The Arp8 and Arp4 module acts as a DNA sensor controlling INO80 chromatin remodeling

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Nuclear actin and actin-related proteins (Arps) are key components of chromatin remodeling and modifying complexes. Although Arps are essential for the functions of chromatin remodelers, their specific roles and mechanisms are unclear. Here we define the nucleosome binding interfaces and functions of the evolutionarily conserved Arps in the yeast INO80 chromatin remodeling complex. We show that the N-terminus of Arp8, C-terminus of Arp4 and the HSA domain of Ino80 bind extranucleosomal DNA 37–51 base pairs from the edge of nucleosomes and function as a DNA-length sensor that regulates nucleosome sliding by INO80. Disruption of Arp8 and Arp4 binding to DNA uncouples ATP hydrolysis from nucleosome mobilization by disengaging Arp5 from the acidic patch on histone H2A-H2B and the Ino80-ATPase domain from the Super-helical Location (SHL) -6 of nucleosomes. Our data suggest a functional interplay between INO80’s Arp8-Arp4-actin and Arp5 modules in sensing the DNA length separating nucleosomes and regulating nucleosome positioning.

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The ATP-dependent chromatin remodeling complex INO80 has important roles in transcription, heterochromatin maintenance, DNA replication, repair, and recombination. These functions reflect the ability of INO80 to shape promoter nucleosome landscape by exchanging histone H2A.Z for H2A in nucleosomes and changing nucleosome positions. Disruption of INO80 function is associated with cancers as well as coronary and other vascular diseases. Recently, the structures of fungal and human INO80 complexes bound to nucleosomes were solved using high-resolution cryo-electron microscopy (cryo-EM). However, these structures did not define the binding interfaces of the evolutionarily conserved nuclein protein and actin-related proteins (Arps) of INO80 or provide insights into their functional properties. Arp4 and actin are shared components of three different chromatin remodeling complexes that mechanically move or alter the composition of nucleosomes in an ATP-dependent manner (INO80 and SWR1) or post-translationally modify histones (NuA4). A similar and less conserved module comprised of Arp7 and 9 is also present in the yeast SWI/SNF and RSC chromatin remodelers. In the INO80 complex, the actin-Arp4 module has an additional Arp subunit, i.e., Arp8. When Arp8 is deleted actin and Arp4 are also lost from the complex, thereby causing the complex to lose its ability to bind DNA. Arp8 is further shown to be associated with DNA by ChIP-exo in which Arp8 is broadly crosslinked with formaldehyde to linker and nucleosomal DNA. The Arp8-Arp4-actin module binds to the helicase/SANT-associated (HSK) domain of the Ino80 subunit. The crystal structures of the actin-Arp4-HSA(Swr1) trimer and the actin-Arp4-HSA(Ino80)-Arp8 tetramer have been solved, and mutational analysis have shown that the HSA domain binds DNA near the barbed end of the actin-fold. Since Nhp10 and Taf14 have similar electrophoretic mobilities, we interpreted the peptide mapping as showing that two distinct sites are crosslinked, because cleavage at residue 541 appears to be blocked and the other cleavage products indicate crosslinking between residues 915 and 1080. The Ino80 HSA domain could be expected to bind the same region as Arp4 and Arp8 given the HSA domain binds DNA in vitro to bind with histones and actin. Here we describe that in addition to the HSA domain Arp8 and Arp4 bind extranucleosomal/linker DNA at a discrete distance from the entry site of nucleosomes and function as a DNA-length sensor to allosterically regulate INO80-mediated nucleosome spacing. In the crystal structure of the actin-Arp4-HSA-Arp8 tetramer (Knoll et al. in preparation), Arp8 is missing its N-terminus from residues 1–254, and our data shows this missing region associates with DNA and is required for the Arp8-Arp4-actin module to linker DNA. Binding of the INO80 Arp8 module to linker DNA is required to couple ATP hydrolysis to nucleosome repositioning and for the proper docking of the Ino80 catalytic subunit and Arp5 to nucleosomes.

