Compartmentalization-aided interaction screening reveals extensive high-order complexes within the SARS-CoV-2 proteome

Graphical abstract

Highlights
- CoPIC emerges as a simple and efficient method to investigate PPI in cellulo
- CoPIC screening reveals extensive high-order complexes within the SARS-CoV-2 RTC
- Profiling intraviral interactions provides targets for therapeutic intervention

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In brief
The unprecedented COVID-19 pandemic is caused by infection with severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2). Xu et al. establish an innovative interaction screening strategy based on phase separation in cellulo and reveal comprehensive high-order complexes within the SARS-CoV-2 proteome, thus providing more effective therapeutic opportunities.
Compartmentalization-aided interaction screening reveals extensive high-order complexes within the SARS-CoV-2 proteome

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SUMMARY

Bearing a relatively large single-stranded RNA genome in nature, severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) utilizes sophisticated replication/transcription complexes (RTCs), mainly composed of a network of nonstructural proteins and nucleocapsid protein, to establish efficient infection. In this study, we develop an innovative interaction screening strategy based on phase separation in cellulo, namely compartmentalization of protein-protein interactions in cells (CoPIC). Utilizing CoPIC screening, we map the interaction network among RTC-related viral proteins. We identify a total of 47 binary interactions among 14 proteins governing replication, discontinuous transcription, and translation of coronaviruses. Further exploration via CoPIC leads to the discovery of extensive ternary complexes composed of these components, which infer potential higher-order complexes. Taken together, our results present an efficient and robust interaction screening strategy, and they indicate the existence of a complex interaction network among RTC-related factors, thus opening up opportunities to understand SARS-CoV-2 biology and develop therapeutic interventions for COVID-19.

INTRODUCTION

The current pandemic of COVID-19 (coronavirus disease-2019), a respiratory disease leading to more than 164 million confirmed cases and 3.4 million deaths globally (World Health Organization, 2021), is caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), an enveloped, positive-sense, single-stranded RNA betacoronavirus of the family Coronaviridae (Zhu et al., 2020). SARS-CoV-2 possesses a remarkably large RNA genome, with a genome size of approximately 30 kb, flanked by 5' and 3' untranslated regions. The genomic RNA at the 5' end features the partly overlapping open reading frames (ORFs) 1a and 1ab, which occupy two-thirds of the capped and polyadenylated genome (Lu et al., 2020; Tan et al., 2005). The expression of ORF1ab requires a ribosomal frameshift near the end of ORF1a. In this manner, SARS-CoV-2 genome translation yields the large replicase polyproteins pp1a and pp1ab (Abduljalil and Abduljalil, 2020; Perlman and Netland, 2009). Pp1a and pp1ab go through extensive autoproteolytic processing, mediated by two ORF1a-encoded proteases and ultimately generating 16 nonstructural proteins (Nsps), which make up the primary constituents of the replication/transcription complexes (RTCs). These include a variety of key enzymes functioning for viral RNA synthesis, i.e., the primase Nsp8 (Kirchdoerfer and Ward, 2019; Konkoloova et al., 2020), RdRp-Nsp12 (Cheng et al., 2005; Shannon et al., 2020), the helicase Nsp13 (Jiang et al., 2020; Lee et al., 2010), the exoribonuclease Nsp14 (Minskaia et al., 2006), and the endoribonuclease NendoU-Nsp15 (Bhardwaj et al., 2004); several multi-spanning membrane proteins (Nsp3, Nsp4, and Nsp6) that presumably contribute to the formation of double-membrane vesicles (DMVs) (Cottam et al., 2011; Kanjanahaluethai et al., 2007; Oostra et al., 2007); proteases for polyprotein cleavage (Nsp5) (Stobart et al., 2013); and the accessory modulating factors hijacking host defense (Nsp1 and Nsp2) (Antonin et al., 2020; Cornillez-Ty et al., 2009). For the 3' one-third of the genome, ORFs that encode four structural proteins and nine accessory proteins are transcribed to form a nested set of subgenomic mRNAs (Abduljalil and Abduljalil, 2020).

The synthesis of SARS-CoV-2 RNA within host cells depends on the efficient and correct assembly of RTCs under the coordinated cascade of transcription and replication. This process takes place in intracellular DMVs (Klein et al., 2020; Snijder et al., 2020), which are actually remodeled to serve as a platform for protein-protein interactions (PPIs) and provide a favorable environment for replication. Although SARS-CoV-2 Nsps (e.g., Nsp7, Nsp8,
(legend on next page)
Nsp12, and Nsp13) are associated with the viral RTCs during infection (Yan et al., 2020), the molecular details of which Nsps and how they come together to form the viral RTCs are not clear. Notably, it is also reported that nucleocapsid (N) protein functions as the only structural protein shuttling inside/outside RTCs and facilitating the transcription and replication of coronavirus RNA (Cong et al., 2020; Sunjit and Lal, 2008). Recently released structural information revealed the interaction of the C-terminal domain of N protein with the pore factor of DMVs (Khan et al., 2021). As a critical step toward understanding this process, it is necessary to investigate the interaction network among Nsps. Information to date in this regard has been scarce, with only a limited number of interactions identified (Imbert et al., 2008; Pan et al., 2008; von Brunn et al., 2007).

Although multiple methods have been developed to detect PPIs, few are suited for high-throughput protein interaction analysis in cells. The most commonly used methods for investigating PPIs of viruses, including coronaviruses (He et al., 2004; Pan et al., 2008; von Brunn et al., 2007), yeast two-hybrid (Y2H) screening, and coimmunoprecipitation (coIP), have their own limitations and both suffer severe false-positive and false-negative results (Berggård et al., 2007; Miernyk and Thelen, 2008; Pizicky and Fields, 1995). To alleviate these drawbacks, in this study we developed a method to study PPIs in living cells, namely compartmentalization of protein-protein interactions in cells (CoPIC). Essentially, CoPIC is the adapted version of a phase separation-based method for PPIs in vitro in our previous study, called phase-separated condensate-aided enrichment of biomolecular interactions in test tubes (CEBIT) (Zhou et al., 2020). One major requirement for CEBIT is the availability of large quantities of well-behaved purified materials, which is difficult to achieve for intrinsically ill-behaved subjects such as large dynamic protein complexes. Correspondingly in CoPIC, one interaction partner is fused with the scaffolds that drive the formation of phase-separated condensates, and the other partner (named the client) is fused with a fluorescent protein, and can be recruited to the condensate via specific interactions in cells. Overall, CoPIC emerges as a simple, sensitive, and efficient method to investigate PPIs in mammalian cells.

To elucidate molecular mechanisms of viral replication and transcription, there is a need to systematically examine possible intraviral interactions. Although relatively unexplored among the virological studies, intraviral interactions have significant and unexpected effects on host range, transmissibility, immunology, and therapeutic effectiveness (Dao et al., 2020; Häsuer et al., 2012; Lee et al., 2011; Yin et al., 2019). We therefore established an interaction map for RTC-related viral proteins by CoPIC matrix screening and identified a total of 47 binary interactions among 14 proteins governing replication, discontinuous transcription, and translation of coronaviruses. Moreover, comprehensive high-order complexes within the SARS-CoV-2 proteome were also comprehensively deduced. Overall, increased awareness of intraviral interactions as described herein is a necessary step for achieving a better understanding of SARS-CoV-2 and thus providing more effective therapeutic opportunities.

RESULTS
Establishment of robust scaffolds for phase separation in cells
The design strategy of CoPIC is as follows: (1) a system (called the scaffold) capable of forming membraneless phase-separated compartments in cells is generated; (2) one component of the PPI of interest (POI) is fused with the scaffold and hence is enriched in the compartments; and (3) the recruitment of another component (called the client) into the compartments serves as the readout for the interaction (Figure 1A). For simplicity, we chose phase separation-prone low-complexity domains (LCDs) as potential scaffolds to form the compartmental hubs for interactions (Malinovska et al., 2013; Wang et al., 2018; Weber and Brangwynne, 2012). To this end, we generated a series of expression vectors with a green fluorescent protein (GFP) fused with a number of LCD candidates (Figure 1B), derived from Nup98 (Schmidt and Görlich, 2015), FIB1 (Berry et al., 2015), hnRNPD (Batlle et al., 2020), RBM14 (Henning et al., 2015), TAF15 (Couthouis et al., 2011), TDP-43 (Saini and Chauhan, 2011), CPEB2N (Kato et al., 2012), and HnfPH1 (Wang et al., 2018). In addition, several optimized LCD combinations were also designed, including Hfq-GFP-FUSN, in which the weak multivalent interactions of FUSN to phase separate are enhanced by the homo-hexamer of Hfq (Murakami et al., 2015; Someya et al., 2012), and DDX4GFP, of which the DEAD-box helicase domain within DDX4 was replaced by GFP as previously described with modifications (Nott et al., 2015).

As shown by the fluorescent images in Figures 1C and S1A–S1G, GFP-Nup98N, DDX4GFP, or Hfq-GFP-FUSN formed green micrometer-scale membraneless spherical compartments while others exhibited poor performance of either irregularly shaped...
condensates or few condensates. Further analyses indicated
higher sphericity of compartments of GFP-Nup98N and Hfq-GFP-FUSN (Figure S1H) while there was a higher proportion of spherical compartments in DDX4GFP (Figure S1I). All compartments formed by GFP-Nup98N, DDX4GFP, or Hfq-GFP-FUSN showed fluidity when examined by fluorescence recovery after photobleaching (FRAP) assays (Figures S1D–S1F). Collectively, we chose GFP-Nup98N, DDX4GFP, and Hfq-GFP-FUSN as potential protein scaffolds of CoPIC for further investigation.

