A proteome-scale map of the SARS-CoV-2–human contactome

Understanding the mechanisms of coronavirus disease 2019 (COVID-19) disease severity to efficiently design therapies for emerging virus variants remains an urgent challenge of the ongoing pandemic. Infection and immune reactions are mediated by direct contacts between viral molecules and the host proteome, and the vast majority of these virus–host contacts (the ‘contactome’) have not been identified. Here, we present a systematic contactome map of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) with the human host encompassing more than 200 binary virus–host and intraviral protein–protein interactions. We find that host proteins genetically associated with comorbidities of severe illness and long COVID are enriched in SARS-CoV-2 targeted network communities. Evaluating contactome-derived hypotheses, we demonstrate that viral NSP14 activates nuclear factor κB (NF-κB)-dependent transcription, even in the presence of cytokine signaling. Moreover, for several tested host proteins, genetic knock-down substantially reduces viral replication. Additionally, we show for USP25 that this effect is phenocopied by the small-molecule inhibitor AZ1. Our results connect viral proteins to human genetic architecture for COVID-19 severity and offer potential therapeutic targets.

Despite over 200,000 SARS-CoV-2 publications in the past two years, fundamental questions remain about the molecular mechanisms of genetic risk factors for severe and fatal COVID-19, the cause of long-persisting disease symptoms (long COVID) and the challenge to identify therapeutic targets. These issues remain urgent in light of incomplete vaccination rates, continuously emerging variants and anticipated future pathogens. Fundamentally, infections are initiated by physical contacts between viral proteins and cellular receptors that set off molecular rearrangements culminating in viral entry and unpacking, followed by cellular reprogramming and host defense response triggering. Each of these steps is mediated by contacts between viral and host molecules that determine functional consequences, including proteolytic cleavage or inflammatory signaling, and ultimately clinical manifestations (Fig. 1). Therefore, understanding the mechanisms by which human genetic variation affects COVID-19, as well as the behavior of newly emerging virus variants such as Delta (δ) and Omicron (ο), requires knowledge of these contacts to enable studies on how variants functionally alter virus–host interactions. For SARS-CoV-2, the contacts between the viral spike and human ACE2 proteins are documented by several hundred structures. In contrast, no direct interaction partners are known for many other viral proteins, precluding even domain-level contact models. Because co-complex assays predominantly detect indirect protein-associations, the virus–host contactome remains largely unexplored and unknown. To address this fundamental research gap, we systematically identified protein–protein contacts between SARS-CoV-2 and the human proteome.

Results
SARS-CoV-2–host contactome mapping

We used a multiasay screening and evaluation framework to generate a high-quality virus–host contactome map. To increase detection sensitivity in the initial screening by yeast two hybrid (Y2H), we
(unpublished). Using Y2HHIS3, 26 viral open reading frames (vORFs; Supplementary Table 1) were screened against 17,472 human ORFs (covering 83% of all pairings of human and viral protein-coding genes, that is 83% ‘search space completeness’) in both orientations; that is, as bait and prey (Extended Data Fig. 1a). Human candidate interactors were pairwise retested in triplicate against every vORF, yielding 118 interactions involving 14 viral and 92 human proteins. We refer to this used two complementary assay versions (Extended Data Fig. 1a): (1) a plate-based version using ‘bait’ and ‘prey’ N-terminal fusion proteins encoded on low-copy plasmids and GAL1-HIS3-based growth selection (Y2HHIS3)2,3, and (2) a new system based on the Barcode Fusion Genetics (BFG)-Y2H technology4, using a C-terminal fusion prey protein encoded from a high-copy plasmid and selecting cells expressing green fluorescent protein (GAL1-GFP) from a pooled liquid culture (Y2HGFP).
Y2Hgrp-based human SARS-CoV-2 interactome database as HuSCIgrp. Using Y2Hgrp, 28 vORFs were screened against 14,627 human ORFs (70% completeness) (Extended Data Fig. 1a). After stringent filtering and HIS3-based verification, this yielded 93 interactions involving 13 viral and 84 human host proteins. We refer to this dataset as HuSCIgrp and to the union with HuSCIhms as HuSCI (Supplementary Table 1). We also carried out a targeted screen with previously identified SARS-CoV-1 host interactors; of the 62 testable orthologous SARS-CoV-2-human pairs, six were found to interact (HuSCIorth) (Supplementary Table 2). Y2Hgrp also yielded an intraviral SARS-CoV-2 interactome of 25 binary interactions among 19 vORFs (IntraSCI; Supplementary Table 1). Having collectively identified a contactome of 204 direct virus–host and 25 intraviral interactions among 170 host and 19 viral proteins, we next assessed data quality.

Seven interactions were identified in both HuSCIgrp and HuSCIhms. Albeit nominally low, this overlap is consistent with the complementary nature of the assays and pipelines. Specifically, the screens interrogated incompletely overlapping protein sets and were each 50%–60% saturated. Each version used for screening has an assay sensitivity of 20%–25% (fraction of detectable interactions); thus, the overlap is consistent with known screening parameters1 and a low false-discovery rate. Moreover, from these parameters we can estimate that HuSCI covers 15%–22% of the complete contactome between SARS-CoV-2 and host proteins (Methods).