**Results**

The Arp8 and Nhp10 modules and Ino80 HSA bind linker DNA. We probed INO80 interactions at eight different positions in extranucleosomal DNA by DNA crosslinking. The Ino80 catalytic subunit along with subunits of the Arp8 module bind to extranucleosomal DNA 110–124 nucleotides (nt) from the dyad axis or 37–51 base pairs (bp) from the edge of nucleosome core particles, and the yeast specific Nhp10 module binds to a wider region on extranucleosomal DNA (nt −92 to −124), 18–51 base pairs (bp) from the edge of nucleosome core particles (Fig. 1a, b). The yeast Arp8 module binds the HSA domain of Ino80 and besides the three conserved subunits contains the yeast specific Ies4 and Taf14 subunits. Of the 5-subunit Arp8 module, we find only the Arp4 and Arp8 subunits directly associating with extranucleosomal DNA as seen by site-directed DNA crosslinking. Immunoprecipitation confirms that Arp4 is crosslinked and not Ies2, which has a similar electrophoretic mobility (Supplementary Fig. 1a). Consistent with our results ChIP-exo shows Arp8, Nhp10, and Ies5 interacting with the linker DNA associated with promoter-proximal +1 nucleosomes, but rather than to a broad stretch of DNA we show that Arp8 is bound to a 14 base pair region of extranucleosomal DNA spanning from 110 to 124 bp from the dyad axis (nt −110 to −124). The Ino80 catalytic subunit associates with this same region and Arp4 to a narrower part spanning from 110 and 111 bp from the dyad axis. Chemical cleavage of photo-crosslinked Arp8 and Arp4 at methionines and cysteines reveal that the N-terminus of Arp8 (residues 1–197) associates with extranucleosomal DNA at nt −124, while the C-terminus of Arp4 (residues 425–489) is associated with extranucleosomal DNA spanning from 110 to 124 bp from the dyad axis of nucleosomes. Deletion of the entire Arp8 subunit causes a loss of the ability of INO80 to shape promoter nucleosomes and changing nucleosome positions.

We find the Nhp10 and Ies5 subunits bind to nearly the same stretch of extranucleosomal DNA as the Arp8 module in two distinct regions, at nt −110 to −124 (Nhp10) and −92 (Ies5) (Fig. 1a, b). The Nhp10 module is comprised of Nhp10, Ies1, Ies3, and Ies5, and this yeast specific subcomplex and its metazoan-specific version bind to the extreme N-terminus of Ino80. The metazoan subcomplex is not required for INO80 remodeling, whereas the yeast Nhp10 module has a role in sensing the DNA length between nucleosomes.

Since Nhp10 and Taf14 have similar electrophoretic mobilities, we confirmed the identity of the crosslinked protein by observing the shift in the mobility of the crosslinked species when three copies of an epitope tag (HA, hemagglutinin) are attached to the C-terminus of Nhp10 (Supplementary Fig. 1b, c). Either the Ies4 or Ies6 subunit also crosslink at nt −92 and was not differentiated at this time (Supplementary Fig. 1a).
shown previously to reduce the association of actin, Arp4 and Arp8 with the rest of the INO80 complex, similar to \textit{arp8} null\cite{10}. We also find the Arp8ΔN-INO80 strain to behave similar to those in which the spacer (or insert) region from amino acids 1018 to 1299 between the two ATPase lobes is deleted (ΔSpacer), causing the loss of the Arp5, Ies6, Ies2, and Rvb1/2 subunits and disrupting the in vivo activity of INO80\cite{25}. Our data suggest that the biochemical activity of the Arp8ΔN-INO80 complex is distinct from that in which Arp8, Arp4 and actin are absent\cite{11,19}, as the nucleosome binding and ATPase activities of Arp8ΔN-INO80 are not significantly reduced compared to wild-type INO80; however, nucleosome mobilization is reduced 6-fold (Fig. 2c, d, and Supplementary Fig. 3a, b). This uncoupling of ATPase activity from nucleosome mobilization indicates that the Arp8ΔN-INO80 complex retains particular contacts with nucleosomes critical to stimulate its ATPase activity and has specifically lost other key interactions required to move DNA through nucleosomes.

We examined the defects in nucleosome mobilization further by mapping changes in histone-DNA contacts as DNA tracks across the octamer surface\cite{17,26}. We find that the Arp8ΔN
complex moves the first 11 bp of DNA at nearly the same rate as wild-type INO80, indicating that it can readily complete this early step in remodeling but cannot efficiently move DNA farther (Fig. 2e, f, and Supplementary Fig. 3c). Besides moving DNA along the surface of the histone octamer, INO80 can persistently displace DNA from the surface of the H2A-H2B dimer in an ATP-dependent manner\(^\text{17}\). We track DNA displacement at the H2A-H2B interface by examining the sum of all INO80-remodeled products including new and starting positions and whether that sum decreases over time (Fig. 2g and Supplementary Fig. 3c lanes 1–9). The Arp8ΔN complex displaces DNA from the H2A-H2B dimer two times less efficiently than wild-type INO80 despite an increased rate of ATP hydrolysis (Fig. 2h, Supplementary Fig. 3c lanes 10–18 versus lanes 1–9, and Fig. 2d).

