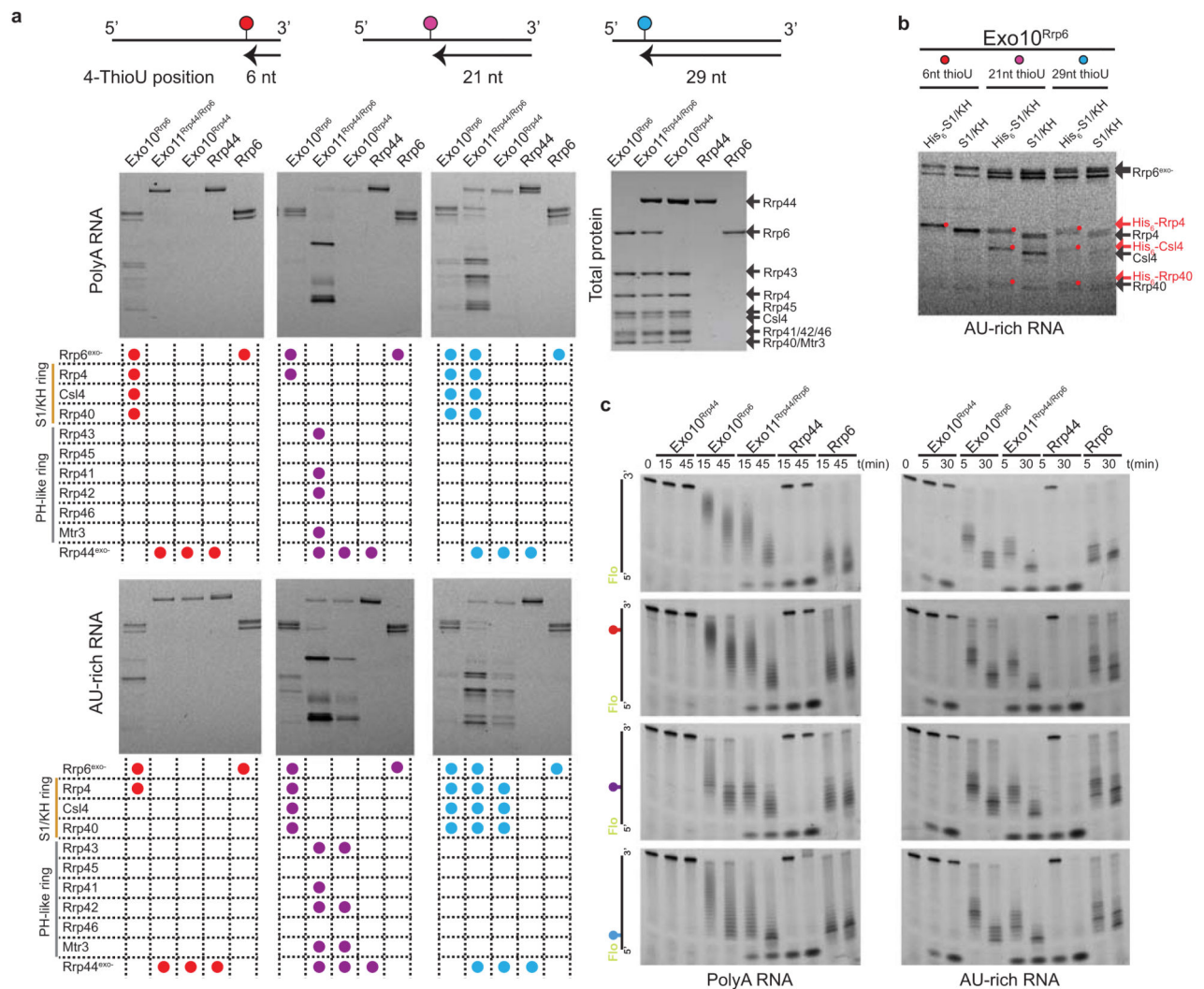
a,b, View of RNA nucleotides 24 to 20 from the 3' end in the active site of *S. cerevisiae* Rrp6 from the structure of Exo10^{Rrp6} (**a**) and the same RNA modeled into the active site of *H. Sapiens* Rrp6 (PDB: 3SAF) based on superposing the respective EXO domains (**b**). Residues involved in RNA contacts in the structure of yeast Exo10^{Rrp6} are shown as teal sticks in (**a**); the corresponding residues in human are shown in (**b**). In both structures, a catalytic aspartate is mutated to asparagine and labeled in red (N238 in yeast; N313 in human). RNA is shown as orange sticks, and magnesium ions are represented as green

spheres. **c**, Structure of the yeast Rrp6 catalytic domain in complex with zinc (Zn; blue sphere) and manganese (Mn; yellow sphere) in complex with AMP (PDB 2HBL). Active site residues are depicted as in **a,b**. The conserved Tyr361 side chain was mutated to alanine in this structure (labeled Y361A in red). **d**, Model of the Rrp6 active site constructed by superposing the 3' nucleotide in the Exo10^{Rrp6} polyA structure to the AMP ligand from PDB 2HBL. Amino acid side chains are labeled and putative interactions between metal ligands, active site residues and RNA are depicted by dashed lines.