**Nup98N condensates compartmentalize client PPI-mediated recruitment**

To investigate whether the aforementioned scaffold-derived compartments support client PPI-mediated recruitment, we chose a well-known PPI pair, MDM2/p53 (Momand et al., 1992), as an example for the test. First, we fused MDM2 to the C terminus of GFP-Nup98N, DDX4GFP, or Hfq-GFP-FUSN, yielding the scaffolding vectors GFP-Nup98N-MDM2, DDX4GFP-MDM2, or Hfq-GFP-FUSN-MDM2, respectively. Among the constructs, GFP-Nup98N-MDM2 and Hfq-GFP-FUSN-MDM2 formed robust compartments upon overexpression in HEK293 cells (Figures 1D and S1J), while DDX4GFP-MDM2 failed to form dynamic phase-separated compartments (Figure S1L). When GFP-Nup98N-MDM2 was co-expressed with the transactivation helix of p53 (15–29 aa, abbreviated to p53 hereafter) labeled with mCherry (mCherry-p53) in cells, the mCherry red fluorescence signal was enriched within the green compartments (Figures 1D and 1E). Moreover, the enrichment was MDM2-dependent, as mCherry-p53 was not recruited into GFP-Nup98N-derived compartments (Figures 1F and 1G). Similar results were obtained for Hfq-GFP-FUSN-MDM2 and mCherry-p53 (Figures S1J and S1K).

To further confirm that the enrichment of red fluorescence signal was due to the cognate interaction between MDM2 and p53, we treated the cells co-expressing GFP-Nup98N-MDM2 and mCherry-p53 with Mi-773, a potent inhibitor specifically targeting the MDM2/p53 interaction (Wang et al., 2014). Upon treatment with Mi-773, the mCherry signal within the puncta immediately started to decrease, and little remained by 46 s post-treatment (Figure 1H). Correspondingly, the ratio between the intensity of mCherry over that of GFP and Pearson’s correlation coefficient also decreased (Figures 1I and S1M). Notably, the rapid response to Mi-773 suggests the potential application of CoPIC in high-throughput screening of specific inhibitors for PPI, where the effectiveness of inhibitors is evaluated by the recruitment of clients into the compartments or not. Taken together, our results indicate that GFP-Nup98N can yield robust compartments for recruitment-based PPI detection in cells and thus works as the scaffold for engineered membraneless compartments used in CoPIC for subsequent studies.

**Assessment of direct and indirect interactions within the PRC2 complex using CoPIC**

One of the most challenging obstacles in characterizing PPIs is to distinguish direct and indirect interactions. Next, we evaluated whether CoPIC can fulfill the task of discriminating direct or indirect interactions. To this end, we used the well-studied polycomb repressive complex 2 (PRC2) as the testing example. PRC2 is comprised of four core subunits including enhancer of zeste 1 or 2 (EZH1/2), embryonic ectoderm development (EED), suppressor of zeste 12 (SUZ12), and retinoblastoma-binding protein 4 or 7 (RBBP4/7) (Figure S2A) (Margueron and Reinberg, 2011). We generated vectors expressing all PRC2 core components fused with mCherry- or GFP-Nup98N. Based on extensive biochemical and structural studies in the literature, it is known that RBBP4/7 directly interacts with SUZ12 but not with EZH2 (Glancy et al., 2021; Tan et al., 2014). We performed binary co-expression experiments to assess the interaction relationship of these components using CoPIC. Consistently, the co-expression of mCherry-RBBP4 with GFP-Nup98N-SUZ12, but not that of mRFP-RBBP7 with GFP-Nup98N-EZH2, resulted in client enrichment in the compartments (Figures 2A–2D).

Further pairwise interactions between all components of PRC2 were conducted. The results were consistent with the previous studies, where the interactions between EED and EZH1/2, between EZH1/2 and SUZ12, and between SUZ12 and RBBP4/7 were direct and all other interactions were indirect (Figure 2E). As a key component of PRC2, SUZ12 is reported to adopt an extended conformation with the C terminus binding EZH2 and the N terminus binding RBBP4/7 and two accessory proteins, PHF1 and AEBP2 (Figures 2F and 2G). Subsequent CoPIC assays utilizing the C-terminal part of SUZ12, SUZ12C, showed that only EZH2, but not RBBP7, PHF1, or AEBP2, was enriched in the GFP-Nup98N-SUZ12C compartments (Figure S2B). On the contrary, CoPIC assays utilizing the full-length SUZ12 showed substantial enrichment of the signals of RBBP7, PHF1, or AEBP2 within the compartments of GFP-Nup98N-SUZ12 (Figure S2B).

Because we demonstrated that CoPIC can robustly detect direct PPIs, we conducted further investigation into the indirect PPIs within the PRC2 complex. Among the mutual interactions of PRC2 complex components, there are two distinctive interaction patterns: (1) direct contact of RBBP4/7 with SUZ12 mediates the indirect interaction of RBBP4/7 with EZH2; and (2) the
interaction of SUZ12 with EED requires the involvement of EZH2 (Ciferri et al., 2012; Kasinath et al., 2018). Next, we sought to characterize the typical indirect interactions via CoPIC and introduced a ternary co-expression system expressing GFP-Nup98N-EZH2, miRFP-RBBP7, and mCherry-SUZ12. Both miRFP and mCherry signals were enriched and co-localized within the compartments of GFP-Nup98N-EZH2, indicating the bridging role of SUZ12 in recruiting RBBP7 into EZH2-containing compartments (Figures 2H and 2I). On the contrary, the co-localization and enrichment were not observed with co-expression of mCherry-SUZ12C (Figures 2J and 2K). Similar results of CoPIC also verified the indirect interaction of SUZ12 with EED via EZH2 (Figure S2C). Taken together, these data indicate that indirect interactions can be unambiguously demonstrated by combining binary and ternary co-transfection experiments via CoPIC.

**CoPIC screening of direct PPIs among SARS-CoV-2 RTC-related viral proteins**

Taking advantage of the established CoPIC system, we screened PPIs among SARS-CoV-2 RTC-related viral proteins. The genes encoding selected viral proteins (Table S1; Figures S3A) were cloned into the expression vectors fused with fluorescent protein tags such as mCherry, BFP, or GFP-Nup98N. Considering the unique properties of Nsp4 and Nsp6 as transmembrane proteins and the difficulty to synthesize Nsp3, these three Nsps were not included in our analysis. First, the expression and subcellular localization of these proteins was examined in Vero E6 cells, which have been reported to be susceptible to SARS-CoV-2 replication in vitro (Chu et al., 2020). Confocal imaging was carried out to detect the individual proteins at 24 h post-transfection (p.t.). With adequate protein levels expressed in Vero E6 cells, representative images of cellular localization were taken with a confocal microscope. As shown in Figure S3B, most Nsps except Nsp1 and Nsp2 were localized both in the nucleus and cytoplasm. Additionally, the correct sizes of all constructs were further verified by western blot analyses (Figure S3C).

After verifying the proper expression of selected viral proteins, CoPIC screening was conducted for pairwise interactions between all combinations of the 14 proteins. Among the 196 protein pairs tested, a total of 47 interaction combinations were identified as positive and summarized in a pairwise matrix (Figure 3A). In total, 26 interactions showed one directionality, implying a possible influence of fusion domains on the interactions. Nevertheless, 14 pairs of interactions were detected in both directions. In addition, self-interactions were observed for Nsp5, Nsp7, Nsp8, Nsp9, Nsp13, Nsp16, and N protein, suggesting that these proteins could form dimeric or multimeric complexes by interacting with themselves. However, CoPIC failed to detect the previously reported self-interactions for Nsp10 and Nsp15 (Ricagno et al., 2006; Su et al., 2006), a failure that could be due to an adverse effect of partner fusion on the self-interaction. Taken together, the SARS-CoV-2 RTC-related viral factors interactome-based CoPIC screening suggests concentrated interactions of ORF1b-encoded Nsps that were mainly connected by Nsp12, Nsp13, Nsp14, Nsp15, and Nsp16 (Figures 3B and 3C). The second-most connected proteins were Nsp5, Nsp7, Nsp8, and Nsp10, whereas none was identified for Nsp1 and Nsp2 (Figures 3B and 3C).

Although employing a similar expression strategy for replication, 31 interaction pairs in SARS-CoV-2 were identified in addition to 16 overlapped hits shared by SARS-CoV (Figure 3A; Table S2). Among the discovered interaction combination in our study, co-expression of mCherry-Nsp9 and GFP-Nup98N-Nsp5 showed the enrichment of client proteins into the compartments and strong co-localization compared with the negative group (Nsp8-Nsp5) (Figures 4B, 4C, 4F, and 4G). The overlapped positive pairs in SARS-CoV and SARS-CoV-2, i.e., Nsp5-Nsp12 and Nsp9-Nsp9, were also well characterized via CoPIC analyses, implying the possible critical roles for maintaining fundamental functions in the coronavirus family (Figures 4D, 4E, 4H, and 4I). More positive pairwise interactions are exhibited in Figures S4 and S5.