**Arp8-DNA interaction regulates key INO80-DNA contacts.** Next, we explored why loss of the N-terminus of Arp8 uncouples nucleosome movement and DNA displacement from ATP hydrolysis by re-examining the INO80-nucleosome binding interfaces. Loss of the N-terminus of Arp8 does not reduce the affinity of INO80 complex for nucleosomes, likely reflecting the multiple interactions that are involved in INO80 binding (Fig. 3a and Supplementary Fig. 4a). DNA crosslinking confirmed that the N-terminus of Arp8 is required for stable binding of both Arp8 and Arp4 to extranucleosomal DNA (nts −124 to −75/−73) (Fig. 3b lanes 2–11 comparing odd and even lanes, and Supplementary Fig. 4b–d). We also show that attachment of a HA-tag at the C-terminus of Arp4 interferes with Arp4 and Arp8 binding at nts −110/−111, consistent with the C-terminus of Arp4 binding DNA (Supplementary Fig. 1c lanes 6 and 7). Binding of the Nhp10 subcomplex to extranucleosomal DNA is however not blocked and its crosslinking increases when the N-terminus of Arp8 is absent or the C-terminus of Arp4 is tagged (Fig. 3b lanes 2–7 plus the lower panel, Supplementary Figs. 1c, lanes 6 and 7, and 4e).

DNA crosslinking experiments revealed that stable binding of the ATPase domain of Ino80\(^\text{17}\) and Ies2 to DNA at the superhelical location (SHL)-6 in nucleosomes (nt −58) is reduced by deletion of the N-terminus of Arp8 (Fig. 3b lanes 14–17, Fig. 3c, and Supplementary Fig. 4d). Previously, Ies2 was shown to bind to the ATPase domain of Ino80, consistent with our DNA crosslinking data for Ies2\(^\text{19}\). Furthermore, Ino80 binding shifts from SHL-6/−5 to extranucleosomal DNA up to 92 nucleotides from the dyad axis (Fig. 3b lanes 8–13, compare odd and even lanes, plus the lower panel, and Fig. 3c). There are two possible reasons for these changes interfering with INO80 moving nucleosomes. First, displacement of DNA from a part of the histone octamer surface caused by the ATPase domain binding at SHL-6 is likely absent when the N-terminus of Arp8 is deleted, while this displacement may prime nucleosome movement by the wild-type complex\(^\text{8,9}\). Second, since Ino80 shifts to an external position outside of the core nucleosomes when Arp8 is truncated, DNA translocation near the H2A-H2B dimer required by INO80 to mobilize nucleosomes will also be missing\(^\text{17}\).

**Arp5 binding to H2A-H2B requires Arp8-DNA interaction.** To further investigate changes in INO80-nucleosome interactions we probed interactions with the histone octamer surface through site-specific histone crosslinking\(^\text{27}\). We find that the Arp5 subunit contacts the H2A-H2B dimer near its acidic pocket, consistent with recent cryoEM structures\(^\text{28–30}\), and this contact is diminished when the N-terminus of Arp8 is absent (Fig. 4a, b, d, and Supplementary Fig. 5). Since Arp5 and Ies1 migrate similarly, we confirmed Arp5 photocrosslinking by comparing the electrophoretic mobility of the crosslinked species with and without HA-tagged Arp5 (Fig. 4c). The Ino80 and Ies2 subunits also bind to the H2A-H2B dimer, but to a significantly lesser extent than Arp5. Loss of the Arp5 subunit has been shown previously to uncouple the nucleosome mobilizing ability of INO80 from its ATPase activity\(^\text{22,23,28}\), consistent with changes we observe here with the Arp8ΔN complex. Also in alignment with our observations, interactions with the acidic pocket of H2A-H2B have been shown previously to be important for positively regulating INO80 and other chromatin remodelers\(^\text{22,29–31}\).