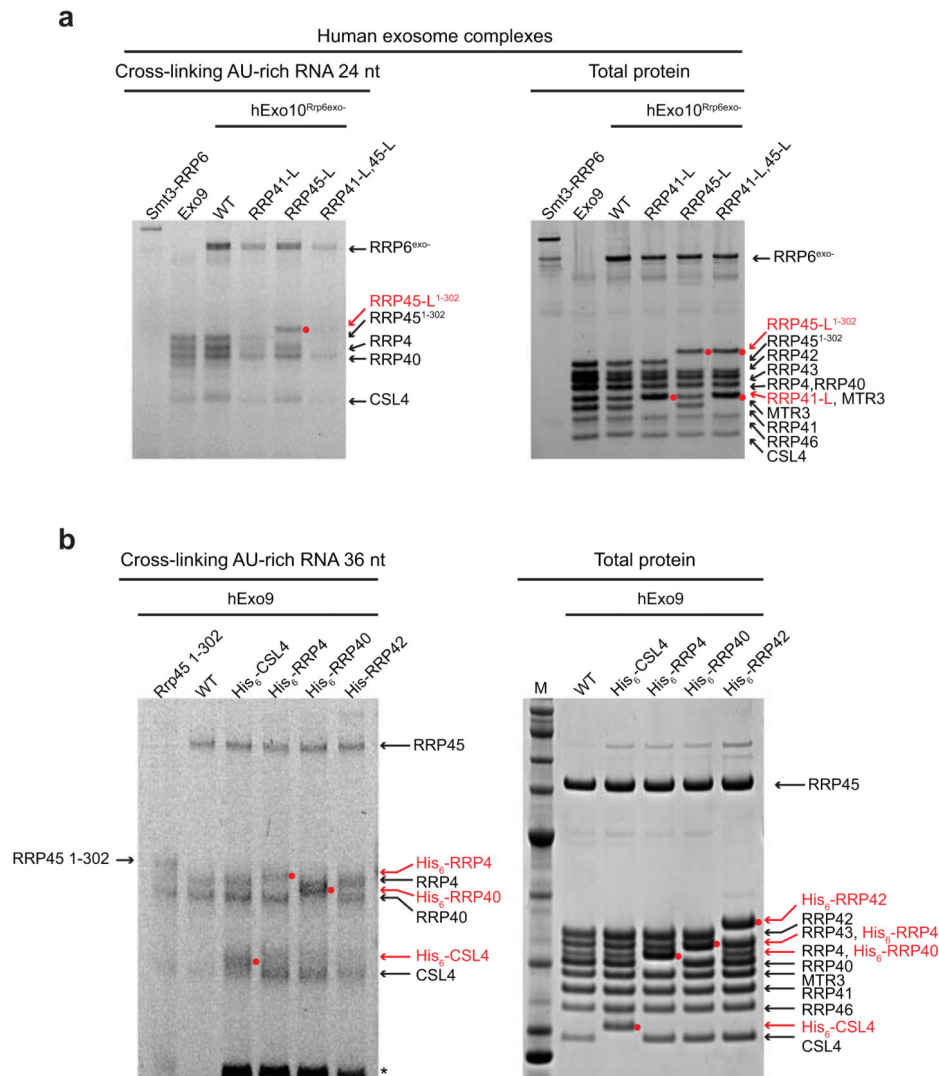
Extended Data Figure 5. Central channel mutations and their impact on Rrp6 degradation of AU-rich RNA

a, 1 nM of reconstituted wild-type and mutant yeast Exo10^{Rrp6} and Exo11^{Rrp44/Rrp6} were incubated with 10 nM 49-nt 5' fluorescein AU-rich RNA at 30°C. Reaction intermediates were analyzed after indicated time points by denaturing PAGE and imaged with a fluorimager. Rrp6 activity decreases in the presence of the Rrp40 K107E/K108E/R110D and Csl4 R145A/R150A/R202D mutations in both Exo10^{Rrp6} and Exo11^{Rrp44/Rrp6} while is mostly unaffected by the channel occlusion in the Rrp45^{Large} exosome. Rrp44 exoribonuclease activity is attenuated by mutations in both the S1/KH and PH-like ring in Exo11^{Rrp44/Rrp6}. **b**, Exoribonuclease activities of wild-type and mutant forms of human Exo10^{RRP6} were performed in a 100-μL reaction mixture with 5 nM 49-nt 5' fluorescein AU-rich RNA and 50 nM protein at 37°C. Reaction intermediates were analyzed after indicated time points as described in **(a)**. Mutation studies reveal that the human RRP6 exoribonucleolytic activity decreases in the presence of a channel occlusion near the S1/KH ring (RRP41^{Large}), while a channel occlusion deeper in the central channel (RRP45^{Large}) of HsExo10^{RRP6} has little impact on activity.



Extended Data Figure 6. UV crosslinking with 36-nt 4-thioU RNAs for yeast exosome complexes
a, Raw data used to generate Fig. 4a in main text for polyA and AU-rich RNA. See Methods for crosslinking details. Presence or absence of crosslinks to given subunits is summarized in the table and denoted by color-coded dots, with red corresponding to 4-thioU 6 nt from the 3' end; purple for 4-thioU 21 nt from the 3' end; and blue for 4-thioU 29 nt from the 3' end. **b**, RNA-protein adducts observed in UV crosslinking to Exo10^{Rrp6} correspond to the S1/KH cap proteins. Exo10^{Rrp6} bearing N-terminal hexahistone tags on the S1/KH cap proteins Rrp4, Rrp40, and Csl4 (ref. 13) or tag free variants were crosslinked to the three 4-thioU RNAs described above, and resolved by SDS-PAGE and imaged with a fluorimager. In comparing the His₆-tagged and tagless complexes, mobility shifts are apparent in crosslinks corresponding to the S1/KH cap proteins. **c**, 4-thioU RNAs are bona fide decay substrates for exosome complexes and catalytic subunits, Rrp6 and Rrp44. Conditions using substrate excess (10 nM RNA, 1 nM enzyme), 4-thioU polyA RNAs (left) and AU-rich RNAs (right) are comparably degraded to corresponding 36-nt RNAs lacking 4-thioU. Reactions were performed at 30°C and stopped after indicated time points and resolved by

denaturing PAGE and imaged with a fluoroimager. 4-thioU RNAs are color-coded as in (a) and (b).