A series of coimmunoprecipitation assays were performed to validate the selected candidates. Briefly, HEK293 cells were co-transfected with two expression plasmids fused with mCherry or GFP tag. At 48 h p.t., cells were lysed and proteins were immunoprecipitated with an anti-GFP monoclonal antibody (mAb). The expression of both tagged proteins was confirmed by western blotting using respective antibodies. As demonstrated in Figures 4J–4M, one of the positive candidate pairs could be immunoprecipitated with another in co-expression samples but not mCherry/GFP-tagged control alone expression samples, further verifying the specific interactions among the above combinations. More pairwise interactions validated by coIP are exhibited in Figure S5B.

**Characterization of in-depth interactions within SARS-CoV-2**

To fulfill a variety of critical functions during the viral life cycle, viral proteins tend to form binary, ternary, or larger complexes with different combinations. Considering that more than half of the interaction pairs turned out to be negative, we speculated that there might be extensive indirect interactions. The corresponding interaction patterns could be discerned, and the detailed strategy of selection is as follows: for virus-encoded proteins, A, B, and C, (1) A interacts with both B and C while B does not interact with C; and (2) it is not until the coexistence of A that B could interact with C, from which we can deduce that A, B, and C are capable of forming a ternary complex (Figures 5A and 5B). Moreover, a large proportion of pairwise interactions with each other within multimeric complexes should also be mentioned (Figures 5A and 5C). Based on the above strategy, we designed the potential combinations and further investigated the detailed patterns of interactions. Among 137 potential protein combinations tested in our study, 31 positive pairs of indirect interactions were eventually identified (Figures 5 and S6). Taking the key pairs of Nsp7-Nsp8 as an example, there are recently published reports of structural analyses underlying the association of this combination with other Nsps, including Nsp12, Nsp9, and Nsp13 (Gao et al., 2020; Hillen et al., 2020; Wang et al., 2020; Yan et al., 2020). However, these interactions were mainly characterized by in vitro assays and have not been tested within mammalian cells. Herein, we performed CoPIC assays combined with coIP, which further supported the existence of high-order complexes.
As shown in Figures S7A and S7B, a combination of CoPIC and coIP analyses showed a validated direct interaction between Nsp8 and Nsp12, Nsp7 and Nsp8, and Nsp7 and Nsp12, thus assembling into the core complex of RTCs to facilitate efficient replication. A ternary complex based on primase (Nsp8) and methyltransferase (Nsp16) was identified, whose formation was dependent on the involvement of Nsp7 (Figures 6A and 6B).

The contrary results were shown in the combination of Nsp10-N-Nsp13, in which Nsp13 failed to mediate the indirect interaction between Nsp10 and N (Figure 6C). Collectively, extensive high-order complexes within SARS-CoV-2 were identified via the unique advantage of the CoPIC system, providing clues for further functional studies.

Validation of intraviral interactions in the context of infection
Although a large number of intraviral interactions were detected for SARS-CoV-2 in virtue of the large-scale screening analysis in the CoPIC system, ectopic expressions in the absence of virus infection make artificial defects inevitable. Moreover, note that almost all of the fusion proteins might lack an authentic N terminus as even C-terminally tagged expression plasmids in our assay would incorporate an N-terminal methionine while these proteins are generated through proteolytic cleavage and will lack methionine during the infection. Therefore, it would be helpful if the identified interactions were validated in the context of SARS-CoV2-infected cells.

The biosafety level-3 (BSL-3) classification of SARS-CoV-2, however, impedes research and antiviral development. Fortunately, the development of transcription and replication-competent SARS-CoV-2 virus-like-particles (trVLPs) (Ju et al., 2021) made it possible to demonstrate the potential interactions of Nsps derived from the proteolytic cleavage of the polyprotein. This trVLP expresses a reporter gene (GFP) replacing the viral N gene, which is required for viral genome packaging and virion assembly (SARS-CoV-2 GFP/$\Delta N$ trVLP). The complete viral life cycle can be achieved and exclusively confined in the cells ectopically expressing SARS-CoV-2 N protein.
Taking advantage of trVLP infection, we analyzed intraviral interaction patterns of N protein with a FLAG tag at the C terminus for detection based on the immunoprecipitation results of infected cell lysates. The precipitate was probed with different antibodies against Nsps. As demonstrated in Figure S7C, N protein precipitated not only a well-known partner Nsp3, but also other viral factors, including Nsp5, Nsp9, and Nsp16, consistent with the above results. Taken together, coIP validation under SARS-CoV-2 trVLP infection further supported the screening results, implying CoPIC as a simple and robust method for investigating PPIs in cells.

Analysis of the interaction hub concentrated on Nsp12-Nsp16
It is well known that ORF1ab of coronavirus is translated upon ribosomal frameshift inside ORF1a, implying significantly lower levels in producing viral proteins from Nsp12 to Nsp16 (Nsp12-Nsp16) than ORF1a-encoded products. Even so, Nsp5, Nsp9, and Nsp16, are thought to serve as a platform to recruit their cofactors (Nsp7 to Nsp10) to the central RTC, thus ensuring efficient replication and transcription. Consistent with this idea, the CoPIC screening demonstrated the concentrated interactions of Nsp12-Nsp16 with their partners (Figures 3, 4, and 5), suggesting the potential roles of intraviral interactions in the sophisticated regulation of enzyme activities.

Taking the canonical RNA-dependent RNA polymerase as an example, Nsp12 exhibits poorly processive RNA synthesis in vitro, contrasting with the efficient replication of coronavirus RNA genomes with a large size in vivo (Ahn et al., 2012; Cheng et al., 2005; te Velthuis et al., 2010). However, the interference of the Nsp7/Nsp8 complex can increase the binding of Nsp12 to RNA, thus activating and conferring the processivity of Nsp12 to the RNA synthesis (Subissi et al., 2014). The Nsp7/Nsp8 complex increases the binding of Nsp12 to RNA, resulting in a larger stretch of nucleotide synthesized per binding event. Nsp8 works as RNA primase and Nsp7 might act to stabilize Nsp8 (Gao et al., 2020; Imbert et al., 2006; Yan et al., 2021; Zhai et al., 2005). Correspondingly, our CoPIC analysis combined with colP confirmed direct interactions between Nsp12 and Nsp7/8 (Figure S7), consistent with the revealed atomic structure (Hillen et al., 2020).

For another presumed RTC catalytic core, Nsp13 helicase activity is also stimulated significantly in the presence of Nsp12 by a direct interaction identified both in SARS-CoV (Adedeji et al., 2012) and our study (Figures 3 and S5A). Cryo-electron microscopy observation revealed an architecture of SARS-CoV-2 mini RTCs assembled by the dynamic interactions of Nsp13 with the RdRp complex (Yan et al., 2020), whose reconstruction in vitro was also partially verified by the CoPIC analyses in the mammalian cells (Figures 6, S6, and S7). Likewise, in vitro studies showed that the extra addition of the Nsp10 protein enhances the weak Nsp14 ExoN activity (Baddock et al., 2020; Bouvet et al., 2012). In the case of Nsp16, a strong interaction of Nsp16 with Nsp10 was shown to trigger its 2'-O-methyltransferase (2'-O-MTase) activity (Bouvet et al., 2010; Imbert et al., 2008; Lin et al., 2020; Mahalapbutr et al., 2020). The direct interactions of both Nsp10-Nsp14 and Nsp10-Nsp16 were also validated by CoPIC (Figures 3 and S4), implying the importance of Nsp10 in two distinct regulatory mechanisms (Nsp14-ExoN and Nsp16-2'-O-MTase). Moreover, the CoPIC screening identified the interaction between Nsp14 and Nsp16 (Figures 3 and S4). There might be a tertiary complex of Nsp14-Nsp10-Nsp16, in which Nsp10-dependent activation of both enzymes may happen simultaneously. Nevertheless, more evidence would be required to confirm the non-limiting levels of Nsp10 to simultaneously activate the enzyme activities of Nsp14 and Nsp16.

Combined with the identified interaction of Nsp12 with Nsp14, an N7-MTase that can methylate the first G of viral RNA (Ferron et al., 2018; Ma et al., 2015; Yan et al., 2021), we surmise that Nsp7/Nsp8/Nsp12/Nsp13/Nsp14 might tend to constitute a higher-order complex as the minimal viral replisome. If so, such assembly would be in charge of more capping activities, i.e., RTPase and N7-MTase, as well as the unwinding of RNA secondary structures, RNA polymerization, and even mismatch excision by the 3'-to-5' exoribonuclease activity of Nsp10-Nsp14. Under the last capping step, the Nsp10/Nsp16 complex would undergo methylation of cap-0 to cap-1 through its 2'-O-MTase activity (Krafcikova et al., 2020; Rosas-Lemus et al., 2020), and it might act periodically during Nsp7/Nsp8/Nsp12/Nsp13/Nsp14-mediated replication. Collectively, our interaction studies confirmed the significance and necessity of the key enzymes encoded by replicase ORF1b forming high-order complexes with the cofactors for efficient replication and transcription during the viral life cycle.

