Supplementary Information Text

METHODS

Structural biology software was accessed through the SBGrid consortium (1).

Protein expression and purification.

 SARS-CoV-2 nsp12. SARS-CoV-2 nsp12 was expressed and purified as described (1). A pRSFDuet-1 plasmid expressing SARS-CoV-2 His6-SUMO- nsp12 (Addgene plasmid 159107) was transformed into *Escherichia coli* (*Eco*) BL21-CodonPlus cells (Agilent). Cells were grown, followed by the addition of isopropyl β-d-1-thiogalactopyranoside (IPTG) to induce protein expression overnight. Cells were collected by centrifugation, resuspended and lysed in a continuous-flow French press (Avestin). The lysate was cleared by centrifugation, loaded onto a HiTrap Heparin HP column (Cytiva), and then eluted using a salt gradient. The fractions containing nsp12 were pooled and loaded onto a HisTrap HP column (Cytiva), washed, and eluted. Eluted nsp12 was dialyzed overnight in the presence of His6-Ulp1 SUMO protease. Cleaved nsp12 was passed through a HisTrap HP column (Cytiva). Flow-through was collected, concentrated by centrifugal filtration (Amicon), and loaded on a Superdex 200 Hiload 16/600 (Cytiva) for size-exclusion chromatography. Glycerol was added to the purified 616 nsp12, aliquoted, flash frozen with liquid N_2 , and stored at -80 $^{\circ}$ C. *SARS-CoV-2 nsp7/8*. SARS-CoV-2 nsp7/8 was expressed and purified as described (1). The pCDFDuet-1 plasmid expressing SARS-CoV-2 His6-ppx- nsp7/8 (ppx is a Prescission Protease cleavage site; Addgene plasmid 159092) was transformed into *Eco* BL21(DE3). Cells were grown and protein expression was induced overnight by the addition of IPTG. Cells were collected by centrifugation, resuspended, and lysed in a continuous-flow French press (Avestin). The lysate was cleared by centrifugation, then loaded onto a HisTrap HP column (Cytiva), washed, and eluted. Eluted nsp7/8 was dialyzed overnight in 625 the presence of His $_6$ -Prescission Protease to cleave the His $_6$ -tag. Cleaved nsp7/8 was passed through a HisTrap HP column (Cytiva). Flow-through was collected, concentrated by centrifugal filtration (Amicon), and loaded onto a Superdex 75 Hiload 16/600 (Cytiva). Glycerol was added to the purified nsp7/8, aliquoted,

629 flash frozen with liquid N_2 , and stored at -80 $^{\circ}$ C.

 SARS-CoV-2 nsp13. SARS-CoV-2 nsp13 was expressed and purified as 631 described (1). The pet28 plasmid containing $SARS-CoV-2 His_6-ppx-nsp13$ (Addgene plasmid 159390) was transformed into *Eco* Rosetta(DE3) (Novagen). Cells were grown, followed by the addition of IPTG to induce protein expression overnight. Cells were collected by centrifugation, resuspended, and lysed in a continuous-flow French press (Avestin). The lysate was cleared by centrifugation, then loaded onto a HisTrap HP column (Cytiva), washed, and eluted. Eluted 637 nsp13 was dialyzed overnight in the presence of His_6 -Prescission Protease to cleave His6-tag. Cleaved nsp13 was passed through a HisTrap HP column (Cytiva). Flow-through was collected, concentrated by centrifugal filtration (Amicon), and loaded onto a Superdex 200 Hiload 16/600 (Cytiva). Glycerol was 641 added to the purified nsp13, aliquoted, flash frozen with liquid N_2 , and stored at -80°C.

 Native electrophoretic mobility shift assays. Nsp12 or nsp12-D760A were incubated with 3-fold molar excess of nsp7/8 in transcription buffer (120 mM K- acetate, 20 mM HEPES pH 8, 10 mM MgCl2, 2 mM DTT) to assemble holo-RdRp 647 (2 μ M final). The resulting complex was incubated with 1 μ M of annealed RNA scaffold (Horizon Discovery) for 5 minutes at 30°C. Nsp13 and pre-mixed ADP and AlF³ (Sigma-Aldrich) were added to a final concentration of 2 μM and 2 mM, respectively, and incubated for an additional 5 minutes at 30°C. Reactions were analyzed by native gel electrophoresis on a 4.5% polyacrylamide native gel (37.5:1 acrylamide:bis-acrylamide) in 1X TBE (89 mM Tris, 89 mM boric acid, 1 mM EDTA) at 4°C. The gel was stained with Gel-Red (Biotium).