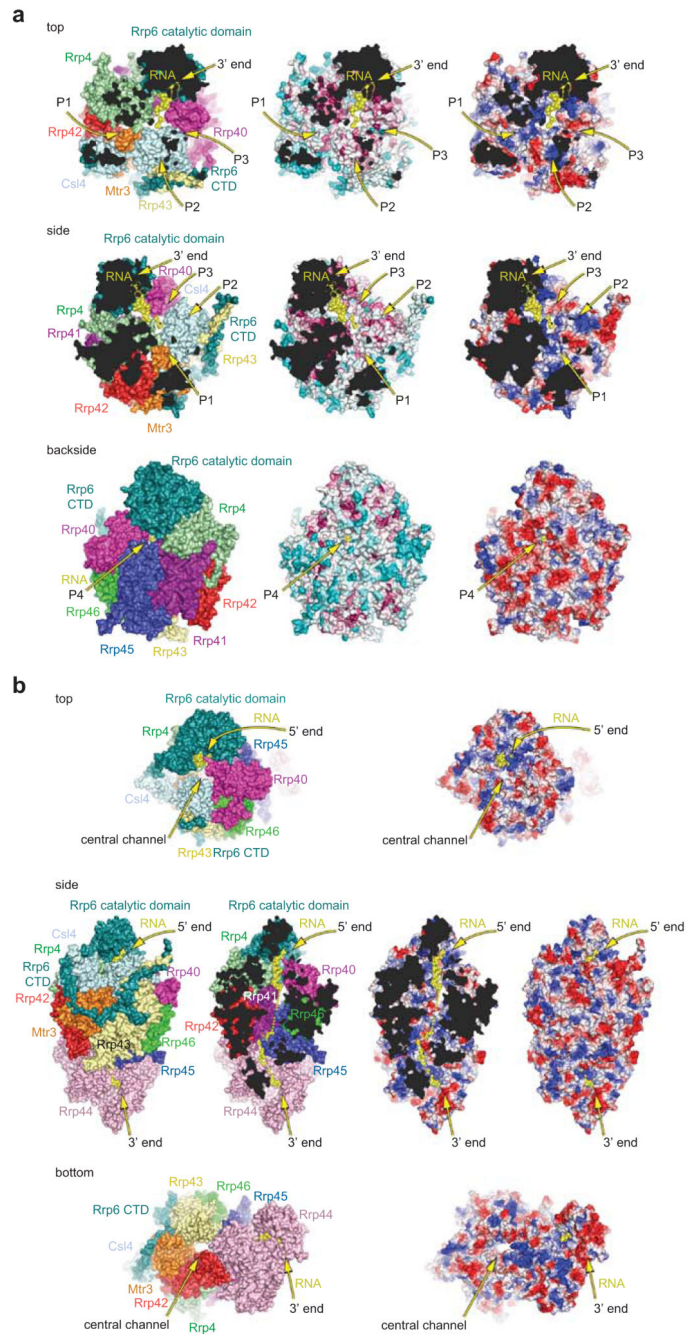


Extended Data Figure 7. UV crosslinking to AU-rich RNA to components of human Exo10^{RRP6}

a, UV crosslinking of RNA to HsExo10^{RRP6} with open and occluded channels. Human RRP6 (Smt3-RRP6¹⁸⁰⁻⁸⁰⁴), HsExo9, and wild type as well as three different pore occlusion mutants of HsExo10^{RRP6} were incubated and UV crosslinked to a 24-nt fluorescein-labeled AU-rich RNA substrate. Samples were analyzed for the presence of RNA-protein crosslinks (left panel) or integrity of protein via Sypro staining (right panel, Bio-Rad). Mutant forms of HsExo10^{RRP6} with insertion loops in RRP41-L and RRP45-L are labeled with red asterisks.

b, Identification of UV-induced crosslinks. Identification of RNA-protein crosslinks between RNA to cap components of the human Exo9 core (RRP4, RRP40, and CSL4) was facilitated by reconstituting His₆-tagged versions of these subunits into HsExo9 complexes, performing crosslinking assays with a 36-nt fluorescein-labeled AU-rich RNA, and detecting slower mobility for the His₆-tagged crosslinked adducts. Similarly, identification

of RRP45-RNA adducts was based on changes in adduct mobility when using isoforms with wild-type carboxy termini (RRP45¹⁻⁴⁵⁶) and the long insertions (RRP45-L¹⁻³⁰²). RRP6-RNA adducts were determined by crosslinking of an Smt3-RRP6¹⁸⁰⁻⁸⁰⁴ fusion and comparison of 9- and 10-subunit exosomes (in **a**).