Exploration of potential higher-order complexes
Based on the screening by CoPIC, we constructed a comprehensive high-order interactome for SARS-CoV-2 RTC-related viral proteins (Figure 3). The resulting map reveals a focus on ORF1b-centered interactions that are mainly connected by Nsp12, Nsp13, Nsp14, Nsp15, and Nsp16, with associated enzymatic activities that are essential for viral RNA replication and protein translation. Further extensive interactions combined with the corresponding high-order complexes imply a crucial role for Nsps in recruiting core replication/transcription proteins for the
assembly of the SARS-CoV-2 RTC and thus performing diverse functions during the viral life cycle. Integrated analyses of tertiary complexes identified by CoPIC revealed the various possibilities to trigger indirect interactions to disrupt the intraviral PPIs. Examples include the indirect interaction of Nsp7 with Nsp9 dependent on the involvement of intermediates including Nsp8, Nsp12, Nsp13, or Nsp16. Since the coronavirus RTCs engage in a variety of RNA synthesis processes, one type of complex may be converted into another by the binding or release of specific protein partners. In this manner, such protein factors may either regulate the balance between different processes or directly form more sophisticated complexes with the core enzymes. Correspondingly, we derived several potential higher-order complexes comprising direct and indirect interactions. For example, the tandem mode of Nsp8-Nsp9-Nsp14-Nsp10 illustrates the cascaded linkage of indirect interactions Nsp8-Nsp9-Nsp14 and Nsp9-Nsp14-Nsp10, in which Nsp8 and Nsp10 bind to the different interfaces of the core Nsp9-Nsp14 and do not share any contacts (Figure 7A). Multiple pairs of indirect interactions shared by the same intermediates result in a Y-shaped mode of Nsp8-Nsp10-Nsp13-N (Figure 7B), which emphasizes the mutual linkages of binding factors with intermediates in different interfaces with no interactive events between themselves (Figure 7B). On the contrary, a quinary tandem combination of Nsp7-Nsp8-Nsp9-Nsp14-Nsp10 comprising different intermediates was identified, exhibiting the successive chain (Nsp7-Nsp8-Nsp9, Nsp8-Nsp9-Nsp14, Nsp9-Nsp14-Nsp10) without forming a closed loop by end-to-end interaction (Figure 7C). More detailed high-order complexes are shown in Figure S8.

The above high-order complexes based on CoPIC also exhibit good potential applications in the high-throughput screening for inhibitors against PPIs (Figure 7D). Simply put, the core intermediate (D) fused with the scaffold acts as the platform to recruit other client components (i.e., A to C) by different interfaces and thus form the high-order complex. The subsequent screening for effective inhibitors disrupting the pairwise interactions would be performed in combination with an imaging-based method, and the activity of each candidate would be determined by assessing the enrichment of a fluorophore-labeled client into condensates enriched with its binding partner. Taken together, these simplified model systems enable more precise clues for the formation of high-order complexes during the different stages of the viral life cycle and potential therapeutic opportunities against SARS-CoV-2 by repurposing specific inhibitors to disrupt the intraviral PPIs.

DISCUSSION

With burgeoning knowledge about SARS-CoV-2 molecular architectures, it is essential to dissect the Nsp interaction network and thus understand the assembly of RTCs. In this regard, our study established an innovative interaction screening strategy based on phase separation in cellulo, that is, CoPIC. In the past years, various fluorescence-based methods for in cellulo visualization of protein-protein interactions have been introduced. Although it seems easy for visualization of PPIs to observe the colocalization of proteins tagged with different fluorescence, it is difficult to distinguish whether the proteins in the living cells have real interaction or just incidentally co-localized in the same place. Two relatively popular in vivo techniques for studying PPIs, fluorescence resonance energy transfer (Miyawaki, 2003; Sekar and Periasamy, 2003) and bimolecular fluorescence complementation (Kerppola, 2006), are based on the expression of fluorescently labeled proteins or fragments thereof. However, fluorescence resonance energy transfer requires costly instrumentation and advanced technical expertise, whereas bimolecular fluorescence complementation is based on the irreversible complementation and slow maturation of fluorophores, which does not allow real-time detection of PPIs. Even though alternative strategies based on the relocation of prey proteins have been developed, i.e., mitochondria-docking (Mito-docking) (Shao et al., 2019) and a differential cytolocalization assay (DCLA) (Blanchard et al., 2006), they are unlikely to distinguish between direct protein-protein interactions or those bridged by other peptides.

Since CoPIC detects the enrichment of client proteins in membraneless compartments versus the surrounding environment, discernible enrichment means the establishment of a concentration gradient. Therefore, only authentic interactions can overcome the associated entropy loss. As such, detection of nonspecific PPIs, i.e., false-positives, can be minimized. Nevertheless, if the binding partners to be recruited prefer the specific scaffold-derived condensates versus the surrounding environments, false-positive results can be generated. However, with proper controls, such false-positive hits can be detected, and a change of scaffold can be exploited to circumvent this issue. There is another caveat associated with CoPIC. If the partner to be recruited can also undergo phase separation, the potentially complicated interaction relationship between the scaffold phase and the client phase might hinder the straightforward recruitment of the clients. For example, for the PPI screening involving the N protein, there were hardly any positive interactions detected except for its self-association when N protein serves as the client; however, multiple PPIs were detected when N was fused with the scaffold, GFP-Nup98N (Figure 3A). We reasoned that the phase separation property of N protein (Iserman et al., 2020; Perdikari et al., 2020; Zhao et al., 2020) interferes with its recruitment into the phase-separated compartment of the scaffolds. To circumvent this caveat, binding partners with phase separation propensity should be fused with...
the scaffold, and those without phase separation propensity should serve as the clients.

In previous studies with the yeast two-hybrid system for identifying PPIs within SARS-CoV (Imbert et al., 2008; von Brunn et al., 2007), Nsp2 was shown to have a wide range of interactions with other viral proteins, including Nsp3N, Nsp6, Nsp8, Nsp11, and Nsp16, but these phenomena could not be observed in our CoPIC assays. Eliminating the misgivings on the inappropriate expression (Figure S3C), we surmise that the difference between the current results and previous data can be explained as follows: (1) The absence of protein modifications in the Y2H system. Since protein modifications that are likely to influence protein interactions might be different for certain proteins in the context of yeast and mammalian cells, our CoPIC system might exhibit better performance in reflecting genuine interactions for human viruses. Moreover, the results performed by mammalian two-hybrid assays against SARS-CoV (Pan et al., 2008) seemed not to have any overlap with those results from the Y2H system. (2) The poor reproducibility of the Y2H system. It was striking that none of the interactions revealed in the two independent studies (Imbert et al., 2008; von Brunn et al., 2007) was overlapping, although both used the Y2H system for the screening. (3) The potential interference of Nsp2 with the intracellular milieu. It has been shown that Nsp2 interacts with two host proteins, prohibitin 1 (PHB1) and PHB2 (Cornillez-Ty et al., 2009; Davies et al., 2020), which have been implicated in cellular functions, including cell-cycle progression, cell migration, cellular differentiation, apoptosis, and mitochondrial biogenesis. More notably, continuous investigations on host interactors for SARS-CoV-2 proteins further profile eukaryotic translation initiation factor 4E family member 2 (EIF4E2 or 4EHP) and GRB10 interacting GYF protein (Figure 3, S4B, and S4C). Notably, CoPIC allows the compartmentalization of multiple PPIs simultaneously, which is hard to achieve by other approaches and is important for the dissection of direct and indirect PPIs, hence the potential high-order complexes (Watanabe et al., 2017; Zhang et al., 2018). Better yet, an extra benefit is that CoPIC is especially suitable for PPI detection in a high-throughput fashion using an imaging-based method, e.g., high-content imaging techniques.

When applied to investigate the complicated interaction within the SARS-CoV-2 proteome, the availability of CoPIC allowed the revelation of several salient findings: (1) SARS-CoV-2 shares conserved interactions with SARS-CoV and might have also evolved more interactions to facilitate its efficient replication/transcription; (2) most of the 16 NspS are associated with viral RTC; and (3) the interactions are centered on ORF1b-encoded NspS that were mainly connected by Nsp12, Nsp13, Nsp14, and Nsp16.

Although utilizing a similar expression strategy for efficient replication/transcription, SARS-CoV-2 exhibits a quite divergent pattern from other coronaviruses, sharing only a limited number of homologs (Figure 4A). By comparative analyses of the interactions between the two subspecies, SARS-CoV and SARS-CoV-2, in combination with independently published data (Li et al., 2021; Pan et al., 2008; von Brunn et al., 2007), the conserved interactions can be classified in the following categories: (1) complex formation among the scaffold proteins, where Nsp7 and Nsp8 support the RNA-dependent RNA polymerase function of nsp12, and nsp10 facilitates the methyltransferase activities of nsp14 and nsp16 by the respective associations; (2) interaction between RNA-dependent RNA polymerase (Nsp12) and other key enzymes, including primase (Nsp8), helicase (Nsp13), and guanine-N^1-MTase (Nsp14); (3) a connection of the main protease (Nsp5) to RdRp (Nsp12) and guanine-N^2-MTase (Nsp14); and (4) self-interaction of the accessory proteins such as Nsp5, Nsp7, Nsp8, Nsp9, Nsp13, and Nsp16, and structural protein N. These conserved interactions highlight the evolutionary convergence on the replication strategies at the protein-protein interaction level and point to its potentially critical role in regulating viral replication.