To further assess data quality experimentally, we compared detection rates of our datasets in the yeast-based nanoluciferase complementation assay (yN2H) to those of established human positive and random reference sets (hsPRS-v2 and hsRRS-v2)2. As additional benchmarks, we derived a set of 55 well-documented binary interactions between human and coronavirus proteins from the curated literature (virus–host interaction binary multiple reference set; vhLit-BM) and a virus–host random reference set (vhRRS) (Supplementary Table 3). We tested HuSCI, IntraSCI and each benchmark set by yN2H (Fig. 1b and Extended Data Fig. 1b). At a stringent scoring threshold of 1% vhRRS, the validation rates of both HuSCI alone and the union of HuSCI with IntraSCI (UnionSCI) were statistically indistinguishable from the two positive control sets (hsPRS-v2, P = 0.76; vhLit-BM, P = 0.06; Fisher’s exact test versus UnionSCI), and each was significantly higher than those of the negative control sets (hsRRS-v2, P = 4 × 10−7; vhRRS, P = 1 × 10−7; Fisher’s exact test versus UnionSCI; Fig. 1b and Supplementary Table 3). Thus, the biophysical quality of our virus–host contactome map is at least on par with high-quality interactions supported by multiple experiments in the curated literature. Although IntraSCI is too small for a separate evaluation by yN2H, 5 of 25 interactions overlap with a previous study (P = 4.6 × 10−5, empirical test; Extended Data Fig. 1c).

The biological relevance of our virus–host contactome map is suggested by the observations that the identified host proteins are enriched for (1) known targets of other viruses (P < 1 × 10−4, empirical test; Fig. 1c), (2) proteins that change phosphorylation status upon SARS-CoV-2 infection (P < 1 × 10−4, empirical test; Extended Data Fig. 1d), and (3) proteins that directly interact with SARS-CoV-2 RNA (P = 0.007, empirical test; Fig. 1d) and (4) proteins that change RNA-binding status during SARS-CoV-2 infection (P = 0.022, empirical test; Extended Data Fig. 1e). These results demonstrate that IntraSCI and HuSCI (Fig. 2a) are of high biophysical quality and enriched for host proteins relevant to SARS-CoV-2 biology.

Complementarity of contactome and co-complex datasets

Previous studies investigating host and SARS-CoV-2 proteins used either affinity purification followed by mass spectrometry (AP-MS) to identify co-complex associations13–15 or biotin identification (BioID) to find proteins in spatial proximity16–18. However, co-complex maps capture largely indirect associations in stable complexes that persist through affinity purification and, likely due to experimental differences, the datasets exhibit limited agreement among each other (Extended Data Fig. 2a). For a subset of such co-complex associations, contacts can be computationally modeled19. In contrast, binary interactome maps provide direct contact partners and are enriched for regulatory interactions6. Despite these differences, 20 of the 204 HuSCI-interacting pairs were found in co-complex and BioID studies, and 38 (34%) of the 170 HuSCI host proteins were associated with a SARS-CoV-2 protein by these studies (Supplementary Table 1). Thus, the contactome map is consistent with previous indirect association datasets while providing substantial novelty.

Although SARS-CoV-2 primarily infects lung and airway tissues, it can spread to additional tissues and this expanded tropism is characteristic for COVID-19 and important for long COVID symptoms20. As previous SARS-CoV-2 interaction datasets could only detect host proteins expressed in the specific assay cell lines, we wondered whether HuSCI was also complementary in terms of the tissue specificity of identified host proteins. Using the Human Protein Atlas (HPA)21, we defined ‘tissue-specific’ and ‘common’ human proteins (Supplementary Table 4). Whereas the AP-MS and BioID data are biased toward common host proteins, HuSCI is more representative of the human proteome and shows good coverage of proteins expressed in the diverse tissues in which SARS-CoV-2 RNA has been detected22 (Fig. 2b, Extended Data Fig. 2b, c and Supplementary Table 4). Thus, the HuSCI contactome has unique advantages for understanding tissue-specific perturbations by SARS-CoV-2.

SARS-CoV-2 targeted functions

To understand which host functions are directly perturbed by the virus, we investigated SARS-CoV-2 targeted pathways. Broad functions enriched among host proteins include (1) immune response, (2) viral process, (3) protein ubiquitination, (4) cytoskeleton and (5) vesicle-mediated transport (Fig. 2c). These largely agree with functions identified in association and proximity datasets13–18 (Supplementary Table 5). Focusing on immune pathways, we noticed that NSP9, NSP14 and NSP16 contact key regulators of cytokine production such as REL (c-REL proto-oncogene, NF-κB subunit), IKBKRG (inhibitor of NF-κB kinase regulatory subunit gamma, also known as IKKγ or NEMO) and TRAF2 (tumor necrosis factor (TNF) receptor-associated factor 2). HuSCI interactors of the membrane-spanning NSP6 were enriched in small boxes according to legend in panel a. Full statistical details are listed in Supplementary Table 5. ER-assoc. ubiq.-dependent prot. cat., Endoplasmic reticulum-associated ubiquitin-dependent protein catabolic. d. Proportion of virus–host interactions in which the human protein has domains that are present in other interactors of the viral protein (shared), not present in other interactors of the viral protein (unique) or no annotated domains (left) and number of shared-domain interactions in HuSCI (arrow) compared to n = 1,000 randomized control networks (gray distribution) (right). One-sided empirical test. Fisher’s exact test with FDR correction. P < 0.001. PPL protein-protein interactions. e. Exemplary ‘shared-domain interaction’ between the viral nucleocapsid protein and four interactors containing a double-stranded RNA binding motif. Domain colors according to legend; gray parts lack domain annotations.
for immune receptors (P < 0.01, empirical test), including CD40 and IL27RA (IL27 receptor subunit alpha). Intriguingly, NSP6 also directly interacts with LY6E, a host restriction factor that limits viral entry for SARS-CoV-2 and other coronaviruses. Several other targets are RNA-binding proteins that function in innate immunity and response to viral infection. MKRN2, together with G3BP1/2, has been suggested...
to regulate olfactory signaling mRNAs\(^{25}\), pointing to potential molecular linkages underlying anosmia in COVID-19. Thus, direct SARS-CoV-2 protein interactors function in immune pathways and viral processes relevant to COVID-19.