**Discussion**

Taken together, our data show that Arp8 and Arp4 associate with extranucleosomal/linker DNA 37–51 bp from the edge of nucleosomes to properly orient INO80 on the nucleosomes for remodeling. These interactions promote proper docking of the ATPase domain of Ino80 inside of nucleosomes, as well as stable binding of Arp5 to the histone H2A-H2B acidic-pocket region (Fig. 4f). The loss of Arp5 contacts with nucleosomes and subsequent loss of coupling of ATP hydrolysis to nucleosome movement is consistent with the prior model of Arp5 being required as a histone and DNA anchor to stabilize Ino80 translocation on DNA and provide sufficient torque to displace and move DNA on the surface of the histone octamer\(^\text{8,9}\). The role of the Arp8 module binding to DNA is therefore not simply to recruit INO80 to nucleosomes as the affinity of INO80 for nucleosomes do not depend on Arp8 binding to linker DNA, but rather acting as a DNA sensor to effect allosteric changes in INO80 required for it to efficiently remodel nucleosomes. The placement of the Arp8 module onto linker DNA is in concordance with the preferred length of linker DNA required to
Fig. 2 The N-terminus of Arp8 is required for the in vivo and nucleosome mobilization activities of INO80. a Complex integrity is maintained in purified wild-type INO80 with Ino80-2FLAG (lane 1), INO80-Arp8ΔN with 2FLAG-Arp8ΔN (lane 2), and INO80 with 2FLAG-Arp8 (lane 3). b Top panel: Schematic depicting deleted regions in Arp8 (Arp8ΔN) and Ino80 (ΔTELY and ΔSpacer) mutants. Plasmids expressing wild type or mutant Arp8 (Arp8ΔN) were introduced into arp8 null yeast and compared to strains lacking INO80 function (ΔSpacer and ΔTELY). c Plots showing the rates at which wild type and Arp8ΔN INO80 move 70N0 nucleosomes (representative native electrophoretic mobility shift assay (EMSA) images shown in Supplementary Fig. 3a). d Plots based on thin-layer chromatography data showing the rates of ATP hydrolysis of wild type and Arp8ΔN INO80 in the presence of either nucleosomes or naked DNA assayed in the same conditions as in c. e, f DNA movement on the histone octamer surface of nucleosomes was measured using nucleosomes modified at residue 53 of H2B (see Supplementary Fig. 3c). Plots show the fraction of nucleosomes that moved 11 (e) and 20 (f) base pairs from the starting position versus time by wild type and Arp8ΔN INO80. g, h The amount of DNA cleaved at the starting position (initial crosslinking), the total amount of cleaved DNA that accumulates during remodeling (new positions), and sum of the intensities of the initial and remodeled positions (sum of all positions) track the retention of histone-DNA contacts versus time for wild type (g) and Arp8ΔN INO80 (h). Error bars in c-h denote s.e.m. from ≥3 experiments.
mobilize nucleosomes and the observed nucleosome spacing activity of INO80. It appears unlikely that Arp8 promotes nucleosome movement by pushing DNA into nucleosomes as DNA gaps and nicks between where the Arp8 module bind and the entry site of nucleosomes do not interfere with INO80-mediated nucleosome movement.

Sensing linker DNA length is a critical property of INO80 as this remodeler is a major factor in establishing nucleosome free or depleted regions (NFR or NDR) at promoters. Genomic studies indicate that Arp5 and Ies6 occupancies correlate with nucleosome positioning at transcription start sites and this positioning is changed when either these subunits or Ino80 are deleted. INO80 is the only ATP-dependent chromatin remodeler that alone can properly reposition nucleosomes in yeast promoter regions when genomic DNA is reconstituted in vitro. Both yeast and human INO80 exhibit DNA sensing activity that likely helps establish promoter architecture. Our data along with recent studies from the Narlikar lab indicate that two seemingly distinct mechanisms involving the Nhp10 and Arp8 modules can be used by INO80 to slide nucleosomes in a linker DNA length dependent manner, while only the Arp8-Arp4-actin module is likely evolutionarily