It is striking to identify 31 interactions among RTC-related factors in our study, several of which were confirmed using coIP (Figures 4 and 5). However, note that the negative results may be due to the limitations of the assay itself and do not mean that the interaction cannot occur during infection. Of these, Nsp1 and Nsp2 have the fewest binding partners, which might result from their effect on host viability and suppression of protein synthesis (Angeletti et al., 2020; Gamashova et al., 2006; Rao et al., 2020; Yuan et al., 2020). Among the identified interactions, perhaps the most interesting are those taking place between ORF1b-encoded core enzymes (Nsp12, Nsp13, Nsp14, Nsp15, and Nsp16), responsible for the efficient genome replication and transcription. Compared with other well-known core enzymes encoded by SARS-CoV-2, the role of Nsp15 in the viral lifecycle is still unknown. Nsp15, also named NendoU (nidovirus endoribonuclease, specific for U), preferentially cleaves the 3′ end of U and forms 2′-3′ cyclic phosphate ends (Kim et al., 2020). Previous reverse genetics analyses suggested the dispensable role of NendoU activity during viral replication (Ivanov et al., 2004). Mutations of the catalytic site with His and Lys residues in MHV were shown to promote a subtle defect in subgenomic mRNA accumulation, while mutation of a conserved Asp residue both in HCoV-229E and MHV almost abolished viral RNA replication (Ivanov et al., 2004; Kang et al., 2007).

**Figure 6.** Characterization of representative indirect interactions of Nsp7-Nsp8-Nsp16 and Nsp10-N-Nsp13
(A) CoPIC analysis of the pairwise interactions within the positive combinations of Nsp7-Nsp8-Nsp16. Scale bars, 5 μm. All assays were performed in triplicate.
(B) Fluorescent intensity profiles of the lines with white arrows from (A).
(C) CoPIC analysis of the pairwise interactions within the negative combinations of Nsp10-N-Nsp13. Scale bars, 5 μm. All assays were performed in triplicate.
Figure 7. The outlook for higher-order complexes and a potential HTS scheme for PPI inhibitors
(A–C) Simplified model systems deciphering the potential higher-order complexes.
(D) Schematic diagram of high-throughput screening for specific inhibitors against intraviral complexes.
In addition, Nsp15 might be responsible for evading the host innate immune system, which is also independent of its endonuclease activity (Liu et al., 2019; Zheng and Perlman, 2018). Intriguingly, our CoPIC screening identified multiple binding partners, mainly concentrated on Nsp9, Nsp10, Nsp13, Nsp14, and Nsp16, suggesting the potential involvement of Nsp15 in multiple sophisticated processes. These findings lay the foundation for further functional studies.

In summary, the current work established a sensitive method for identifying PPIs in living cells and thus constructed an interaction network of SARS-CoV-2 RTC-related factors. Among the identified interactions in combination with the data obtained by other methods, the most intriguing and striking are those between the viral core enzymes and the corresponding high-order complexes. While the functional importance of most intraviral PPIs remains less well understood, our network provides a powerful resource to dissect their roles in mediating the genome replication/transcription processes of SARS-CoV-2. The extensive interaction landscape among intraviral proteomes seems to be the norm (Bartel et al., 1996; Choi et al., 2000; Dimitrova et al., 2003; Guo et al., 2001; McCraith et al., 2000; Zell et al., 2005), underscoring the crucial roles of PPIs in the viral life cycle. Hence, the disruption of these intraviral PPIs would likely provide therapeutic opportunities against SARS-CoV-2. Notably, we have been already actively engaging in high-throughput screening for drugs disrupting essential PPIs within SARS-CoV-2 using CoPIC.

**STAR METHODS**

**Detailed methods are provided in the online version of this paper and include the following:**

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**SUPPLEMENTAL INFORMATION**

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**AUTHOR CONTRIBUTIONS**

Conceptualization, P.L.; methodology, H.L. and J.W.; investigation, W.X., and G.P.; writing – original draft, W.X. and H.L.; writing – review & editing, G.P., J.W., X.J., Q.D., and P.L.; funding acquisition, P.L. and H.L.

**DECLARATION OF INTERESTS**

The authors declare no competing interests.

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## KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| mCherry (E5D8F) Rabbit mAb | Cell Signaling Technology | Cat #43590; RRID: AB_2799246 |
| GFP (D5.1) Rabbit mAb | Cell Signaling Technology | Cat #2956; RRID: AB_1196615 |
| DYKDDDDK Tag (D6W5B) Rabbit mAb | Cell Signaling Technology | Cat #14793; RRID: AB_2572291 |
| SARS-CoV-2 NSP3 Rabbit pAb | ABclonal Technology | Cat #A20236, RRID: AB_2891174 |
| SARS-CoV-2 3CLpro Rabbit pAb | ABclonal Technology | Cat #A20198, RRID: AB_2891175 |
| SARS-CoV-2 NSP9 Rabbit pAb | ABclonal Technology | Cat #A20308, RRID: AB_2891176 |
| SARS-CoV-2 NSP16 Rabbit pAb | ABclonal Technology | Cat #A20283, RRID: AB_2891177 |
| F(ab')2-Goat anti-Rabbit IgG (H+L) Cross-Adsorbed Secondary Antibody, HRP | Thermo Fisher Scientific | Cat #A24537; RRID: AB_2536005 |
| **Bacterial and virus strains** |        |            |
| *Escherichia coli* DH5α | Tiangen Biotech | CB101-02 |
| **Chemicals, peptides, and recombinant proteins** |        |            |
| Kanamycin | Inalco | Cat #1758-9316; CAS: 70560-51-9 |
| Ampicillin | Inalco | Cat #1758-9314; CAS: 69-52-3 |
| Phenylmethylsulfonyl fluoride (PMSF) | Amresco | Cat #0754; CAS: 329-98-6 |
| PBS | Solarbio | Cat P1020 |
| 4% Paraformaldehyde | Beyotime Biotechnology | Cat #P0099; CAS: 30525-89-4 |
| Hoechst 33258 | Beyotime Biotechnology | Cat #C1017; CAS: 23491-45-4 |
| DMEM | Hyclone | Cat #SH30022.01 |
| FBS | GIBCO | Cat #10099-141 |
| GFP-Trap Magnetic Agarose | Chromotek | Cat #gtma-100 |
| Protease inhibitor cocktail | Bimake | Cat #B14002 |
| MI-773 (SAR405838) | Selleck | Cat #S7649 |
| FuGENE® HD Transfection Reagent | Promega | Cat #E2311 |
| Rabbit IgG | Beyotime | Cat #A7016 |
| **Critical commercial assays** |        |            |
| Total Protein Extraction Kit for Animal Cultured Cells/Tissues | Invent | Cat #SD-001 |
| BCA Protein Assay Kit | Thermo Fisher Scientific | Cat #23227 |
| Chemiluminescence reagent | Thermo Fisher Scientific | Cat #34077 |
| **Deposited data** |        |            |
| Microscopy images | This paper | https://doi.org/10.17632/gtswbc65nv.2 |

**Experimental models: Cell lines**

|                      |             |            |
|----------------------|-------------|------------|
| Vero E6              | A gift from J.Z. Lab, Tsinghua University | N/A |
| Caco-2-NFlag         | A gift from O.D. Lab, Tsinghua University | N/A |
| HEK293               | Cell Bank/Stem Cell Bank, Chinese Academy of Sciences | Cat #GNHu 43 |

**Oligonucleotides**

|                      |             |            |
|----------------------|-------------|------------|
| GFP-Nup98N (forward primer for PCR) | gtacgctggtctcctggccaaagtct | N/A |
| GFP-Nup98-3.1 (reverse primer for PCR) | ccttccttcctggccaaagtct | N/A |