Viral proteins contact shared host-protein domains

The restricted size of viral genomes limits their coding potential. We therefore wondered to what extent this limitation yielded viral proteins that bind multiple human proteins via target-shared domains, thus offering opportunities for structure-based drug discovery. We sought domains shared by multiple human targets of each viral protein. In the contactome, SARS-CoV-2 proteins engaged in 43 interactions involving such shared domains (21% of HuSCI; \(P < 0.001\), empirical test; Fig. 2d, Extended Data Fig. 2d and Supplementary Table 6), corresponding to 22 significant virus-protein-domain pairs (\(P < 0.001\), empirical test; Supplementary Table 6). Although the difference was not statistically significant, the 21% proportion of the virus–host contactome with shared-domain interactions was numerically higher than the corresponding 17% in the human reference interactome network (HuRI)\(^{26}\).

Specific examples in HuSCI include four interactors of the nucleocapsid protein sharing the double-stranded RNA-binding motif (\(P < 0.05\), Fisher’s test; Fig. 2e) and the recently confirmed finding that viral nucleocapsid protein interacts with the NTF2 domains of G3BP1 and G3BP2\(^{27}\). Disease-causing mutations are located in the interaction interfaces of the enriched domains of several human proteins (for example, TNF receptor domain of CD40 (ref. \(^{28}\)) or zf-CCCH in MKRN3 (ref. \(^{29}\)). Thus, recurrent structural themes may reflect binding mechanisms that are subject to modulation by human coding variants affecting infection outcome\(^{30,31}\), or by rationally designed drugs.

HuSCI links to COVID-19 risk loci

The severity of COVID-19 symptoms and outcomes are highly variable, and understanding the underlying molecular mechanisms may enable effective treatments. Recently, two independent meta-studies identified genetic loci that are associated with severe COVID-19 illness\(^{22,23}\) (Fig. 3a and Extended Data Fig. 3a), but mechanistic links to viral infection remain unknown. Similarly, several preconditions increase the risk of severe COVID-19, but for these, the molecular links are also poorly understood. At least two models can help to conceptualize how this genetic variation relates to virally targeted host proteins. In a ‘direct’ model, genetic variation in targeted host proteins modulates disease outcome, exemplified by the interaction of adenovirus E1A oncoprotein with the tumor suppressor protein pRb\(^{34}\). In an alternative ‘indirect’ model, genetic variation in the network neighborhood of targeted host proteins modulates the downstream effects and thereby influences disease outcome. A precedent for this model was observed in a plant system, where pathogen-targeted host proteins tend to interact with proteins relevant to disease severity and fitness (encoded by highly variable genes under balancing selection)\(^{35}\). The availability of a high-quality contactome map enabled us to address this fundamental question for COVID-19. Because bias toward well-studied proteins in the SARS-CoV-2 literature\(^{26}\) (Fig. 3b and Extended Data Fig. 3b) limits mechanistic understanding and can cause artifacts, we focused our analyses on systematic protein interaction datasets. The direct model was not supported, given that no targeted host protein from HuSCI was encoded from a critical illness associated locus\(^{22,23}\) (‘critical illness proteins’), and only one (HLA-G, associated with ORF3) was found in a single co-complex study\(^1\). Investigating the indirect model, we sought contacts between targeted host proteins and critical illness proteins, finding 20 (\(P = 0.002\), empirical test; Fig. 3c)\(^{16}\) and 8 (\(P = 0.012\), empirical test; Extended Data Fig. 3a, c)\(^{16}\) in the binary HuRI host network map. In contrast, the virus-associated host-protein sets from AP-MS studies\(^{12,13}\) interact with no more critical illness proteins than expected by chance (Extended Data Fig. 3d). Functionally, the HuSCI host-target proteins linking critical illness to SARS-CoV-2 proteins are enriched in microtubule organization, membrane trafficking and TNF signaling annotations (Supplementary Table 7). Intriguingly, three of seven direct OAS1 interactors are targeted by NSP14 and NSP16, and all three have Golgi- and membrane trafficking-related functions, providing protein contacts that support the finding that the Neanderthal-derived protective OAS1 variant promotes degradation of viral RNA in endoplasmic reticulum- and Golgi-derived virus replication organelles\(^{37}\). These observations indicate that, consistent with the indirect model, clinically relevant genetic variation acts in the local network neighborhood of viral contact proteins.