Fig. 3 Binding of Arp8 and Arp4 to extranucleosomal DNA mediates the proper docking of the Ino80 catalytic subunit onto nucleosomes. a Plots showing the fractions of end-positioned (70N0) nucleosomes bound by wild type and Arp8ΔN INO80 versus increasing concentrations of the enzymes (representative native electrophoretic mobility shift assay (EMSA) images shown in Supplementary Fig. 4a). Error bars denote s.e.m. from three experiments. b Site-specific interactions of wild type and Arp8ΔN INO80 with nucleosomal and linker DNA were mapped by DNA crosslinking as in Fig. 1a. ADP was added in lanes 16 and 17 (Fig. 1a and ref.17). c Crosslinking efficiencies of the Ino80 catalytic subunit in wild type and Arp8ΔN INO80 (from b) at specified nucleotide positions. The schematic below shows the changes in Ino80, Ap8, Arp4, and Nhp10 crosslinking between wild type and Arp8ΔN INO80. Circle sizes are proportional to crosslinking efficiencies at the specified DNA positions. Values are mean of three experiments, normalized to the Ino80 subunit of wild type at nt −58 without ADP (lane 14 in panel b). Error bars denote ± s.d.
Fig. 4 Arp5 interactions with the acidic pocket of the histones H2A-H2B dimer requires Arp8 and Arp4 binding to extranucleosomal DNA. a Locations of the aspartate (D) and glutamate (E) residues constituting the acidic pocket on histone H2A-H2B dimer surface are highlighted in red. The sites of the modified cysteine residues used to map histone interactions of wild type and Arp8ΔN INO80 are shown as black spheres. b–d Phosphorimages of INO80 subunits tagged by crosslinking and [125I] radiolabel-transfer. Crosslinked Arp5, Ino80, and les2 subunits are indicated on the left. c Crosslinked Arp5 was distinguished from co-migrating les1 using INO80 with C-terminally HA-tagged Arp5 (Arp5-HA). d The crosslinking patterns of wild type and Arp8ΔN INO80 are compared. e Plots showing the crosslinking efficiencies of wild type and Arp8ΔN INO80 to the H2A-H2B dimer surface. Values are mean of three experiments, normalized to the Ino80 subunit of wild type at H2B44. Error bars denote ± s.d. f The role of the Arp8 module in regulating INO80 chromatin remodeling in a linker DNA-length dependent manner is illustrated. The Arp8 and Arp5 modules are depicted along with the ATPase domain of Ino80, as a part of the INO80 complex. As nucleosomes are moved closer together, binding of the Arp8 module to linker DNA is disrupted, which in turn causes the Arp5 module to lift off from the histone dimer surface and the ATPase domains moves away from the core nucleosome. These conformational changes uncouple the ATP hydrolysis from the nucleosome mobilization activity of INO80.
conserved. The Nhp10 and Arp8 modules appear to work by distinct mechanisms since the absence of the Nhp10 module causes INO80 to move nucleosomes regardless of extra-nucleosomal DNA length23 and the lack of Arp8 module binding to extranucleosomal DNA, from our data, arrests sliding even when there is sufficient extranucleosomal DNA. It is not clear how Nhp10 negatively regulates INO80 when linker DNA is shorted, but for the Arp8 module it appears that loss of Arp8 binding to linker DNA changes critical contacts of the Ino80 and thus tunes its DNA translocation activity or makes crucial interactions with Actin-related proteins in chromatin remodelers therefore when there is sufficient extranucleosomal DNA, from our data, arrests sliding even through direct interaction with their ATPases domains12,34 which is quite distinct from how Arps regulate RSC and SWI/SNF for proper docking of the catalytic subunit and ATPase domain,12 For INO80, the Arp8 module is a DNA-binding sensor required for proper docking of the catalytic subunit and ATPase domain, which is quite distinct from how Arps regulate RSC and SWI/SNF for proper docking of the catalytic subunit and ATPase domain.

**Methods**

**Strain construction and enzyme purification.** INO80 subunits Arp5, and Nhp10 were tagged at their C-terminus using polymerase chain reaction (PCR)-mediated gene replacement in yeast. A 3HA peptide coding sequence and URA3 selection marker were amplified from a pRS416-based plasmid (kind gift from Dr. Xue Tong Shen). Purified PCR products were transformed into *Sacharomyces cerevisiae* BY4743, with the INO80 subunit bearing a 2FLAG at the C-terminus25. Putative clones were selected by growth on rich agar plates. Homologous recombination junctions were verified by PCR and expression of HA-tagged proteins was confirmed by western blotting. Yeast expressing INO80-2FLAGNhp10 and a version of wild-type INO80, with 2FLAG-tag at the N-terminus of Arp8N or full-length Arp8 respectively, were constructed by expressing 2FLAG-601-DNA or 2FLAG-Arp8 from a pRS416-based centromeric plasmid containing the native arp8 promoter, in *Sacharomyces cerevisiae* BY4743 with deletion of the endogenous arp8 (INO80-Aarp8, a kind gift from Dr. Xue Tong Shen)11. Protein expression was confirmed by western bloting. Oligonucleotide primers used for epitope tagging and Arp8 N-terminal deletion are listed in Supplementary Table 1.

**Nucleosome reconstitution.** 70N0 nucleosomes (N = nucleosome core particle with 147 bp of 601-nucleosome positioning sequence DNA, and number on either side indicate length(s) of extranucleosomal DNA) were assembled at 37°C by salt-dilution with 7–10 μg of recombinant *Xenopus laevis* histone octamers (wild type or cysteine mutant octamer, see below), 100–200 fmoi of 5’-[32P]-labeled 217 bp 601-DNA (or 601-DNA containing site-specific photo-reactive labels, see below), 5–10 μg of sheared salmon sperm DNA (for heterogeneous nucleosomes) or 601-DNA without [35S]-radiolabel as carrier (for homogeneous nucleosomes used in ATPas and nucleosome binding assays, see below). PCR oligonucleotides listed in Supplementary Table 1. DNA and histone octamer were mixed at ~1:1 w/w ratio in 2 M NaCl. Nucleosomes were dialyzed in steps to reach a final concentration of 280 mM NaCl in 25 mM Tris-HCl (pH 8). Samples were examined on 4% native polyacrylamide gels and staining with Coomassie Blue. The activities of the purified complexes were determined by EMSAs to test for nucleosome binding and remodeling, and thin-layer chromatography (TLC) for ATP hydrolysis (see below).