(Continued on next page)
| REAGENT or RESOURCE |
|---------------------|
| GFP-Nup98-EZH1 (forward primer for PCR) | taagaagaagagagagagaacggaggaggccggctcccatggagaatccacatcatctatagtga |
| N/A |
| GFP-Nup98-EZH1 (reverse primer for PCR) | aacatataataggccctcttagaataagagtagtgctctctctctcat |
| N/A |
| mCherry-EZH1 (forward primer for PCR) | cgggtgtagcggcgccttcctacgactggaaataccaaatcccccta |
| N/A |
| mCherry-EZH1 (reverse primer for PCR) | aacatataagagagccctcttagacttaaaggacgtcgctccctctctccag |
| N/A |
| GFP-Nup98-RBBP4 (forward primer for PCR) | taagaagaagagagagagaacggaggaggccggctcccatggcgagtagcggcggctccatgggagtaaagagatgtt |
| N/A |
| GFP-Nup98-RBBP4 (reverse primer for PCR) | aacatataataggccctcttagagcatgctcctccttagatccttgagtt |
| N/A |
| mCherry-RBBP4 (forward primer for PCR) | cgggtgtagcggcgccttcctacgactggaaataccaaatcccccta |
| N/A |
| mCherry-RBBP4 (reverse primer for PCR) | aacatataataggccctcttagacttaaaggacgtcgctccctctctccag |
| N/A |
| GFP-Nup98-EED (forward primer for PCR) | aagaagagagagagagagagacggaggaggccggctcccatggcgagtagcggcggctccatgggagtaaagagatgtt |
| N/A |
| GFP-Nup98-EED (reverse primer for PCR) | aacatataataggccctcttagagcatgctcctccttagatccttgagtt |
| N/A |
| mCherry-EED (forward primer for PCR) | aagaagagagagagagagagacggaggaggccggctcccatggcgagtagcggcggctccatgggagtaaagagatgtt |
| N/A |
| mCherry-EED (reverse primer for PCR) | aacatataataggccctcttagagcatgctcctccttagatccttgagtt |
| N/A |
| BFP-EED (forward primer for PCR) | aagaagagagagagagagagacggaggaggccggctcccatggcgagtagcggcggctccatgggagtaaagagatgtt |
| N/A |
| BFP-EED (reverse primer for PCR) | aacatataataggccctcttagagcatgctcctccttagatccttgagtt |
| N/A |
| GFP-Nup98-EZH2-3.1 (forward primer for PCR) | gtagcggcggctccctcctctagagatgggcctccatggcgagtagcggcggctccatgggagtaaagagatgtt |
| N/A |
| GFP-Nup98-EZH2-3.1 (reverse primer for PCR) | aacatataataggccctcttagagcatgctcctccttagatccttgagtt |
| N/A |
| mCherry-EZH2-3.1 (forward primer for PCR) | gtagcggcggctccctcctctagagatgggcctccatggcgagtagcggcggctccatgggagtaaagagatgtt |
| N/A |
| mCherry-EZH2-3.1 (reverse primer for PCR) | aacatataataggccctcttagagcatgctcctccttagatccttgagtt |
| N/A |
| GFP-Nup98-RBBP7 (forward primer for PCR) | aagaagagagagagagagagacggaggaggccggctcccatggcgagtagcggcggctccatgggagtaaagagatgtt |
| N/A |
| GFP-Nup98-RBBP7 (reverse primer for PCR) | aacatataataggccctcttagagcatgctcctccttagatccttgagtt |
| N/A |
| MiRFP-RBBP7 (forward primer for PCR) | aagaagagagagagagagagacggaggaggccggctcccatggcgagtagcggcggctccatgggagtaaagagatgtt |
| N/A |

(Continued on next page)
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| MiRFP-RBBP7 (reverse primer for PCR) | atagaatagggccccctcatagcatgttaagatccttgtccctccagttc | N/A |
| GFP-Nup98-SUZ12 (forward primer for PCR) | tggtaagatccttgtccctccagttc | N/A |
| GFP-Nup98-SUZ12 (reverse primer for PCR) | ctatagaatagggccccctcatagcatgttaagatccttgtccctccagttc | N/A |
| mCherry-SUZ12 (forward primer for PCR) | tggtaagatccttgtccctccagttc | N/A |
| mCherry-SUZ12 (reverse primer for PCR) | ctatagaatagggccccctcatagcatgttaagatccttgtccctccagttc | N/A |
| GFP-Nup98-AEBP2 (forward primer for PCR) | tggtagcggcggctccctcgagatggcgcctcagaagcacggcggtgggga | N/A |
| GFP-Nup98-AEBP2 (reverse primer for PCR) | ctatagaatagggccccctcatagcatgttaagatccttgtccctccagttc | N/A |
| mCherry-AEBP2 (forward primer for PCR) | tggtagcggcggctccctcgagatggcgcctcagaagcacggcggtgggga | N/A |
| mCherry-AEBP2 (reverse primer for PCR) | ctatagaatagggccccctcatagcatgttaagatccttgtccctccagttc | N/A |
| GFP-Nup98-PHF1 (forward primer for PCR) | tggtagcggcggctccctcgagatggcgcctcagaagcacggcggtgggga | N/A |
| GFP-NupN-PHF1 (reverse primer for PCR) | ctatagaatagggccccctcatagcatgttaagatccttgtccctccagttc | N/A |
| mCherry-PHF1 (forward primer for PCR) | tggtagcggcggctccctcgagatggcgcctcagaagcacggcggtgggga | N/A |
| mCherry-PHF1 (reverse primer for PCR) | ctatagaatagggccccctcatagcatgttaagatccttgtccctccagttc | N/A |
| GFP-NupN-SUZ12C (forward primer for PCR) | tggtagcggcggctccctcgagatggcgcctcagaagcacggcggtgggga | N/A |
| GFP-NupN-SUZ12C (reverse primer for PCR) | ctatagaatagggccccctcatagcatgttaagatccttgtccctccagttc | N/A |
| mCherry-p53 (forward primer for PCR) | tggtagcggcggctccctcgagatggcgcctcagaagcacggcggtgggga | N/A |
| mCherry-p53 (reverse primer for PCR) | ctatagaatagggccccctcatagcatgttaagatccttgtccctccagttc | N/A |
| DDX4 (forward primer for PCR) | tggtagcggcggctccctcgagatggcgcctcagaagcacggcggtgggga | N/A |
| DDX4GFP 208-251-a (reverse primer for PCR) | ctatagaatagggccccctcatagcatgttaagatccttgtccctccagttc | N/A |
| DDX4GFP 208-251-b (forward primer for PCR) | tggtagcggcggctccctcgagatggcgcctcagaagcacggcggtgggga | N/A |
| DDX4GFP 675-724-a (reverse primer for PCR) | tggtagcggcggctccctcgagatggcgcctcagaagcacggcggtgggga | N/A |
| DDX4 675-724-a (reverse primer for PCR) | tggtagcggcggctccctcgagatggcgcctcagaagcacggcggtgggga | N/A |
| DDX4 675-724-a (reverse primer for PCR) | tggtagcggcggctccctcgagatggcgcctcagaagcacggcggtgggga | N/A |

(Continued on next page)
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| DDX4 675-724-b (reverse primer for PCR) | acactataagggcgccccctagatgcatgtaagctgagatccatcactacatctactgatggagcagtctagtgaagaaatccacagctgtgttcagtaa | N/A |
| GFP-CPEB2 (forward primer for PCR) | gtgtqpccgccccgctctctgaaatgggggcatagctgctgctgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgcttgctg
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| pCDNA3.1-GFP-Nup98N-Nsp1 | This paper | N/A |
| pCDNA3.1-BFP-Nsp1 | This paper | N/A |
| pCDNA3.1-mCherry-Nsp2 | This paper | N/A |
| pCDNA3.1-GFP-Nup98N-Nsp2 | This paper | N/A |
| pCDNA3.1-BFP-Nsp2 | This paper | N/A |
| pCDNA3.1-mCherry-Nsp5 | This paper | N/A |
| pCDNA3.1-GFP-Nup98N-Nsp5 | This paper | N/A |
| pCDNA3.1-BFP-Nsp5 | This paper | N/A |
| pCDNA3.1-mCherry-Nsp7 | This paper | N/A |
| pCDNA3.1-GFP-Nup98N-Nsp7 | This paper | N/A |
| pCDNA3.1-BFP-Nsp7 | This paper | N/A |
| pCDNA3.1-mCherry-Nsp8 | This paper | N/A |
| pCDNA3.1-GFP-Nup98N-Nsp8 | This paper | N/A |
| pCDNA3.1-BFP-Nsp8 | This paper | N/A |
| pCDNA3.1-mCherry-Nsp9 | This paper | N/A |
| pCDNA3.1-GFP-Nup98N-Nsp9 | This paper | N/A |
| pCDNA3.1-BFP-Nsp9 | This paper | N/A |
| pCDNA3.1-mCherry-Nsp10 | This paper | N/A |
| pCDNA3.1-GFP-Nup98N-Nsp10 | This paper | N/A |
| pCDNA3.1-BFP-Nsp10 | This paper | N/A |
| pCDNA3.1-mCherry-Nsp11 | This paper | N/A |
| pCDNA3.1-GFP-Nup98N-Nsp11 | This paper | N/A |
| pCDNA3.1-BFP-Nsp11 | This paper | N/A |
| pCDNA3.1-mCherry-Nsp12 | This paper | N/A |
| pCDNA3.1-GFP-Nup98N-Nsp12 | This paper | N/A |
| pCDNA3.1-BFP-Nsp12 | This paper | N/A |
| pCDNA3.1-mCherry-Nsp13 | This paper | N/A |
| pCDNA3.1-GFP-Nup98N-Nsp13 | This paper | N/A |
| pCDNA3.1-BFP-Nsp13 | This paper | N/A |
| pCDNA3.1-mCherry-Nsp14 | This paper | N/A |
| pCDNA3.1-GFP-Nup98N-Nsp14 | This paper | N/A |
| pCDNA3.1-BFP-Nsp14 | This paper | N/A |
| pCDNA3.1-mCherry-Nsp15 | This paper | N/A |
| pCDNA3.1-GFP-Nup98N-Nsp15 | This paper | N/A |
| pCDNA3.1-BFP-Nsp15 | This paper | N/A |
| pCDNA3.1-mCherry-Nsp16 | This paper | N/A |
| pCDNA3.1-GFP-Nup98N-Nsp16 | This paper | N/A |
| pCDNA3.1-BFP-Nsp16 | This paper | N/A |
| pCDNA3.1-GFP-Nup98N | This paper | N/A |
| pCDNA3.1-DDX4-GFP | This paper | N/A |
| pCDNA3.1-Hfq-GFP-FUSN | This paper | N/A |
| pCDNA3.1-GFP-RBM14LCD | This paper | N/A |
| pCDNA3.1-GFP-TAF15N | This paper | N/A |
| pCDNA3.1-GFP-FIB1N | This paper | N/A |
| pCDNA3.1-GFP-TDP-43C | This paper | N/A |
| pCDNA3.1-GFP-CPEB2N | This paper | N/A |
| pCDNA3.1-GFP-hnRNPDLC | This paper | N/A |
| pCDNA3.1-GFP-hnRNPH1C | This paper | N/A |
| pCDNA3.1-GFP-Nup98N-EZH2 | This paper | N/A |

(Continued on next page)
RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact Pilong Li (pilongli@mail.tsinghua.edu.cn).