To further explore the local subnetworks surrounding targeted host proteins and their links to human genetic variation, we identified 204 subnetwork communities in HuRI\(^{26}\) (Fig. 3o) that were significantly targeted by SARS-CoV-2 (nominal \(P < 0.05\), Fisher’s exact test; Supplementary Table 8). Examples include community 28, enriched for ‘negative regulation of viral transcription’ (false discovery rate (FDR) = 0.0018; Fig. 3d) and community 52, enriched for ‘Arp2/3 complex-mediated actin nucleation’ (FDR = 0.0002; Supplementary Table 8). The Arp2/3 complex enables human respiratory viruses to spread among adjacent cells without forming virions\(^{38}\), and ARPC3 scored among the top 50 in two CRISPR screens for SARS-CoV-2 host factors\(^{39,40}\). We then asked whether direct viral target proteins and proteins in each community are encoded by genes associated with human traits of 114 uniformly processed genome-wide association studies (GWAS)\(^{41}\). Variation in genes encoding direct viral targets was only associated with ‘depression’ (FDR = 0.03, MAGMA). In contrast, among the communities, genetic variation associated with severe COVID-19 illness was associated with ten virus-targeted communities, more communities than any other human trait. In contrast, host-protein sets from AP-MS studies were enriched in fewer communities (nominal \(P < 0.05\), Fisher’s exact test; Extended Data Fig. 3e and Supplementary Table 8), and only one host-protein-enriched community each from two AP-MS datasets was enriched for genetic variation associated with severe COVID-19 (refs. \(^{12,13}\); (Lit et al. community 14 and Gordon et al. community 11; Extended Data Fig. 3f). Intriguingly, of the 14 human traits (from 15 studies) associated with 20 additional HuSCI-target-enriched communities, 8 traits (from 9 studies) are clinical risk factors for severe COVID-19 and long COVID, including high body mass index (BMI)\(^{42}\), hypothyroidism\(^{43}\) and schizophrenia\(^{44}\) (\(P = 0.01\), Fisher’s exact test; Fig. 3d, Extended Data Fig. 3e, f and Supplementary Table 8). These links between viral targets and genetic variation associated with COVID-19 comorbidities open the possibility that this genetic variation may impact the course of infection and severity of COVID-19 independent of trait manifestation. Other traits associated with host-target-enriched communities, such as neuroticism, have not been linked to COVID-19 symptoms, possibly because the genetic influence is masked by confounding parameters such as behavior\(^{45}\), and should be considered in the future. Together, these results suggest that the HuSCI contactome map is a powerful and unique resource for studying molecular mechanisms by which human genetics affect the outcome of SARS-CoV-2 infection.

Validation of pathways and host targets

We next explored specific hypotheses for viral proteins and human target functions. Both literature reports and our analyses suggest a role for NF-κB immune signaling in SARS-CoV-2 infection. Because we observed multiple interactions of viral proteins with different members of the NF-κB signaling pathway, we used reporter assays to determine whether and in which direction (that is, activating or inhibiting) viral factors modulate pathway activity. Transfection of NSP14, which interacts with multiple positive NF-κB regulators, resulted in dose-dependent transcriptional activation of NF-κB and even further augmented NF-κB activity following proinflammatory TNFα stimulation in HEK293 cells (Fig. 4a, Extended Data Fig. 4a, b and Supplementary Table 9). This finding suggests that SARS-CoV-2 can induce a proinflammatory state during COVID-19 via direct interaction of NSP14 with NF-κB activators.
Fig. 3 | HuSCI host targets link to genetic variation for severe COVID-19.

a. HuRI interactors (gray nodes) of COVID-19 ‘critical illness proteins’ loci (seed, red nodes) indicate a significant \( (P = 0.0009, \text{empirical test}) \) number of direct SARS-CoV-2 targeted proteins (purple nodes). A total of 144 additional seed protein interactors are not resolved individually. Node and edge colors according to legend. b. Genes in indicated COVID-19 datasets ranked across the human genome by number of publications. Number of publications is indicated by the top panel on log scale. Asterisks indicate significant differences relative to COVID-19-associated genes\(^{30,31}\). NS, \( P = 1 \) (HuSCI) and \( P = 0.36 \) (Stuckalov et al.); *\( P = 0.047; \) ****\( P = 0.00004, \) two-sided Mann–Whitney U test, Bonferroni correction, from top to bottom: \( n = 45, 170, 383, 876 \) and 849; error bars are boxes according to legend (Supplementary Table 8); relation of indicated immune response and viral transcription. Functional descriptors in squared boxes according to legend (Supplementary Table 8); relation of indicated community clusters by shared proteins.

c. Virus interactors enrichment: number of direct SARS-CoV-2 protein interacting in HuSCI interactors in the subnetwork formed by proteins encoded by seed proteins\(^{45,46}\) and their first-level interactors (arrow) compared to \( n = 10,000 \) randomized control networks (gray distribution). d. Of 3,603 communities in Human Reference EntroNoCE (HuRI) with \( n \geq 4 \) members (step 1), 204 are significantly targeted by SARS-CoV-2 (two-sided nominal \( P < 0.05; \) Fisher’s exact test) (step 2); Gene Ontology (GO) enrichment identifies functions associated with each community (step 3); and MAGMA identifies 31 communities significantly associated with human traits (FDR < 0.05) (step 4), the great majority of which are COVID-19 comorbidities. Example community 28 is significantly targeted by SARS-CoV-2 in HuSCI (two-sided \( P = 0.0078; \) Fisher’s exact test, uncorrected) and enriched for negative regulation of adaptive immune response and viral transcription. Functional descriptors in squared boxes according to legend (Supplementary Table 8); relation of indicated traits to COVID-19 is indicated in rightmost column as general link (+) (e.g., via immunity) and clinical evidence for modulation of diseases symptoms and risk for severe or long COVID (+ +; Extended Data Fig. 3f). BMI, body mass index.