Extended Data Figure 8. RNA paths to Rrp6 and Rrp44

Structure of an Exo11^{Rrp44/Rrp6} model derived by superposing the PH-like ring subunits in Exo10^{Rrp6} and Exo10^{Rrp44+6Cterm} (PDB 4IFD). **a**, RNA paths to the Rrp6 active site. RNA

(yellow) shown as derived from the Exo10^{Rrp6} polyA complex. Three views are provided from the top, side and backside of the complex with subunits colored as in Fig. 1 in the main text (left), surfaces colored according to conservation as calculated by ConSurf³⁵ (middle) from red (highly conserved) to blue (variable), and surfaces colored according to electrostatic potential as calculated by Pymol²⁷ (right). Subunits that are visible are labeled in the left panel as is the RNA. Three putative paths that traverse the S1/KH ring into the Rrp6 active site are denoted by yellow arrows and labels (P1, P2, P3) in the top and side views and a fourth path (P4) under the Rrp6 catalytic domain is denoted in the backside view. Based on conservation, electrostatics and crosslinking we deem P1 or P2 as the most likely path of ingress for the incoming RNA substrate. The top and side views utilize cutaway surfaces to illustrate the putative paths and surface properties. **b**, RNA paths to the Rrp44 active site. Subunits and surfaces depicted as in (a) with the exception that the RNA is now derived from the structure of Exo10^{Rrp44+6Cterm} (PDB 4IFD) with the RNA edited to remove the stem loop after nucleotide 36. Top, side and bottom views are depicted with the RNA path indicated by a yellow arrow with 5' and 3' ends labeled. The position of the central channel is also labeled and indicated by a yellow arrow in the top and bottom views. The top and side views show that RNA could pass by Rrp6 to penetrate the S1/KH and PH-like ring central channel.

Extended Data Table 1
Data collection and refinement statistics

Exo10 ^{Rrp6} /polyA	
Data collection	
Space group	P2 ₁ 2 ₁ 2
Cell dimensions	
<i>a</i> , <i>b</i> , <i>c</i> (Å)	193.3, 200.1, 97.4
<i>α</i> , <i>β</i> , <i>γ</i> (°)	90.0, 90.0, 90.0
Resolution (Å)	50-3.3 (3.42-3.3) *
<i>R</i> _{merge}	6.1 (65.3)
<i>I</i> / <i>σI</i>	15.5 (1.4)
Completeness (%)	97.7 (96.9)
Redundancy	5.2 (3.1)
Refinement	
Resolution (Å)	50-3.3
No. reflections	56332
<i>R</i> _{work} / <i>R</i> _{free}	22.7/26.5
No. atoms	21595
Protein	21450
Ligand/ion	145
Water	0
B-factors	
Protein	125
Ligand/ion	156

Exo10 ^{Rrp6} /polyA	
Water	n/a
R.m.s deviations	
Bond lengths (Å)	0.002
Bond angles (°)	0.47

One crystal was used.

* Highest resolution shell is shown in parenthesis.

Extended Data Table 2 Amino acid and RNA residues

Subunit	Residues	Non-native N-terminal residues	Residues observed in structure
Rrp45	305	None	3-206, 213-301
Rrp41	246	GDPH	4-242
Rrp43	394	None	6-101,121-180, 184-191, 209-249, 271-309, 327-394
Rrp46	223	GS	1-221
Rrp42	265	GDPH	2-161, 170-264
Mtr3	250	None	5-20, 43-147, 163-248
Rrp40	240	GDPH	2-45, 54-235
Rrp4	359	GDPH	5-14, 50-145, 158-249, 277-357
Csl4	292	GDPH	5-70, 105-113, 128-162, 186-291
Rrp6 (128-685)	558	SL	128-516, 525-628
RNA	24	n.a.	19-24

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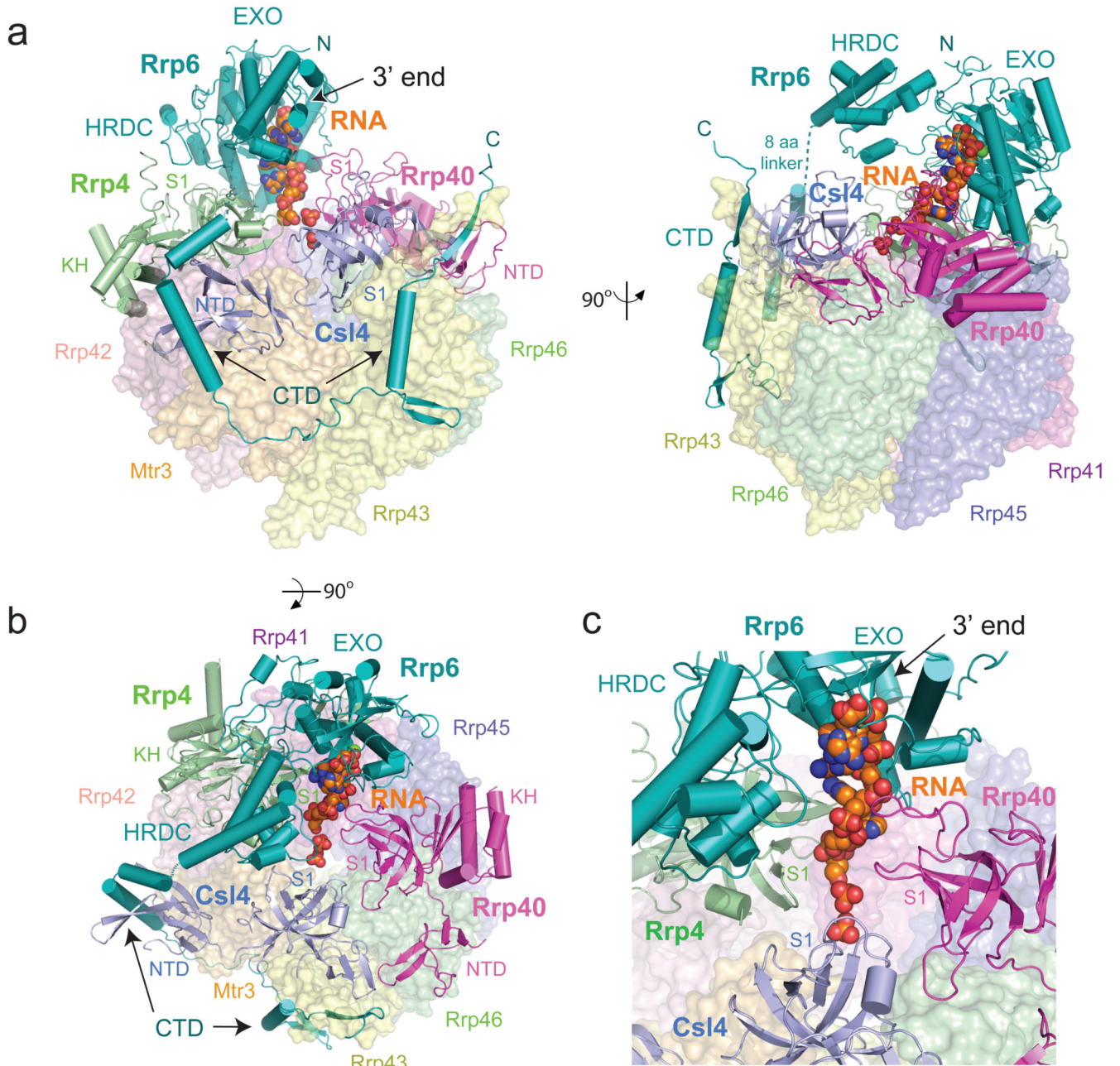