Native mass spectrometry (nMS) analysis. The reconstituted sample

656 containing 4 μ M RTC and 8 μ M nsp13 incubated with 2 mM ADP-AIF₃ was

buffer-exchanged into 150 mM ammonium acetate, 0.01% Tween-20, pH 7.5

using a Zeba microspin desalting column with a 40 kDa MWCO (ThermoFisher

Scientific). For nMS analysis, a 2–3 µL aliquot of the buffer-exchanged sample

 was loaded into a gold-coated quartz capillary tip that was prepared in-house and then electrosprayed into an Exactive Plus with extended mass range (EMR) instrument (Thermo Fisher Scientific) with a static direct infusion nanospray source (2). The MS parameters used: spray voltage, 1.2 kV; capillary temperature, 150 °C; in-source dissociation, 0 V; S-lens RF level, 200; resolving 665 power, 17,500 at m/z of 200; AGC target, 1 x 10⁶; maximum injection time, 200 ms; number of microscans, 5; injection flatapole, 6 V; interflatapole, 4 V; bent flatapole, 4 V; high energy collision dissociation (HCD), 200 V; ultrahigh vacuum 668 pressure, 7.2×10^{-10} mbar; total number of scans, at least 100. Mass calibration in positive EMR mode was performed using cesium iodide. For data processing, the acquired MS spectra were visualized using Thermo Xcalibur Qual Browser (v. 4.2.47). MS spectra deconvolution was performed either manually or using the software UniDec v. 4.2.0 (3, 4). The following parameters were used for the UniDec processing: m/z range, 7,000 – 10,000 Th; background subtraction, subtract curved at 100; smooth charge state distribution, enabled; peak shape function, Gaussian; Beta Softmax distribution parameter, 20.

 The expected masses for the component proteins based on previous nMS experiments (1) include nsp7: 9,137 Da; nsp8 (N-terminal Met lost): 21,881 Da; 678 nsp13 (post-protease cleavage, has three Zn^{2+} ions coordinated with 679 9 deprotonated cysteine residues): 67,464 Da, and nsp12 (has two Zn^{2+} ions coordinated with 6 deprotonated cysteine residues): 106,785 Da. The mass of the assembled RNA duplex scaffold is 30,512 Da.

682 Experimental masses were reported as the average mass \pm standard deviation (S.D.) across all the calculated mass values within the observed charge state series. Mass accuracies were calculated as the % difference between the measured and expected masses relative to the expected mass. The observed mass accuracies ranged from 0.016 – 0.035%.

 Preparation of SARS-CoV-2 nsp13-BTC⁵ for Cryo-EM. Purified nsp12 and 689 nsp7/8 were mixed in a 1:3 molar ratio and incubated at 22° C for 15 minutes.