**Site-specific DNA photoaffinity crosslinking.** Site-specific photo-reactive DNA probes with 601-nucleosome positioning sequence for 70N0 nucleosomes were synthesized by enzymatic incorporation of modified nucleotides into double stranded DNA.27,28, DUMP and DCMG analogs coupled with p-azidophenylalanine bromide were incorporated along with [α-32P]dGTP/dATP, in tandem, at specific positions. Photo-reactive DNA was reconstituted into nucleosomes and bound with saturating amounts of wild type or mutant INO80 at 30 °C for 30 min. The extent of enzyme binding was assessed on 4% native polyacrylamide gels. In all experiments >90% of nucleosomes were bound by the enzyme. After binding, INO80 was crosslinked to DNA by UV irradiation (3 min at 310 nm, 2.65 mW cm−2), and DNA was digested with DNaseI and S1 nuclease for transfer of the radioactive label into Klenow linked subunits (or protein-DNA adducts) was synthesized and reduced, in the presence of pre-cooled IP buffer containing 25 mM Na-HEPES (pH 7.6), 300 mM NaCl, 10% glycerol, 0.1% NP-40, 10 mM 2-mercaptoethanol (BME), 1 μg ml−1 leupeptin, 1 μg ml−1 pepstatin A, and 1 mM PMSE. One microfilament and three microtubes of anti-α-FLAG antibodies were pre-incubated by Drs. Bruce Stillman and Ewa Kowalczyk, with the reactions, which were incubated at 4°C overnight, followed by an additional hour at 4°C with equilibrated EZView Protein A-agarose (Sigma) with gentle agitation. Agarose beads were washed three times with IP buffer and resuspended in SDS buffer. Samples were analyzed on 4–20% gradient SDS-polyacrylamide gels.

**Isolation of Arp8 and Arp4 subunits for peptide mapping.** INO80 photoaffinity-labeling reactions were scaled up for peptide mapping. Following digestion of DNA with DNaseI and S1 nuclease and preparation of radiolabeled subunit(s) were visualized by phosphorimaging. Data images in figures are representative of ≥3 experiments.

Peptide mapping with cyanogen bromide. Dried, photoaffinity-labeled Arp8 and Arp4 samples were resuspended in 2% SDS and 0.2 mM 2-mercaptoethanol, and divided into several 30 μl aliquots. For each sample, 2 μl of 1 M NaOH, 2 μl of various concentrations (20 mM to 2 M) of freshly prepared cyanogen bromide (CNBr) solution (in acetone) was mixed with the protein samples and incubated at 25°C in the dark for up to 2 h to vary the degree of digestion. CNBr reactions were stopped by diluting the samples with 170 μl of water and drying in a Centrivap concentrator (Labconco). The dried CNBr cleavage samples were resuspended in SDS sample buffer and neutralized by adding 0.5 M Tris-base. Samples were analyzed by 4–20% gradient SDS-PAGE for Arp8 and 10–20% gradient SDS-PAGE for Arp4. The sizes of the [32P]-labeled fragments were compared to [35S]-labeled, in vitro synthesized peptide markers corresponding to the expected protein fragments. Data images in figures are representative of three experiments.

Peptide mapping with 2-Nitro-5-thiocyanatobenzoic acid. Dried, photoaffinity-labeled Arp8 and Arp4 samples were resuspended in freshly prepared denaturing buffer containing 8 M urea, 10 mM 2-mercaptoethanol, and 10% D2O at 37°C for 30 min or 100 μM Tris-acetate (pH 8.5) by incubating at 37°C for 3 hours prior to dividing into several 36 μl aliquots. A volume of 2 μl of various concentrations (30 mM to 1 M) of NTCA (dissolved in D2O) were added and incubated at room temperature for 15 minutes. At this stage, NTCA modifies cysteine without cleavage. To initiate cleavage, the pH of the solution was adjusted to ~9.0–9.5 with 1 M NaOH as measured by a micro pH-probe, followed by incubation at 37°C for 4 hours. The cleavage reactions were stopped by adding SDS sample buffer and heating at 90°C for 10 min. Samples were analyzed on 4–20% gradient SDS-PAGE gels for Arp8 and 10–20% gradient SDS-PAGE gels for Arp4. The sizes of the [32P]-labeled fragments were compared to [35S]-labeled, in vitro synthesized peptide markers corresponding to the expected protein fragments. Data images in figures are representative of three experiments.