Figure 1. Overall structure of a RNA bound Rrp6 exosome
a,b,c, Cartoon and surface representation of *S. cerevisiae* Exo10^{Rrp6} bound to polyA₂₄ RNA. Views from the (a) sides, (b) top, and (c) close-up of RNA engaged by the Rrp6 EXO domain and S1 domains of Rrp40, Rrp4, and Csl4. RNase PH-like ring as transparent surfaces, S1/KH ring and Rrp6 as cartoons, RNA as spheres.

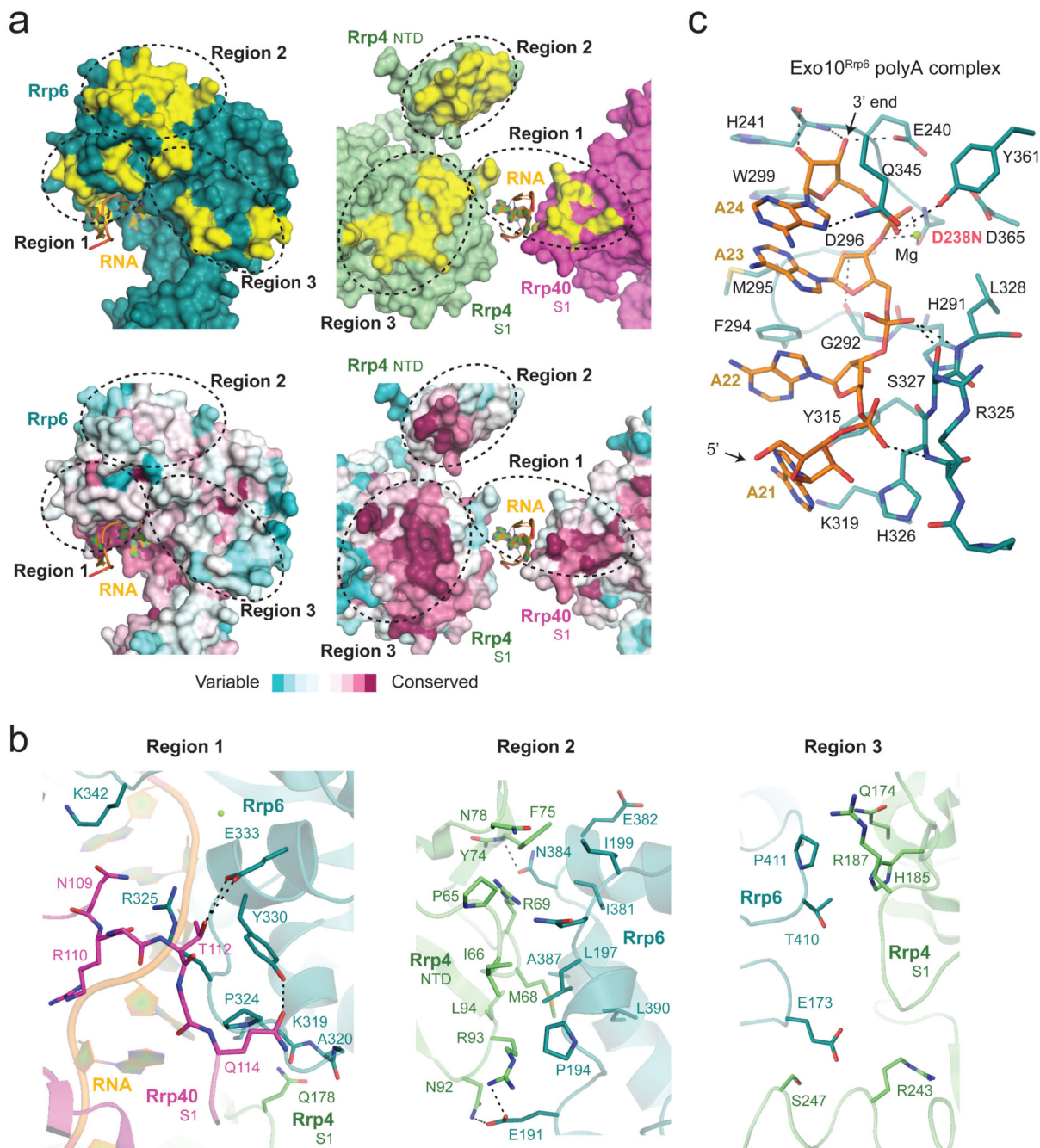


Figure 2. Rrp6 contacts the S1/KH ring and RNA

a,b, Rrp6 EXO domain interface with Rrp4 and Rrp40 as yellow buried surfaces (**a**) and in atomic detail (**b**). **a,** Rrp6 (teal; left), Rrp4 (pale green; right) and Rrp40 (magenta; right) in open book representation (160°). Below, surfaces colored by sequence conservation from red (conserved) to blue (variable). **b,** Contacts between Rrp6, Rrp4 and Rrp40 in stick representation colored as in (**a**). **c,** Contacts between RNA (orange), magnesium (green) and Rrp6 active site residues (teal) in Exo10^{Rrp6}-RNA. A catalytic aspartate, D238, is mutated (D238N; red). Potential hydrogen bonds depicted as dashed lines.