- The mixture was buffer-exchanged into cryo-EM buffer (20 mM HEPES pH 8.0,
- 150 mM K-acetate, 10 mM MgCl2, 1 mM DTT) using Zeba desalting columns
- (ThermoFisher Scientific) and incubated with annealed BTC5-scaffold (Fig. 1A) in
- a 1:1.5 molar ratio. Purified nsp13 was concentrated by centrifugal filtration
- (Amicon) and buffer exchanged into cryo-EM buffer using Zeba desalting
- columns. Buffer exchanged nsp13 was mixed with ADP and AlF³ and then added
- to nsp7/8/12/RNA scaffold at a molar ratio of 1:1 with a final concentration of
- 697 $\,$ 2 mM ADP-AIF₃. Complex was incubated for 5 minutes at 30 $\,^{\circ}$ C and further
- concentrated by centrifugal filtration (Amicon).
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- **Cryo-EM grid preparation.** Prior to grid freezing, 3-([3-
- cholamidopropyl]dimethylammonio)-2-hydroxy-1-propanesulfonate (CHAPSO,
- Anatrace) was added to the sample (8 mM final), resulting in a final complex
- concentration of 10 µM. The final buffer condition for the cryo-EM sample was
- 20 mM HEPES pH 8.0, 150 mM K-acetate, 10 mM MgCl2, 1 mM DTT,
- 2 mM ADP-AlF3, 8 mM CHAPSO. C-flat holey carbon grids (CF-1.2/1.3-4Au,
- Electron Microscopy Sciences) were glow-discharged for 20 s prior to the
- application of 3.5 μL of sample. Using a Vitrobot Mark IV (ThermoFisher
- Scientific), grids were blotted and plunge-frozen into liquid ethane with 90%
- chamber humidity at 4°C.
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 Cryo-EM data acquisition and processing. Structural biology software was accessed through the SBGrid consortium (5). Grids were imaged using a 300 kV Titan Krios (ThermoFisher Scientific) equipped with a K3 camera (Gatan) and a BioQuantum imaging filter (Gatan). Images were recorded using Leginon (6) with a pixel size of 1.065 Å/px (micrograph dimensions of 5,760 x 4,092 px) over a nominal defocus range of -0.8 μm to -2.5 μm and 30 eV slit width. Movies were recorded in "counting mode" (native K3 camera binning 2) with ~30 e-/px/s in dose-fractionation mode with subframes of 50 ms over a 2.5 s exposure (50 frames) to give a total dose of ~66 e-/Å. Dose-fractionated movies were gain-

 normalized, drift-corrected, summed, and dose-weighted using MotionCor2 (7). The contrast transfer function (CTF) was estimated for each summed image using the Patch CTF module in cryoSPARC v2.15.0 (8) Particles were picked and extracted from the dose-weighted images with box size of 256 px using cryoSPARC Blob Picker and Particle Extraction. The entire dataset consisted of 10,685 motion-corrected images with 4,961,691 particles. Particles were sorted using cryoSPARC 2D classification (N=100), resulting in 2,412,034 curated particles. Initial models (Ref 1: decoy 1, Ref 2: complex, Ref 3: decoy 2; SI Appendix; Fig. S2) were generated using cryoSPARC *ab initio* Reconstruction on a subset of 85,398 particles. Particles were further curated using Ref 1-3 as 3D templates for cryoSPARC Heterogeneous Refinement (N=6), resulting in the following: class1 (Ref 1), 258,097 particles; class2 (Ref 1), 263,966 particles; class3 (Ref 2), 668,743 particles; class4 (Ref 2), 665,480 particles; class5 (Ref 3), 280,933 particles; class6 (Ref 3), 274,815 particles. Particles from class3 and class4 were combined and further curated with another round of Heterogeneous 735 Refinement (N=6), resulting in the following: class1 (Ref 1), 67,639 particles; class2 (Ref 1), 61,097 particles; class3 (Ref 2), 553,368 particles; class4 (Ref 2), 554,581 particles; class5 (Ref 3), 42,114 particles; class6 (Ref 3), 55,424 particles. Curated particles from class3 and class4 were combined, re-extracted with a box size of 320 px, and further classified using Ref 2 as a 3D template for cryoSPARC Heterogeneous Refinement (N=4). Classes from this round of Heterogeneous Refinement (N=4) were as follows: class1 (Ref 2), 871,163 particles; class2 (Ref 2), 77,769 particles; class3 (Ref 2), 61,489 particles; class4 (Ref 2), 64,026 particles. Particles from class1 and class2 were combined and further sorted using Heterogeneous Refinement (N=4) using class maps as templates, resulting in the following: class1, 134,536 particles; class2, 270,170 particles; class3, 294,162 particles; class4, 172,295 particles. Classification 747 revealed two unique classes: $nsp13₁-BTC$ (class1 and class2) and $nsp13₂-BTC$ (class3 and class4). Particles within each class were further processed using RELION 3.1-beta Bayesian Polishing(9, 10). Polished particles were refined using cryoSPARC Non-uniform Refinement, resulting in structures with the

 following particle counts and nominal resolutions: nsp131-BTC (404,706 particles; 752 3.40 Å) and nsp13₂-BTC (466,457 particles; 3.45 Å).

 To improve the resolution of the RNA in the BTC, particles from both classes were combined in a cryoSPARC Non-uniform Refinement and density corresponding to nsp13 was subtracted. Subtracted particles were further refined with cryoSPARC Local Refinement using a mask encompassing the BTC and a 757 fulcrum point defined on the backtracked RNA. This map, BTC₅(local), contained 871,163 particles with a nominal resolution of 3.23 Å.