Peptide mapping of Ino80 subunit with ArgC protease. Photoaffinity-labeled INO80 complex (after digestion of DNA and label transfer) was denatured with 8 M urea, 10 mM 2-mercaptoethanol, and 10% D2O at 37°C for 3 hours prior to dividing into several 36 μl aliquots. A volume of 2 μl of various concentrations (30 mM to 1 M) of NTCA (dissolved in D2O) were added and incubated at room temperature for 15 minutes. At this stage, NTCA modifies cysteine without cleavage. To initiate cleavage, the pH of the solution was adjusted to ~9.0–9.5 with 1 M NaOH as measured by a micro pH-probe, followed by incubation at 37°C for 4 hours. The cleavage reactions were stopped by adding SDS sample buffer and heating at 90°C for 10 min. Samples were analyzed on 4–20% gradient SDS-PAGE gels for Arp8 and 10–20% gradient SDS-PAGE gels for Arp4. The sizes of the [32P]-labeled fragments were compared to [35S]-labeled, in vitro synthesized peptide markers corresponding to the expected protein fragments. Data images in figures are representative of three experiments.
Western blots. Pro tease (ArG) digested Ino80 fragments were resolved in 20 cm x 20 cm 4–20% Tris-glycine SDS-polyacrylamide gels and transferred onto PVDF membranes using Bio-Rad Trans-Blot® electrophoretic transfer cell for 3 hours at 4°C, using a transfer buffer containing 25 mM Tris, 192 mM glycine, 20% methanol, 0.1% SDS (pH 8.3) at 20 V constant voltage. The membranes were blocked with 5% fat-free milk in TBST (20 mM Tris-HCl pH 7.5, 150 mM NaCl, 0.1% Tween-20), overnight at 4°C, washed with TBST, and incubated with mouse monoclonal ANTI-FLAG® M2–Peroxidase (HRP) antibody (Sigma Aldrich catalog # A8592) diluted 1:1000 for 1 hour at room temperature. The blots were washed with TBST, and developed with SuperSignal West Femto Maximum Sensitivity Substrate (Thermo Fisher) before visualization using an Image Quant LAS 4000 (GE healthcare Life Sciences). Uncropped images of western blots are shown in Supplementary Fig. 6.

Synthesis of Arp8 and Arp4 peptide-mapping markers. The arp8 and arp4 genes, plus 100 bp of DNA flanking each end of the coding sequences, were PCR-amplified from genomic DNA isolated from the same yeast strain used for INO80-FLAG purification. This DNA was used as templates for PCR amplification of expression cassettes with coding sequences corresponding to Arp8 and Arp4 fragments expected from NTC or CNBr cleavage, under the control of 17 promoter and terminator. Arp8 and Arp4 peptides of specific lengths and molecular weights were synthesized by coupled in vitro transcription/translation using the Tn1® Quick Coupled Transcription/Translation System (Promega) with [35S]-Methionine (Perkin Elmer). Transcription/translation reactions were carried out at 30 °C for 3 hours at 4°C, using a transfer buffer containing 25 mM Tris, 192 mM glycine, 0.1% Tween-20, and 0.1% SDS (pH 8.3) at 50 V constant voltage. The membranes were equilibrated with an equal volume of 100 mM sodium phosphate (pH 7.4) and 0.1 leupeptin. Samples were analyzed on 4-20% Tris-glycine gradient SDS-polyacrylamide gels followed by phoshorimaging. Peptide standards corresponding to both incomplete or limited digestion as well as complete digestion conditions were prepared. Oligonucleotide primers for Arp8 and Arp4 peptide synthesis are listed in Supplementary Table 1.

Synthesis of INO80-FLAG peptide-mapping markers. The ino80 gene with 2FLAG coding sequence at the 3’ end, plus 100 bp of DNA flanking each end of the coding sequence, was PCR-amplified from genomic DNA isolated from the same yeast strain used for INO80-FLAG purification. This DNA was used as template for PCR amplification of expression cassettes with coding sequences corresponding to truncated INO80-FLAG fragments of varying lengths from the C-terminus. Peptides were synthesized by coupled in vitro transcription/translation as described above. Peptide standards of all sizes were pooled and run alongside INO80 peptide-mapping reactions. Relative electrophoretic mobilities (Rf) of the peptide marker from three technical replicates were calculated and plotted against log10 of their molecular weights to generate a standard curve17. Molecular weights and sizes of protein fragments generated by ArgC cleavage were estimated from the standard curve based on their Rf. Protease cleavage maps were mapped to arginine residues closest to the estimated cleavage sites. Oligonucleotide primers for Ino80 peptide synthesis are listed in Supplementary Table 1.