Materials availability
This study did not generate new unique reagents. Plasmids generated in this study are available from the Lead Contact without restriction.

Data and code availability
All microscopy image data have been deposited on Mendeley Data via the following link: https://doi.org/10.17632/gtswbcb5nv.2. This study did not generate any unique code. Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Cell lines
HEK293 cells were obtained from Cell Bank/Stem Cell Bank, Chinese Academy of Sciences (Shanghai, China), and Vero E6 cells were kindly provided by Dr. Jianyang Zeng and Dr. Dan Zhao from the Institute for Interdisciplinary Information Sciences (Tsinghua University, Beijing, China). Both HEK293, Vero E6, and Caco-2-NFlag cells were cultured in Dulbecco’s modified Eagle’s medium (HyClone, Inc.) supplemented with 10% fetal bovine serum (Invitrogen, Inc.) at 37°C with 5% CO2. FuGENE HD (Promega, Inc.) was used for transient transfection according to the manufacturer’s instructions.

REAGENT or RESOURCE SOURCE IDENTIFIER

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| pCDNA3.1-mCherry-EZH2 | This paper | N/A |
| pCDNA3.1-GFP-Nup98N-EZH1 | This paper | N/A |
| pCDNA3.1-mCherry-EZH1 | This paper | N/A |
| pCDNA3.1-GFP-Nup98N-EED | This paper | N/A |
| pCDNA3.1-mCherry-EED | This paper | N/A |
| pCDNA3.1-BFP-EED | This paper | N/A |
| pCDNA3.1-GFP-Nup98N-SUZ12 | This paper | N/A |
| pCDNA3.1-mCherry-SUZ12 | This paper | N/A |
| pCDNA3.1-mCherry-SUZ12C | This paper | N/A |
| pCDNA3.1-GFP-Nup98N-RBBP4 | This paper | N/A |
| pCDNA3.1-mCherry-RBBP4 | This paper | N/A |
| pCDNA3.1-GFP-Nup98N-RBBP7 | This paper | N/A |
| pCDNA3.1-mCherry-RBBP7 | This paper | N/A |
| pCDNA3.1-miRFP-RBBP7 | This paper | N/A |
| pCDNA3.1-GFP-Nup98N-MDM2 | This paper | N/A |
| pCDNA3.1-mCherry-p53 | This paper | N/A |

Software and algorithms

NIS-Elements AR Analysis | Nikon Corporation Healthcare Business Unit | https://www.microscope.healthcare.nikon.com/en_AOM/products/software/nis-elements
Adobe Illustrator CC | Adobe Systems | https://www.adobe.com/products/illustrator.html?promoid=RL89NGY7&mv=other
Prism 8 | GraphPad Software | https://www.graphpad.com/scientific-software/prism/
ImageJ | National Institutes of Health | https://imagej.nih.gov/ij/download.html
Cytoscape | Cytoscape Software | https://cytoscape.org/
METHOD DETAILS

Construction of recombinant plasmids
To construct CoPIC-related vectors, the LCD regions, including the N-terminal half of Nup98 (aa 1-515); the N-terminal half of FIB1 (aa 1-82); the N-terminal half of CPEB2 (aa 2-138); the C-terminal half of TDP-43 (aa 258-414); the low complexity region of RBM14 (aa 257-573); the N-terminal half of FUS (aa 1-212); Bacillus subtilis Hfq (BsHfq); full-length (FL) EZH1; FL EZH2; FL SUZ12 (aa 1-739); truncated SUZ12 (aa 561-739); FL EED; FL RBBP4; FL RBBP7; FL PHF1, and truncated AEBP2 (aa 209-503), were amplified using the cDNA derived from HEK293 cells as a template and then cloned into pCDNA3.1 expression vectors. The detailed primer sequences are listed in Key Resources Table. The ORFs of SARS-CoV-2 viral proteins were synthesized de novo and cloned into the pCDNA3.1 plasmid by Synbio Technologies (Monmouth Junction, NJ, USA) (General Biosystems).

Microscopy
Imaging was with a NIKON A1 microscope equipped with a 100 × oil immersion objective. NIS-Elements AR Analysis was used to analyze these images.

Fluorescence recovery after photobleaching (FRAP) measurements
In cellulo FRAP experiments were carried out with a NIKON A1 microscope equipped with a 100 × oil immersion objective. Droplets were bleached with the corresponding laser pulse (3 repeats, 70% intensity, dwell time 1 s). Recovery from photobleaching was recorded for the indicated time.

Validation of expression of fusion proteins
To validate the correct expression of SARS-CoV-2-encoded proteins fused with a mCherry tag, Vero E6 cells transfected with the indicated plasmids were harvested and extracted for lysates using Minute™ Total Protein Extraction Kit for Animal Cultured Cells/Tissues (Invent Biotech) according to the manufacturer’s instructions. The cell lysates were subjected to western blot analyses using the antibody against mCherry (1:1,000 dilution).

Co-immunoprecipitation assays
To confirm the interactions of viral protein pairs, Co-IP assays were performed. HEK293 cells (5 × 10⁷) were co-transfected with the respective expression plasmids. Cells were harvested at 48 h.p.t. and washed twice with chilled phosphate-buffered saline (PBS) supplemented with protease inhibitor cocktail (Bimake, Inc.). Cell lysates were obtained using Minute™ Total Protein Extraction Kit for Animal Cultured Cells/Tissues (Invent Biotech) according to the manufacturer’s instructions. Subsequently, the lysates were incubated with GFP-Trap Magnetic Agarose (ChromoTek, Inc.) at 4°C for 1h, with rotation. After incubation, beads were collected with a Magnetic Separator, and the unbounded samples were removed and the rest saved for analysis. After 5 times washing, the samples were boiled for 10 min and subjected to SDS-polyacrylamide gel electrophoresis (PAGE) and examined by western blot to determine if the mCherry-tagged protein was co-immunoprecipitated with the GFP-tagged protein.

To verify the interactions identified by CoPIC under infection, Caco-2-NFlag cells were infected with recombinant SARS-CoV-2 GFP/ΔN trVLP at an M.O.I of 0.05, harvested at 48 h-post-infection (p.i.), and extracted for lysates using Minute™ Total Protein Extraction Kit for Animal Cultured Cells/Tissues (Invent Biotech) according to the manufacturer’s instructions. One-tenth volumes of the supernatants were preserved as the input controls, and the rest cells were divided into two equal parts and incubated overnight at 4°C with Protein A+G agarose beads (Beyotime Biotech) plus either anti-Flag rabbit mAb or rabbit IgG as a control. The beads were washed four times with lysis buffer and then precipitated for western blot analyses. The primary antibodies used were as follows: anti-Nsp3 (1:1,000 dilution), anti-3CLpro (1:1,000 dilution), anti-Nsp9 (1:1,000 dilution), anti-Nsp16 (1:1,000 dilution), and anti-Flag (1:1,000 dilution).

Western blotting
All antibodies used in the study are summarized in the Key Resources Table. Figures 4J–4M, S3C, S5B, S7B, and S7C were directly analyzed by western blot. The cell samples were resuspended using 2 × Laemmli loading buffer. Proteins were fractionated by SDS-PAGE and transferred to a PVDF membrane. The membranes were incubated overnight with primary antibodies, then with the corresponding peroxide-labeled IgG. Finally, enhanced chemiluminescence reagent was used to visualize the results.

QUANTIFICATION AND STATISTICAL ANALYSIS
Number of biological replicates (BR) is stated in each legend. Error bars represent SD. Graphpad Prism version 8 was used for statistical analysis and graph preparation. For confocal image analysis, NIS-Elements AR Analysis (Nikon, Inc.), ImageJ (National Institutes of Health), and Adobe Illustrator CC (Adobe Systems, Inc.) were used.
Supplemental information

Compartmentalization-aided interaction screening reveals extensive high-order complexes within the SARS-CoV-2 proteome

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Supplemental Figure 1

Figure S1. Characterization of scaffold candidates compartmentalizing and validating the direct interaction between p53 and MDM2 (Related to Figure 1)

(A-G) FRAP analysis of scaffold candidates fused with GFP-tag in HEK293 cells. Scale bar, 5 μm. All assays were performed in triplicates.
**H** Histogram of the circularity of compartments formed by testing seven protein scaffolds in cells. \( n \) (Nup98N) = 100; \( n \) (DDX4GFP) = 182; \( n \) (FUSN) = 73; \( n \) (TAF15N) = 179; \( n \) (FIB1N) = 58; \( n \) (TDP-43C) = 60; \( n \) (RBM14LCD (N)) = 120. Each data point was determined by three independent assays, and error bars represent standard deviations. Values are means ± standard deviation.