759 To improve the density of nsp13.2 in the nsp13₂-BTC map, particles were subtracted using a mask defined around nsp13.2, leaving residual signal for only nsp13.2. Subtracted particles were classified (N=4) in RELION 3.1 beta using a mask around nsp13.2, resulting in the following classes: class1, 71,607 particles; class2, 163,540 particles; class3, 176,461 particles; class4, 54,849 particles. Subtracted particles in class1 and class2 were combined and reverted back to the original particles, followed by refinement using cryoSPARC Non-uniform Refinement. The resulting map of nsp132-BTC contains 235,147 particles with nominal resolution of 3.59 Å. Local resolution calculations were generated using blocres and blocfilt from the Bsoft package (11).

 Model building and refinement. Initial models were derived from PDB: 6XEZ (1). The models were manually fit into the cryo-EM density maps using Chimera (12) and rigid-body and real-space refined using Phenix real_space_refine (13). For real-space refinement, rigid body refinement was followed by all-atom and B- factor refinement with Ramachandran and secondary structure restraints. Models were inspected and modified in Coot (14).

 4-thiouridine crosslinking. Nsp12 or nsp12-D760A were incubated with 3-fold molar excess of nsp7/8 to assemble holo-RdRp (2 μM final) in transcription

buffer. The resulting holo-RdRp was added to a modified RNA scaffold

 (SI Appendix; Fig. S5A) containing a photoactivable 4-thiouridine base (Horizon Discovery) which was 5'-labelled by T4-polynucleotide kinase (New England 782 Biolabs) with $y^{-32}P$ -ATP (Perkin-Elmer). The holo-RdRp/RNA complex was left to incubate for 5 minutes at 30°C in the dark. Nsp13 and ATP were added to a final concentration of 2 uM and 2 mM, respectively, and incubated for five minutes at 30°C in the dark. The reaction mixture was transferred to a Parafilm covered aluminum block at 4°C and irradiated with a 365-nm handheld UV lamp. Reactions were quenched with LDS sample loading buffer (ThermoFisher Scientific) and analyzed by gel electrophoresis on a NuPAGE 4-12% Bis-Tris gel (ThermoFisher) at 150 Volts for 1 hour and visualized by autoradioagraphy.

Molecular dynamics simulations

 General simulation setup and parameterization. Proteins, ADP, and ions were parameterized with the DES-Amber SF1.0 force field (15). RNAs were parameterized with the Amber ff14 RNA force field (16) with modified electrostatic, van der Waals, and torsional parameters to more accurately reproduce the energetics of nucleobase stacking (17). The systems were solvated with water parameterized with the TIP4P-D water model (18) and neutralized with 150 mM NaCl buffer. The systems each contained ~887,000 atoms in a 190×190×190 Å cubic box.

 Systems were first equilibrated on GPU Desmond using a mixed NVT/NPT schedule (19), followed by a 1 µs relaxation simulation on Anton, a special- purpose machine for molecular dynamics simulations (20). All production simulations were performed on Anton and initiated from the last frame of the relaxation simulation. Production simulations were performed in the NPT ensemble at 310 K using the Martyna-Tobias-Klein barostat (21). The simulation time step was 2.5 fs, and a modified r-RESPA integrator (22, 23) was used in which long-range electrostatic interactions were evaluated every three time steps. Electrostatic forces were calculated using the *u*-series method (24). A 9-Å cutoff was applied for the van der Waals calculations.