Yeast growth assays. Purified single colonies of wild-type INO80, INO80-ΔArp8, INO80-ΔArp8p(arp8), INO80-ΔArp8ΔIno, INO80-ΔIno, and INO80-ΔTEL1 strains were diluted in water to O.D.600 0.5 M LiCl and 0.5 M formic acid. Experiments were repeated four times. Measurements of inorganic 32P (Pi) released, which is resolved from un-hydrolyzed ATP and ADP with nucleosomes would be completely bound. Nucleosome movement was initiated by adding 80 μM ATP and was stopped with 1.5 mM ATP γ-S and excess EDTA at the indicated time points (Supplementary Fig. 3b). Aliots were analyzed on 5% native polyacrylamide gels (as in Supplementary Fig. 3a). For site-directed histone-DNA crosslinking25, samples were irradiated with UV (3 min at 310 nm, 2.65 mw cm-2). Samples were denatured with 0.1% SDS at 37 °C for 20 min in 30 mM NaCl and 20 mM Tris-HCl (pH 8.0). Crosslinked protein-DNA were enriched and separated in un-crosslinked DNA by electrophoresis. The radioactive phase containing un-crosslinked DNA was discarded. Crosslinked DNA was ethanol precipitated with 1 M LiCl in the presence of sheared salmon sperm DNA as carrier. Crosslinked DNA was cleaved with 1 M pyridoline (Sigma) at 90°C for 20 min. DNA samples were analyzed alongside a sequence ladder made from the same DNA, on denaturing 6.5% polyacrylamide gels containing 8 M urea. Gels were visualized by phosphorimagning and quantified using ImageQuant software (Version 5.2). Total lane intensity was normalized to correct for loading bias using Microsoft Excel. Data images in figures are representative of three experiments.

Nucleosome binding assays. Increasing concentrations of wild type or mutant INO80 as indicated (Supplementary Fig. 4a) were incubated with 40 nM homogeneous 70N0 nucleosomes (discussed above) for 30 min at 30°C in 10 mM Na-HEPES (pH 7.8), 4 mM MgCl2, 60 mM NaCl, 0.2 mM EGTA, 0.04 mM EDTA, 8% glycerol, and 0.1 μg μl-1 bovine serum albumin. Reactions were analyzed by resolving enzyme-bound nucleosomes from free nucleosomes on 4% native polyacrylamide gels in 1x Tris-EDTA buffer. Data images in figures are representative of three experiments.

Histone crosslinking. Recombinant Xenopus laevis histones with amino acids replaced by cysteine at specific positions (Fig. 4a) were expressed, purified, and reconstituted into octamers with other histones as discussed above. Reducing conditions with 5 mM 2-mercaptoethanol were maintained for cysteine mutant histones and octamers. Reduced 50 μM and 100 μM nucleosomes were immobilized on agarose-entrapped 20-fold molar excess of PEAS [N-[[(2-pyridldithio)ethyl]4-azidosalicylamide] (Thermo Fisher) and incubated at room temperature for 30 minutes. Excess unconjugated PEAS was removed by buffer exchange (100-fold) with octamer dilution buffer (5 M Tris-HCl pH 8, 0.5 mM EDTA, 1 M NaCl) and subsequent concentration using a 30 KDa cutoff Amicon ultrafilter column. The octamers were stored at −20°C in 50% glycerol. Homogenous 70N0 nucleosomes formed with the 601-nucleosome positioning sequence containing Cys5 at one end were reconstituted as discussed above. [125I] (Perkin Elmer) was added into IODOGEN tubes (Pierce) in a 2.5 times molar excess over that of the nucleosomes to be iodinated, and pre-equilibrated with an equal volume of 100 mM sodium phosphate (pH 7.4). After 6 minutes of incubation, nucleosomes were added and incubated at room temperature for 5 minutes. The iodination reaction was stopped by adding 1μl of 2.5 mM tyrosine and 1μl of 80 mM methionine. Unconjugated [125I]I was removed using Sephadex G25 spin-columns centrifuged at 800 × g for 2 minutes at room temperature. The amount of nucleosome-conjugated PEAS was determined by measuring the specific activity of [125I] in nucleosomes compared to the specific activities of known molar quantities of unconjugated PEAS-[125I]. Nucleosomes (70N0) with photo-reactive octamers were bound to saturation amounts of wild type or mutant INO80 at 30°C and assessed as before. Samples were irradiated directly in buffer (18 min) using a UV (3 minutes, 100 cm distance from transilluminator). Growth assays were repeated three times, each from separate single colonies of yeast with wild type or mutant INO80.

Data availability. The data sets generated and/or analyzed during the current study are available from the corresponding author on reasonable request.

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**Author contributions**

DNA crosslinking experiments were performed by S.B. and M.U., peptide mapping and site-directed mapping experiments were done by S.B., and histone crosslinking experiments by S.P., M.U. S.B. constructed HA-tagged strains, M.N. constructed the Arp8ΔN strain, performed growth assays, and purified Arp8ΔN INO80. S.B. and M.N. performed all the other biochemical assays with the Arp8ΔN INO80 complex. The experiments were directed by B.B. and the manuscript was written by B.B. and S.B. with contributions from S.P. and M.N.

**Additional information**

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**Competing interests:** The authors declare no competing interests.

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