These results are corroborated by a study that implicates IMPDH2 in NF-κB pathway activation by NSP14 (ref. \(^{43}\)). Moreover, transcriptional profiling experiments have demonstrated NF-κB activation in HEK293 cells and in patients following SARS-CoV-2 infection\(^{44,45}\). As TNF-α has a central role in the cytokine storm that contributes to many COVID-19 deaths\(^{46}\), the observation that SARS-CoV-2 activates this system in a cell-intrinsic manner may have therapeutic implications.

We explored the role of the NSP14 interactor IKBKG/NEMO, an essential mediator of canonical NF-κB signaling\(^{47,48}\), for transcriptional activation. We generated IKBKG HEK293 knockout (KO) clones (Extended Data Fig. 4) and checked for NF-κB activation in three independent clones after NSP14 transfection (Fig. 4c). IKBKG deficiency abolished NF-κB activation in response to TNF-α and severely impaired NSP14-induced NF-κB activation, providing evidence for a functional role of IKBKG in driving NF-κB activation by NSP14. Interestingly, the residual NF-κB reporter induction upon NSP14 expression in the KO cells indicates that other NSP14 interactors (for example, TRAF2 and REL) contribute to the full NF-κB transcriptional response.

We wondered whether NF-κB signaling proteins and virally targeted host proteins in enriched functional groups other than ‘immune response’ (Fig. 2a) are important for viral replication. After generating A549 alveolar basal epithelial adenocarcinoma cells that exogenously express human ACE2 (A549-ACE2), we quantified viral replication in the presence and absence of CRISPR-Cas9-mediated KO of viral-target encoding genes. Of eight genes that were selected from enriched functional groups and successfully knocked out, deletion of five (63%) resulted in a significant decrease of viral replication (Fig. 4d). Intriguingly, deletion of three NSP14-interacting proteins of the NF-κB signaling system (REL, IKBKG and TRAF2) resulted in strong reduction of viral replication (Fig. 4d and Extended Data Fig. 4f,g). This finding is consistent with a model in which SARS-CoV-2 directly activates NF-κB via NSP14, with this activation being required for successful viral replication. Deletion of kinesin light chain 1 (KLC1), a cargo adaptor protein for microtubule mediated transport, caused reduction of replication by ~80% \( (P < 0.0001, \) Kruskal–Wallis test). Beyond this observation, deletion of ubiquitin-specific peptidase 25 (USP25), which has antiviral functions in influenza and herpes infections\(^{50}\), resulted in essentially complete elimination of viral replication without impacting cell growth, suggesting that human USP25 is required by SARS-CoV-2 (Fig. 4d, Extended Data Fig. 4f,g and Supplementary Table 10).