 System preparation. The nsp132-BTC-1U+1C and the nsp132-BTC-1U+1U complexes 811 were prepared from the cryo-EM structure of the $nsp132-BTC_5$. AlF₃ and CHAPSO were removed. Cytosines at the +2 and +3 positions of the p-RNA were removed, and the cytosine at −1 was mutated to uracil. The resulting p- RNA had a matched −1U and a mismatched +1C in nsp132-BTC−1U+1C, and a matched −1U and +1U in nsp132-BTC−1U+1U. Missing loops and termini in proteins were capped with ACE/NME capping groups. The two complexes were prepared for simulation using the Protein Preparation Wizard in Schrödinger Maestro. After a 1 µs relaxation simulation of the nsp132-BTC−1U+1C complex, the −1U of the p-RNA formed a Watson-Crick base pair with the −1A in the t-RNA, and the +1C of p-RNA formed a non-Watson-Crick C-A hydrogen bond with the +1A of the t-RNA in the active site. After a 1 µs relaxation simulation of the nsp132-BTC−1U+1U complex, the −1U and +1U of the p-RNA formed Watson-Crick base pairs with the −1A and +1A of the t-RNA respectively. *Simulation analysis.* All simulations were visually inspected using the in-house

visualization software Firefly. The average root-mean-square deviation (RMSD)

826 was calculated for +1C (or +1U) of the p-RNA between the last frame of the 1 µs

- relaxation simulation and instantaneous structures from the trajectories, aligned
- on the entire nps12 module.

Quantification and statistical analysis. The nMS spectra were visualized using

Thermo Xcalibur Qual Browser (versions 3.0.63 and 4.2.27), deconvolved using

UniDec versions 3.2 and 4.1 (3, 4) and plotted using the m/z software

(Proteometrics LLC, New York, NY). Experimental masses (SI Appendix;

834 Fig. S1B and C) were reported as the average mass \pm standard deviation across

 all the calculated mass values obtained within the observed charge state distribution.

 The local resolution of the cryo-EM maps (SI Appendix; Fig. S3B-D) was estimated using blocres (11) with the following parameters: box size 15, sampling 1.1, and cutoff 0.5. Directional 3D FSC (SI Appendix; Fig. S3H-J) were calculated

- by 3DFSC (25). The quantification and statistical analyses for model refinement
- and validation were generated using MolProbity (26) and PHENIX (13).
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861 **Table S1. Cryo-EM data collection, refinement, and validation statistics.**

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865 **SUPPLEMENTAL FIGURES**

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Mass assignments from nMS analysis of BTC₆ incubated with nsp13 and $MgADPAIF₃$

* Calculated from the average and S.D. of all the measured masses across the charge-state distribution ($n \ge 4$).

Figure S1

Fig. S1. Native gel electrophoresis mobility shift assay and nMS analysis of the

- **BTC.**
- **A.** A native gel electrophoretic mobility shift assay reveals that wt-holo-RdRp requires
- 871 nsp13(ADP-AIF₃) to bind the BTC₅-scaffold efficiently (compare lanes 1, 2, and 6) but
- holo-RdRp with nsp12-D760A does not require nsp13 (lane 4).
- **B.** The nMS spectrum and the deconvolved mass spectrum showing assembly of stable
- 874 nsp13-BTC $_6$ complexes. The peak for the nsp13₂-BTC $_6$ assembly is present at about
- 875 \sim 9% intensity relative to the predominant peak from nsp13₁-BTC₆.
- **C**. Mass assignments of the deconvolved peaks from the nMS analysis.
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Figure S2

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Fig. S3. Cryo-EM analysis.

- **A.** Gold-standard FSC plots for nsp131-BTC5, nsp132-BTC5, and BTC5(local), calculated
- by comparing two independently determined half-maps from cryoSPARC (8). The
- dotted line represents the 0.143 FSC cutoff.
- **B-D.** Cryo-EM reconstructions filtered by local resolution(11). The view on the right is a cross-section.
- *(top)* Colored by subunit according to the color key.
- *(bottom)* Color by local resolution (key on the bottom).
- **B.** Nsp131-BTC5.
- **C.** Nsp132-BTC5.
- **D.** BTC5(local).
- **E – G.** FSC calculated between the refined structures and the half map used for
- refinement (work, red), the other half map (free, blue), and the full map (black).
- 904 **E.** Nsp13₁-BTC₅.
- **F.** Nsp132-BTC5.
- **G.** BTC5(local).
- **H - J,** Directional 3D Fourier shell correlation plots, calculated by 3DFSC(25).
- **H.** Nsp131-BTC5.
- 909 **I.** Nsp13₂-BTC₅.
- 910 **J.** BTC₅(local).
- **K – M.** Particle angular distribution plots calculated in cryoSPARC. Scale shows the
- number of particles assigned to a particular angular bin. Blue, a low number of particles;
- red, a high number of particles.
- **K.** Nsp131-BTC5.
- 915 **L.** Nsp13₂-BTC₅.
- 916 **M.** BTC₅(local).