**I** Histogram of the area ratio of phase-separated puncta to the nucleus, indicating the phase separation capability of the testing scaffolds. \( n \) = 6. Each data point was determined by three independent assays, and error bars represent standard deviations. Values are means ± standard deviation.

**J** FRAP analysis of DDX4GFP-MDM2 in HEK293 cells. Scale bars, 5 μm. All assays were performed in triplicates.

**K and L** Validation of the direct interaction between Hfq-GFP-FUSN-MDM2 and mCherry-p53 using CoPIC. The p53 fusion protein, as indicated by the mCherry signal, was recruited to the green compartment of Hfq-GFP-FUSN-MDM2 by the specific interaction (K). The co-expression of Hfq-GFP-FUSN and mCherry-p53 served as the control (L). Scale bars, 5 μm. All assays were performed in triplicates.

**M** Plots of Pearson’s correlation coefficient for the intensity of and mCherry-p53 versus GFP-Nup98N-MDM2 pre-treatment and after 46 s. Each data point was determined by three independent assays.
Figure S2. CoPIC analysis of the interaction patterns of SUZ12 with the binding factors and EZH2-mediated indirect interaction of SUZ12 with EED (Related to Figure 2)

(A) Schematic diagram depicting the architecture of the PRC2 complex.
(B) CoPIC analysis of pair-wise interactions between GFP-Nup98N fused SUZ12 (full-length SUZ12) or SUZ12C (C-terminal region of SUZ12) and mCherry-fused binding factors (EZH2, RBBP7, PHF1 and AEBP2). Scale bar, 5 μm. All assays were performed in triplicates.

(C) CoPIC analysis of the indirect interaction of SUZ12 with EED mediated by EZH2. Scale bars, 5 μm. All assays were performed in triplicates.
**Supplemental Figure 3**

Figure S3. Genome annotation and subcellular localization of SARS-CoV-2 RTC-related factors (Related to Table 1 and Figure 3)

(A) SARS-CoV-2 genome annotation on replication/transcription-related factors.

(B) Papain-like protease (PLpro, Nsp3) cleavage sites and main protease (Mpro, Nsp5) cleavage sites.

(C) Molecular weight (kDa) comparison of various SARS-CoV-2 RTC-related factors.
(B) Subcellular localization of SARS-CoV-2 Nsp5 and N protein used in CoPIC screening. Scale bar, 5 μm.

(C) Western blot validation of the constructs fused with an mCherry tag expressed in Vero E6 cells. The cell lysates were extracted and examined with the specific antibody against mCherry.
Figure S4. Positive pairwise (self- and bidirectional-) interactions of SARS-CoV-2 identified by CoPIC screening (Related to Figures 3, 4 and 5)

(A) Collections of positive self-interactions identified by CoPIC screening. All assays were performed in triplicates. Scale bar, 5 μm.

(B) Collections of positive bidirectional interactions identified by CoPIC screening. All assays were performed in triplicates. Scale bar, 5 μm.
Figure S5. Positive pairwise (unidirectional-) interactions of SARS-CoV-2 identified by CoPIC screening (Related to Figures 3, 4 and 5)

(A) Collections of positive unidirectional interactions identified by CoPIC screening. All assays were performed in triplicates. Scale bar, 5 μm.

(B) Co-IP analysis of selected pairwise interactions identified by CoPIC screening.
Figure S6. Collections of positive tertiary complexes identified by CoPIC screening (Related to Figure 6)

Positive tertiary interactions identified by CoPIC screening. All assays were performed in triplicates. Scale bar, 5 μm.
Supplemental Figure 7

(A) CoPIC analysis of binary and tertiary interactions among Nsp7, Nsp8, and NSp12. All assays were performed in triplicates. Scale bar, 5 μm.

(B) CoIP validation of pairwise interactions between Nsp7 and Nsp8, Nsp7 and Nsp12, Nsp8 and Nsp12. All assays were performed in triplicates.

(C) CoIP analysis of Nprotein and binding candidates under SARS-CoV-2 GFP/ΔN trVLP infection in Caco-2-NFlag cells.

Figure S7. Characterization of mutual interactions among Nsp7, Nsp8, and Nsp12 (Related to Figure 6)

(A) CoPIC analysis of binary and tertiary interactions among Nsp7, Nsp8, and NSp12. All assays were performed in triplicates. Scale bar, 5 μm.

(B) CoIP validation of pairwise interactions between Nsp7 and Nsp8, Nsp7 and Nsp12, Nsp8 and Nsp12. All assays were performed in triplicates.

(C) CoIP analysis of Nprotein and binding candidates under SARS-CoV-2 GFP/ΔN trVLP infection in Caco-2-NFlag cells.
Figure S8. Higher-order complexes deduced from CoPIC analysis (Related to Figure 7)

(A) List of quaternary complexes deduced from CoPIC analysis.

(B) List of quinary complexes deduced from CoPIC analysis.

(C) List of six-membered complexes deduced from CoPIC analysis.

(D) List of seven-membered complexes deduced from CoPIC analysis.
Table S1 Sequence information for RTC-related viral proteins of SARS-CoV-2 used in the study for plasmid construction. (Related to Figure 3)

| Protein | Accession Number     | Nucleotide positions | Position in polyprotein | Protein length (aa) |
|---------|----------------------|----------------------|-------------------------|---------------------|
| Nsp1    | YP_009725297.1       | 266-805              | M1-G180                 | 180                 |
| Nsp2    | YP_009725298.1       | 806-2719             | A181-G818               | 638                 |
| Nsp5    | YP_009725301.1       | 10055-10972          | S3264-Q3569             | 306                 |
| Nsp7    | YP_009725303.1       | 11843-12091          | S3860-Q3942             | 83                  |
| Nsp8    | YP_009725304.1       | 12092-12685          | A3943-Q4140             | 198                 |
| Nsp9    | YP_009725305.1       | 12686-13024          | N4141-Q4253             | 113                 |
| Nsp10   | YP_009725306.1       | 13025-13441          | A4254-Q4392             | 139                 |
| Nsp11   | YP_009725312.1       | 13442-13480          | S4393-V4405             | 13                  |
| Nsp12   | YP_009725307.1       | 12480-16236          | V4406-Q5324             | 919                 |
| Nsp13   | YP_009725308.1       | 16237-18039          | A5325-Q5925             | 601                 |
| Nsp14   | YP_009725309.1       | 18040-19620          | A5926-6452              | 527                 |
| Nsp15   | YP_009725310.1       | 19621-20658          | S6453-Q6798             | 346                 |
| Nsp16   | YP_009725311.1       | 20659-21555          | S6799-N7096             | 298                 |
| N protein | YP_009724397.2      | 28274-29533          | --                      | 419                 |
Table S2 Collections of reported interaction patterns of SARS-CoV. (Related to Figure 4)

| Number | Positive interactions       | References               |
|--------|----------------------------|--------------------------|
| 1      | Nsp2-Nsp2                  | (von Brunn et al., 2007) |
| 2      | Nsp2-Nsp7                  | (Pan et al., 2008)       |
| 3      | Nsp2-Nsp8                  | (von Brunn et al., 2007) |
| 4      | Nsp2-Nsp11                 | (von Brunn et al., 2007) |
| 5      | Nsp2-Nsp15                 | (Pan et al., 2008)       |
| 6      | Nsp2-Nsp16                 | (von Brunn et al., 2007) |
| 7      | Nsp5-Nsp5                  | (von Brunn et al., 2007) |
| 8      | Nsp5-Nsp7                  | (von Brunn et al., 2007) |
| 9      | Nsp5-Nsp8                  | (von Brunn et al., 2007) |
| 10     | Nsp5-Nsp12                 | (Pan et al., 2008)       |
| 11     | Nsp5-Nsp14                 | (Pan et al., 2008)       |
| 12     | Nsp7-Nsp7                  | (von Brunn et al., 2007) |
| 13     | Nsp7-Nsp8                  | (von Brunn et al., 2007) |
| 14     | Nsp7-Nsp9                  | (von Brunn et al., 2007) |
| 15     | Nsp7-Nsp13                 | (von Brunn et al., 2007) |
| 16     | Nsp8-Nsp8                  | (von Brunn et al., 2007) |
| 17     | Nsp8-Nsp9                  | (von Brunn et al., 2007) |
| 18     | Nsp8-Nsp12                 | (von Brunn et al., 2007) |
| 19     | Nsp8-Nsp13                 | (von Brunn et al., 2007) |
| 20     | Nsp8-Nsp14                 | (von Brunn et al., 2007) |
| 21     | Nsp9-Nsp9                  | (Sutton et al., 2004)    |
| 22     | Nsp10-Nsp10                | (Su et al., 2006)        |
| 23     | Nsp10-Nsp14                | (Pan et al., 2008)       |
| 24     | Nsp10-Nsp16                | (Pan et al., 2008)       |
| 25     | Nsp12-Nsp13                | (von Brunn et al., 2007) |
| 26     | Nsp13-Nsp13                | (von Brunn et al., 2007) |
| 27     | Nsp15-Nsp15                | (Joseph et al., 2007)    |
| 28     | N protein-N protein        | (Chen et al., 2007)      |