Inspired by the strong effect on viral replication, we explored USP25 as an antiviral drug target using the small molecule AZ1, which effectively inhibits USP25 and USP28 enzymatic activity\(^{51}\). Using an infectious clone-derived SARS-CoV-2 (iSARS-CoV-2) harboring a mNeonGreen marker\(^{52}\), we showed that treatment with 10 μM AZ1 effectively inhibited SARS-CoV-2 replication in Vero E6 cells (Fig. 4e). Next, we used an independent iSARS-CoV-2 expressing nanoluciferase\(^{53}\) for dose titrations. The AZ1 compound interfered with SARS-CoV-2 replication with half-maximum effective concentration (EC\(_{50}\)) values of 0.8 μM and 0.1 μM in HEK293-ACE2 and Vero E6 cells, respectively (Fig. 4f and Supplementary Table 11), on par with the effects of the clinically approved remdesivir (Extended Data Fig. 4h). Effective concentrations are in the range of the half-maximal inhibitory concentration determined for inhibition of USP25/28 enzymatic activities\(^{54}\), further supporting that USP25 is necessary for SARS-CoV-2 replication. Although the antiviral activity of AZ1 was independently identified in
**Fig. 4 | Validation of pathways and host targets.** a, Relative NF-κB transcriptional reporter activity in unstimulated (left) and TNF-α-stimulated conditions (one-way analysis of variance (ANOVA) with Dunnett’s multiple comparisons test; *P = 0.0395 and ***P = 0.0047, respectively). Error bars represent standard deviation of the mean, n = 3. b, Relative NF-κB transcriptional reporter activity at different amounts of transfected viral protein-encoding plasmid in unstimulated (top) and TNF-α-stimulated conditions (middle) (one-way ANOVA with Dunnett’s multiple comparisons test, **P = 0.0183, ****P < 0.0001 and **P = 0.0012, respectively). Error bars represent standard deviation of the mean, n = 3 (top) and n = 6 (middle). Representative anti-hemagglutinin (HA) western blot demonstrating levels of tagged viral protein in titration experiments relative to actin beta (ACTB) loading controls (bottom). a and b, Precise P values, biological repeats and n for each test are shown in Extended Data Fig. 4 and Supplementary Table 9. c, Relative NF-κB transcriptional reporter activity under unstimulated (left), TNF-α-stimulated (middle) and NSP14-induced conditions in wild-type (WT) and three independent IKBKG KO clones of HEK293 cells (two-way ANOVA with Dunnett’s multiple comparisons test). Error bars represent standard deviation of the mean, n = 3. Precise P values, biological repeats and n for each test are shown in Extended Data Fig. 4, ctrl., control. d, Schematic of viral replication assay (top) and viral replication in wild-type, mock KO and CRISPR KOs of the indicated HuSCI host targets (bottom) (Kruskal–Wallis with Dunn’s multiple comparisons test, *P = 0.031, **P = 0.0047, ***P = 0.0003, ****P < 0.0001, respectively). Error bars represent standard deviation of the mean, n = 9. Precise P values, biological repeats and n for each test are shown in Extended Data Table 4 and Table 10. e, Fluorescence microscopy images showing replication of icSARS-CoV-2-mNeonGreen in infected Vero E6 cells treated with 10 µM AZ1 or solvent (DMSO, dimethylsulfoxide). f, Cell viability and relative replication of icSARS-CoV-2-nanoluciferase in HEK293 cells (left) and Vero E6 cells (right) at different concentrations of AZ1. The EC50 values were calculated with a variable slope model. Error bars represent standard deviation of the mean, n = 8 biological repeats and full analysis in Supplementary Table II.
a small-molecule screen\(^{35}\), our results inform mechanistic studies by identifying NSP16 as a viral interaction partner. NSP16 and associated complexes methylate viral RNA to prevent its detection and destruction by the innate immune system\(^{56,57}\). The stable recruitment of USP25 may protect this complex from ubiquitination and degradation by the host defense machinery. Although elucidating precise mechanisms will require further studies, these findings illustrate the high potential of the HuSCI contactome map in helping to understand and inhibit the SARS-CoV-2 life cycle.

### Perturbed contactome in SARS-CoV-2 variants

Evaluating the impact of novel viral strains on the contactome has been largely restricted to spike protein interactions with ACE2 and antibodies\(^{36}\). Wondering if coding variants in other viral proteins perturb the contactome and thereby modulate viral effects, we explored the potential of 19 SARS-CoV-2 mutations in 14 variants of 9 proteins from the Alpha, Beta, Gamma and Delta strains to alter interactions with host contact targets in HuSCI (Supplementary Table 12). Indeed, some mutations resulted in perturbed interactions. The Alpha strain mutant combination D3L, S23SF in the nucleocapsid protein reduced interaction with ARPC3, the SARS-CoV-2 host factor discussed above. Similarly, the Beta-strain mutation P71L in the envelope (E) protein diminished the interaction with BAG4, an antipapoptotic protein involved in TNF signaling (Extended Data Fig. 5). Although it is currently unknown whether the respective interactions promote viral replication or facilitate immune recognition, the observed changes demonstrate the plasticity of the contactome and, together with recent reports of increased replication of the Delta strain\(^{39}\), strongly suggest that this dimension of viral evolution should also be monitored to assess the risk posed by emerging variants.

### Discussion

In summary, we present a validated contactome map, HuSCI, which provides direct interactions between SARS-CoV-2 and human target proteins in pathways and tissues relevant to COVID-19. HuSCI enables identification of paths of direct contact between viral target proteins and proteins encoded from loci that modify the risk for critical COVID-19 illness and important comorbidities. Examining specific hypotheses for both viral and host proteins, we demonstrate that NSP14 activates the NF-κB pathway even beyond pathway activation by cytokines. Moreover, the majority of the virally targeted host proteins we evaluated, including key NF-κB regulators, are required for efficient SARS-CoV-2 replication. For one of these targeted host proteins, USP25, we confirm that a small-molecule inhibitor can dramatically reduce viral replication and implicate a mechanism for this potential therapeutic. Last, we demonstrate that coding changes in SARS-CoV-2 strains perturb the intracellular interactome. We anticipate that these findings and the contactome resource will stimulate important research toward characterizing new viral strains, understanding the mechanism of COVID-19 symptoms and developing therapies for current and future pandemics.

### Online content

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41587-022-01475-z.

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Methods
Cloning SARS-CoV-2 ORFs
Two independent SARS-CoV-2 vORF collections were constructed in Gateway entry vectors. The Y2HHIS3 collection includes all but one (NSP11) as well as Gateway cloningoptimized ORF of SARS-CoV-2, synthesized based on a published genome, which were cloned with and without stop codon, to enable C-terminal fusions. The Y2HHIS3 entry clone collection is based on National Center for Biotechnology Information (NCBI) accession number NC_045512.2 and annotation. Y2HHIS3 vORFs were synthesized by Twist Bioscience without codon optimization and included 5’ and 3’ linkers with SfiI restriction sites. The 5’ linker incorporates a translational start ATG flanked by BamHI sites; the 3’ linker provides a stop codon flanked by PacI and AsI SI restriction sites. For Y2HHIS3 vORFs were cloned into pENTR223.1 using SfiI restriction cloning, and the alternative ATG was removed by BamHI digest. A total of 28 vORFs were synthesized for Y2HHIS3 and 27 for Y2HHIS3-NSPI-16 (except NSP11). S, E, M, N and ORFs 3A, 3B, 3D, 6, 7A, 7B, 8, 9, 9C and 10-14 (Supplementary Table 1).