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Figure S4

Fig. S4. Sequence conservation of nsp12 homologs and NTP-entry tunnel environment.

 A. Sequence alignment of nsp12 homologs from six pathogenic and model CoV family members, covering RdRp motifs (27) (motifs F, C, D, and E denoted at the top of the sequence alignment) architecturally important for the NTP-entry tunnel. Selected residues discussed in the text are highlighted (red outlines). Sequence logos(28) for motif F and motif E are shown, with residues that interact with the backtracked RNA highlighted (colored dots above; see Figure 4). The sequence logos were generated 930 from an alignment of 97 RdRp sequences from α -, β -, γ -, and δ -CoVs (Data S1). **B.** Views from the outside into the NTP-entry tunnels of the SARS-CoV-2 BTC (*left*), an

E. coli DdRp BTC [PDB ID: 6RI9; (29)] and an *S. cerevisiae* DdRp BTC [PDB ID: 3GTP;

- (30)]. Protein surfaces are colored by the electrostatic surface potential [calculated
- using APBS; (31)]. Backtracked RNA is shown as atomic spheres with yellow carbon
- atoms.
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Figure S5

- 939 **A.** Protein-RNA crosslinking analysis: The 5'-[³²P]-labelled RTC(4-thio-U)-scaffold and
- the indicated proteins were incubated along with 2 mM ATP (present in every lane),
- exposed to UV as indicated, then analyzed by SDS polyacrylamide gel electrophoresis
- and autoradiography. The positions of nsp8, nsp12, and nsp13 bands are indicated.
- Lanes 1 and 5, containing nsp13 only, are identical controls indicating uniform UV
- exposure across the samples. Holo-RdRp(*) denotes the nsp12-D760A substitution that
- facilitates backtracking (see Figure S1A). The two panels show the same SDS
- polyacrylamide gel (left panel, Coomassie stained; right panel, visualized by
- autoradiography).
- 948 **B.** Protein-RNA crosslinks are specific. Lanes 1, 2; Analysis using the 5'-[³²P]-RTC(4-
- thio-U)-scaffold (RNA-scaffold 'a' shown on the bottom). Crosslinking to nsp12 serves
- as a positive control for the crosslinking reaction. Lanes 3-6; Analysis using RNA-
- 951 scaffold 'b' (RTC-scaffold with 5'-[³²P]-labelled p-RNA). Lanes 7-10: Analysis using
- 952 RNA-scaffold 'c' (RTC-scaffold with 5'-[³²P]-labelled t-RNA). The complete absence of
- protein-RNA crosslinks in lanes 3-10 indicates that the observed protein-RNA crosslinks
- arise from the 4-thio-U site-specifically incorporated in the p-RNA of the RTC(4-thio-U)-
- scaffold.
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Figure S6

Fig. S6. Molecular dynamics simulations of nsp132-BTC1U+1C vs.

nsp132-BTC-1U+1U.

- Molecular dynamics simulations of the nsp132-BTC−1U+1C (*top*) and nsp132-BTC-1U+1U
- (*bottom*) complexes. The schematics illustrate the active-site proximal nucleotides in
- each modeled complex. Each complex was simulated with 3 replicates. RMSD values plotted as a function of time represent the heavy-atom RMSD of the +1 nucleotide of the
- 966 p-RNA (+1C for nsp13₂-BTC-1U+1c or matched +1U for nsp13₂-BTC-1U+1U) compared with
- the starting configuration (see Methods). The RMSD histograms (plotted on the right) are aggregates of all 3 replicates.
- (*top*) Nsp132-BTC-1U+1C. As shown in Figure 5C, the mismatched p-RNA +1C spends
- 970 about 60% of the time frayed from the t-RNA +1A and near or in the NTP-entry tunnel 971 (RMSD \ge ~3.5 Å).
- (*bottom*) Nsp132-BTC-1U+1U. With the p-RNA +1U matched with the t-RNA +1A for
- Watson-Crick base pairing, the p-RNA +1U does not fray and spends all of its time in
- the vicinity of the RdRp active site and base paired with the t-RNA.
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SUPPLEMENTAL DATA FILES

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