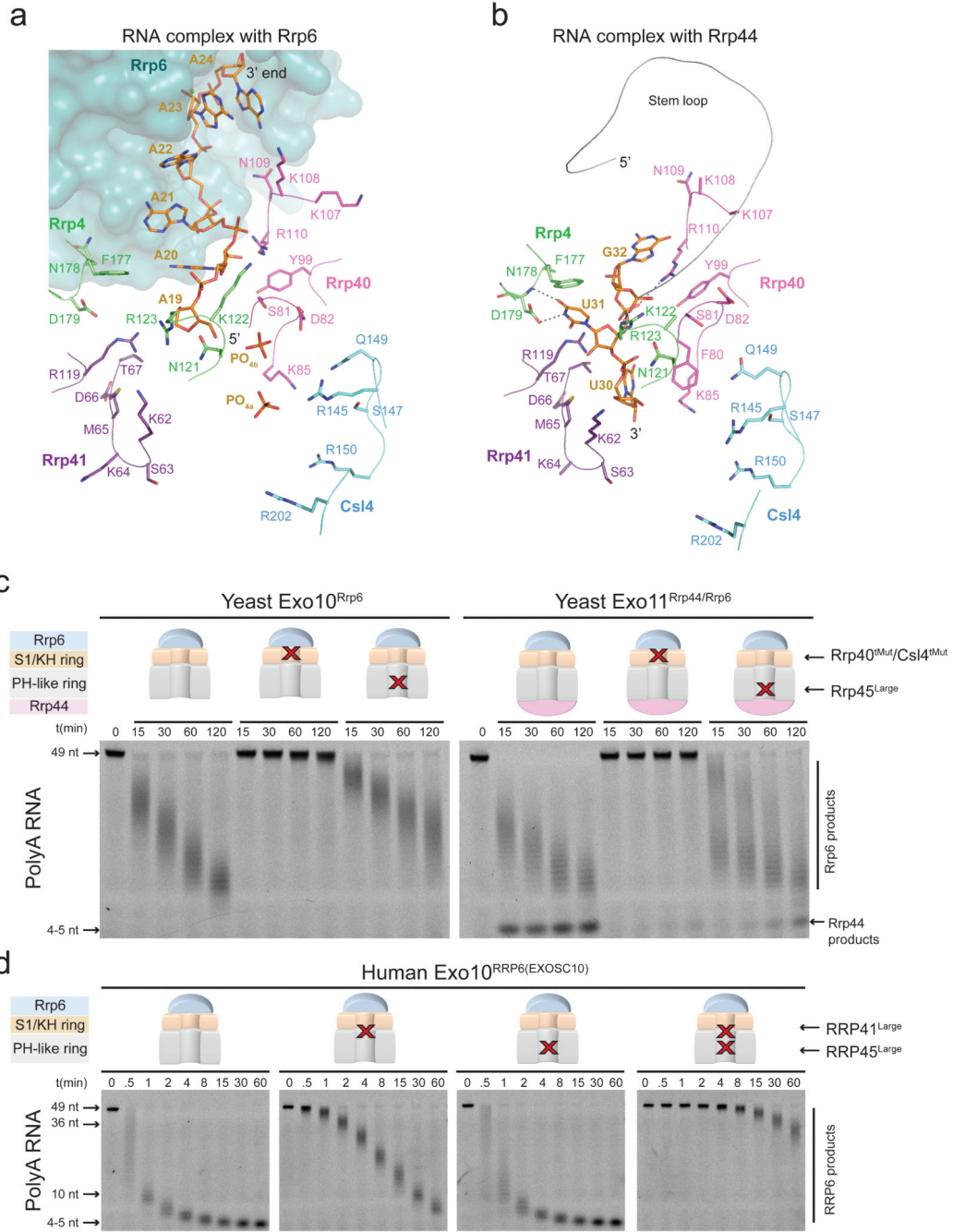


Figure 3. Rrp6 activities are dependent on the S1/KH ring
a,b, Partially overlapping paths for RNA in complex with **(a)** Exo10^{Rrp6} and **(b)** Exo10^{Rrp44+Rrp6Cterm} within the S1/KH ring. RNA and S1/KH side chains depicted as sticks, Rrp6 **(a)** as a surface, the stem loop **(b)** in Exo10^{Rrp44+Rrp6Cterm} as a ribbon. Color-coded labels adjacent to subunits. **c**, RNA decay activities for yeast Exo10^{Rrp6} and Exo11^{Rrp44/Rrp6} with mutations in the S1/KH ring or a channel occluding insertion lower in the PH-like ring (Rrp45^{Large}). **d**, RNA decay activities for human Exo10^{RRP6} with insertions at the top (RRP41^{Large}) or middle of the PH-like ring (RRP45^{Large}). RNA decay assays were

conducted at least three times for each of the samples with typical results depicted in (c) and (d).