Y2HHIS3 vORF entry clones were verified by full-length Sanger sequencing. As NSP10 had a one-base deletion, it was excluded from further experiments. vORFs were moved to the destination vectors pPC86 (N-terminal AD fusion, CEN origin) and pHiDEST-DB (N-terminal DB fusion, CEN origin) by Gateway cloning and confirmed by PCR. For Y2HHIS3, barcoded ‘prey’ (pAR068: C-terminal AD fusion, 2µ origin/pHiDEST-AD: N-terminal AD fusion, CEN origin), and ‘baits’ (pHiDEST-DB: N-terminal DB fusion, CEN origin) destination vectors were generated using published protocols, with the integration of the barcode locus at the SacI restriction site as described. Single barcoded plasmid containing colonies were picked, arrayed into 384-well plates with 80 µl LB agar supplemented with 100 µg/ml carbenicillin and 35 µg/ml chloramphenicol (LB + Carb + CM) per well and incubated at 37°C for 16 h. Barcode sequences were identified using a modified Kiloseq procedure using an Illumina NextSeq 500 and analyzed as previously described. Y2HHIS3 vORFs and human ACE2 were moved by Gateway cloning into barcoded destination plasmids and pHiDEST-AD (N-terminal AD fusion, CEN origin (low copy number)) and pHiDEST-DB (N-terminal DB fusion, CEN origin (low copy number)) such that each ORF was linked to two to six barcodes in every configuration. Gateway cloning was performed individually and for ORF-barcode pairs using Sanger sequencing (TCAG, The Hospital for Sick Children) (Supplementary Table 1).

Generation of HuSCIGFP
Barcoded ORFeomes. The barcoded human ORFeome consisting of 16,747 fully sequence-verified human ORFs with ~95% ORFs represented by two unique barcodes was previously described. The barcoded bait and prey collections were arranged into a 10-by-10 screening matrix consisting of 10 DB and 10 AD groups, each containing ~1,400 ORFs with two distinct sets of unique barcodes, and ~200 ORFs with a single unique barcode set. Barcoded SARS-CoV-2 plasmids were transformed individually into RY3011 (AD plasmids) and RY3031 (DB plasmids) (genotypes in Supplementary Table 1b). Transformed colonies were copied on fresh plates, incubated, scraped off and pooled to make glycerol stocks of all the barcoded SARS-CoV-2 ORFs plus the human ORF ACE2 in each plasmid configuration (with two or more barcodes per ORF).

Mating of pooled haploid yeast. Multiple pooled matings were performed using the frozen haploid pools. Each of the 10 human ORF pools (in C-terminal AD fusion plasmids with 2µ origin; pAR068) were separately mixed with the pool of SARS-CoV-2 ORFs plus human ACE2 in each plasmid configuration (with two or more barcodes per ORF). The model achieved an accuracy >0.9 during all folds of a tenfold cross-fold validation. All positive scores were confirmed by a trained researcher. The verification step was done in triplicate and protein pairs scoring positive in at least two repeats were considered bona fide Y2H interactors. One representative colony of all interaction pairs was picked from selective plates to confirm the identities of X and Y by Sanger sequencing.

Selection of yeast with interacting pair of DB-X and AD-Y by FACS. Pool of glycerol stocks were inoculated into 1-liter flasks with a starting vCFU of 30 M and incubated at 200 rpm for 24 h. Negative controls were started as 10 ml cultures and processed in parallel. ‘Presort’ cultures were prepared for each sample (2 × 10 ml cultures with OD600 10) with doxycycline added (10 µg ml⁻¹) to these cultures to induce barcode swapping while these cultures were incubated for 24 h. To prepare for fluorescence-activated cell sorting (FACS), cells were
concentrated by centrifugation (500 × g, 5 min) and resuspended in PBS to a final OD600 of 10. Propidium iodide (4 mg liter−1) was added to identify dead yeast cells during FACS. Using the diploid negative control, the FACS gate for GFP-positive cells was set to capture 0.1% of GFP-negative cells, yielding a 0.01% false positive rate. Then, 100 million cells per group were sorted, and GFP-positive cells for each sample were plated on 10 SC-Leu-Trp+Ade+10x His (8 mM) plates and incubated for 72 h. Colonies were collected by scraping, centrifuged and resuspended into 2 × 10 ml cultures (OD600 = 10). Doxycycline (10 µg ml−1) was added to induce barcode swapping, and cultures were incubated for 24 h, when plasmid DNA was extracted. Fused barcodes were PCR amplified with primers that attach modified Illumina i5 and i7 adapters to uniquely identify each sample. Following agarose gel analysis of PCR products, the bright band at ~350 bp was purified using a NucleoSpin Gel and PCR Clean-up kit. DNA concentrations were measured for each sample using a Qubit (Invitrogen, Q32851) and, guided by DNA concentration, samples were pooled to ensure equal sequencing depth relative to the number of protein pairs tested.