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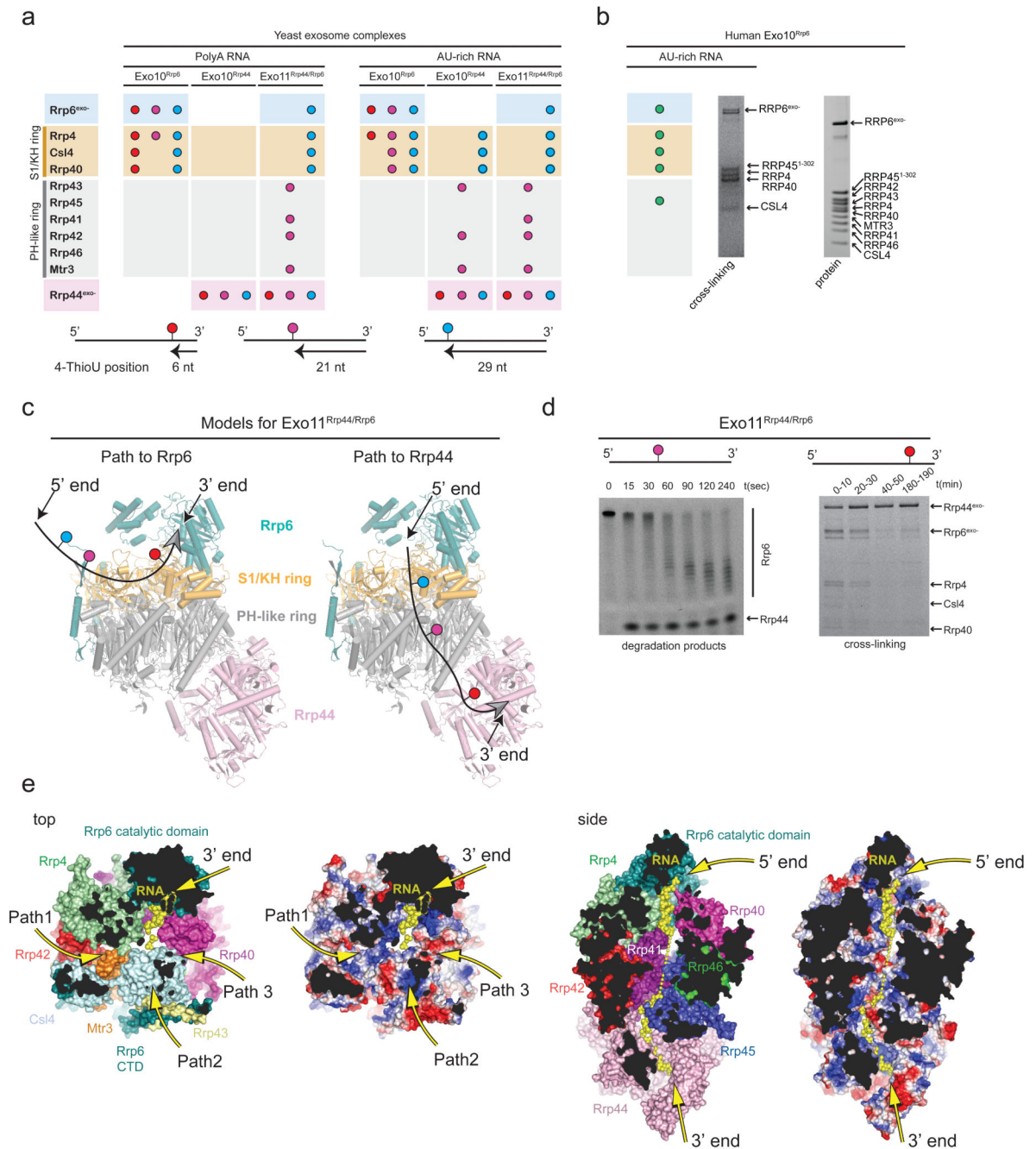


Figure 4. RNA paths to Rrp6 and Rrp44

a, RNA crosslinking pattern schematic for yeast exosomes (Extended Data Fig. 6a). Subunits labeled, background colored with Rrp6 (blue), S1/KH ring (orange), PH-like ring (grey) and Rrp44 (pink). 4-thioU positions indicated by colored circles at 6, 21 or 29 nt from the 3' end. **b**, RNA crosslinking pattern schematic for human Exo10^{RRP6} as in (a). Crosslinks (left); proteins (right) (Extended Data Fig. 7). **c**, Exo11^{Rrp44/Rrp6} cartoon. Subunits and rings colored and labeled. RNA paths depicted by arrows, arrowhead at 3' end with 4-thioU positions indicated as in (a). **d**, RNA degradation at 5-fold enzyme excess with

Rrp6 and Rrp44 products (left) and time course for crosslinking to 4-thioU RNA 6 nt from 3' end (right). **e**, Exo11^{Rrp44/Rrp6} model indicating RNA paths to Rrp6 (left) or Rrp44 (right). Subunits colored as in Fig. 1 next to surfaces colored according to electrostatic potential²⁷. Visible subunits and RNA (yellow) labeled in left panels. Three putative paths past the S1/KH ring to Rrp6 denoted by yellow arrows and labels (P1, P2, P3) (Extended Data Fig. 8a). Side view shows an RNA path to Rrp44 (yellow arrow; 5' and 3' ends labeled) indicating that RNA could pass by Rrp6 (Extended Data Fig. 8b). RNA from PDB 4IFD was edited to remove the stem loop after nt 36. Assays in **(a,b,d)** conducted at least three times with typical results depicted.

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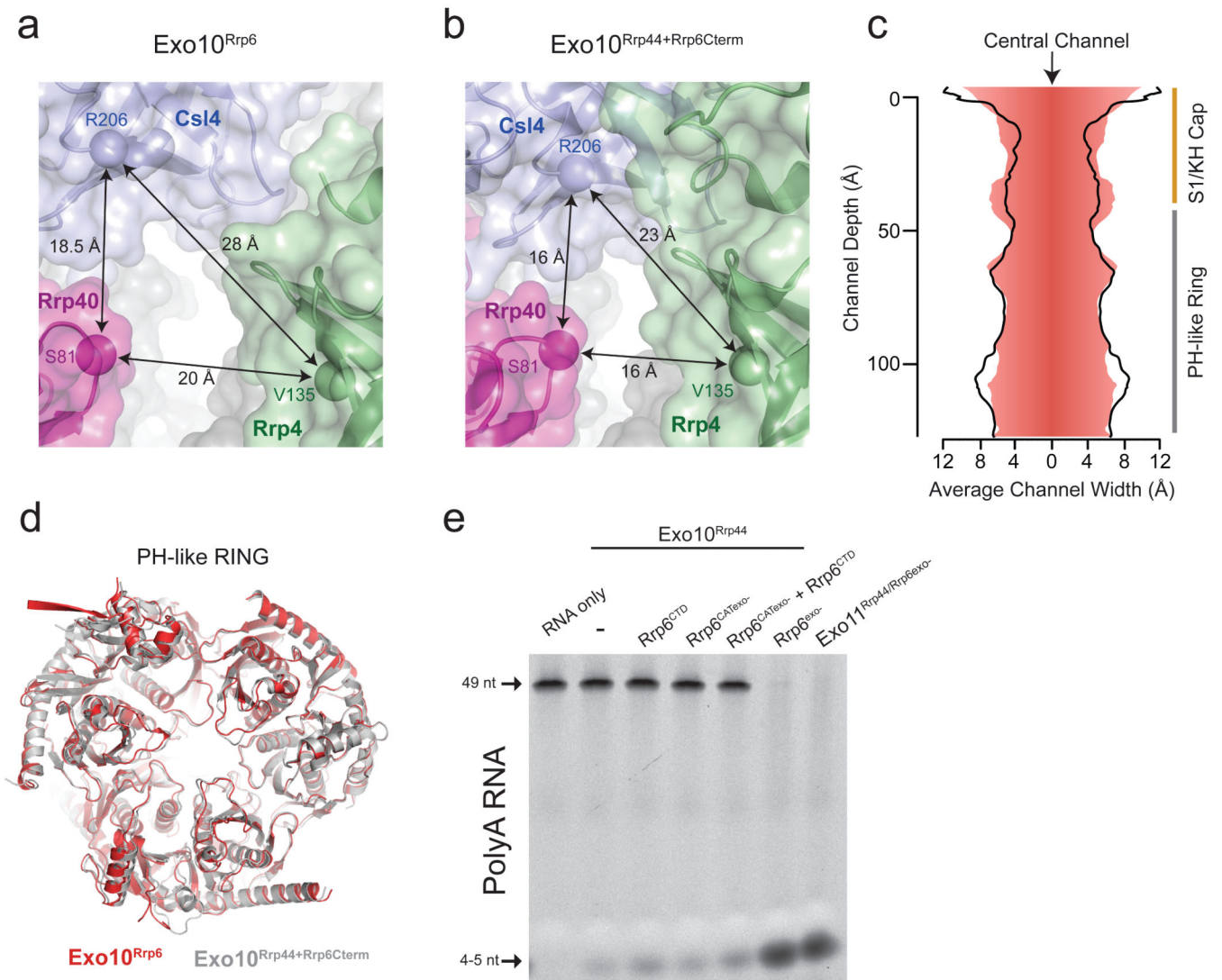


Figure 5. S1/KH ring widening in Exo10Rrp6

a,b, Close-up of the central channel looking down the S1/KH ring. Rrp4 (green), Rrp40 (magenta) and Csl4 (light blue) in cartoon with transparent surfaces from **(a)** Exo10Rrp6 and **(b)** Exo10Rrp44+Rrp6Cterm. Distances indicated between respective labeled amino acids. **c,** Channel dimensions in Exo10Rrp6 (red) and Exo10Rrp44+6Cterm (black) calculated using HOLE²⁸. **d,** Superposition of PH-like rings for Exo10Rrp6 (red) and Exo10Rrp44+6Cterm (grey). **e,** Rrp6^{exo-} (128-733), and not Rrp6^{CTD} (518-733) or Rrp6^{CAT} (128-518), is required to stimulate Rrp44 activity. RNA decay assays conducted at least three times with typical results depicted.