After primer-dimer removal, DNA was quantified by qPCR, and the pooled NGS library was sequenced on an Illumina NextSeq using a mid- or high-output 150-cycles kit.

Read counting based on expected barcodes. The sequencing data were demultiplexed using bcl2fastaq2 (v2.20.0.422) provided by Illumina with the following command: ’bcl2fastq -r 10 -p 20 -w 10 –no-lane-splitting –barcode-mismatches 1 –adapter-stringency 0.7 –ignore-missing-bcls –ignore-missing-filter –ignore-missing-positions’. After alignments, the fastq files were aligned to the group specific reference files using bowtie2 with the following parameters:

For read 1: q –norc –local –very-sensitive-local –t 23 –redoer.

For read 2: q –nofw –local –very-sensitive-local –t 23 –redoer.

Reference files contained expected barcode sequences for the ORFs in each group. After alignments, reads with mapping quality scores <20 were removed. Following successful BFG barcode recombination, paired-end reads map to up-up or dn-dn when an interaction is present. The number of reads mapping to up-up and dn-dn were counted separately and merged as the final read count. The pipeline was implemented in Python v2.7.

Interaction scoring. For virus–host interactions, we used the product of marginal frequencies of bait and prey strains to estimate the abundance of each diploid bait–prey strain in the presort condition (‘PreSort’). The interaction score was defined by

$$f_{ij} = \frac{f_{preSort}}{f_{AD}}$$

$$f_{preSort} = \sum_{i} \sum_{j} f_{ij}$$

$$f_{AD} = 10^{-4}$$

$$f_{GFP} = \sum_{i} \sum_{j} f_{ij}$$

with the following variables: \(c\), read count; \(i\), AD barcode count; \(j\), DB barcode count; \(f\), frequency.

For every DB barcode, we used the 960 AD null barcodes to define the thresholds leading to a 1% false positive rate. An interaction was accepted as positive only if the ORF pair interaction score was above this threshold for two or more barcode pairs. For intraviral screening, we accepted as interactions those protein pairs for which the frequency of barcode pairs was 1,000 times greater than the median frequency of the corresponding DB barcode for three or more independent barcode pairs, similar to the scoring method previously used for BFG-Y2H with HIS3-based growth selection.

Pairwise retesting. Candidate interaction pairs for HuScLGF were verified in a pairwise HIS3 growth-based Y2H assay as described above (Y2H_HIS3 verification step), with minor modifications. Barcode replicates of candidate human AD-Y and viral DB-X were pooled prior to mating. VORFs NSPI and NSPI2 were omitted from this retesting due to DB autoactivation. After mating, colonies were replica plated on SC-Leu-Trp-His and 3AT-plates. After 72–96 h of yeast growth, these pairwise tests were scored according to the standardized scoring method used for the Y2H_HIS3 screen57. Interaction pairs scoring ≥3 were considered bona fide Y2H interactions.

Estimating completeness using the interactome framework

Assay sensitivity (\(S_a\)) is defined as the fraction of true interactions that can be detected by a given assay. Sampling sensitivity (\(S_s\)) is defined as a fraction of detectable true interactions that can be recovered by the pipeline used. Overall sensitivity of a given screen \(S\) can be calculated as \(S = S_a × S_s\). In pairwise settings, \(S = 1\) and the assay sensitivity is given by the fraction of hsPRS-v1/v2 pairs that score positive. Y2H_HIS3 was benchmarked previously1 and has an assay sensitivity of \(S_{his} = 21.7\%\). Sampling sensitivity of Y2H_HIS3 after two repeats in two orientations has been shown to be \(S_{his} = 60\%\), yielding a spotting sensitivity of \(S_{his} = S_{his} × S_{his} = 0.217 × 0.6 = 13\%\). Given that Y2H_HIS3 screen had a search space completeness of 83\% (\(T_{his} = 83\%\)), the overall completion of HuScCHIS3 is \(C_{his} = T_{his} × S_{his} = 0.83 × 0.13 = 10.8\%\).

A different version of Y2H_GFP using low-copy plasmids and N-terminally fused hybrid proteins (Icn2Y2H_GFP) was benchmarked using 84 pairs of hsPRS-v1 and 92 pairs of hsRVS-v1. Flow cytometry was used to score interactions based on percentage of singlets in GFP-positive gate, which was set using empty bait and prey constructs. In addition, Icn2Y2H_GFP was benchmarked in a pooled setting using all possible combinations of proteins constituting 78 hsPRS-v2 and 77 hsRVS-v2 pairs supplemented with a set of 14 pairs of Y2H-positive controls defined as calibration set1. The experiment was carried out and interactions were scored as described above, except that no empirical null distribution was used. Icn2Y2H_GFP recovered 12 out of 82 (\(S_{lsnGFP} = 15\%\)) hsPRS-v1 pairs when tested in a pairwise single bait–prey configuration and 8 of 92 (\(S_{lsnGFP} = 9/15 = 60\%\)) hsPRS-v2 + calibration set pairs when tested in a pooled single bait–prey configuration, yielding \(S_{lsnGFP} = S_{lsnGFP} × S_{lsnGFP} = 0.15 × 0.6 = 9\%\). It has been previously shown that using high-copy C-terminal fusions increases sensitivity by ~33% without affecting precision2. Thus, screening sensitivity of Y2H_GFP was modeled from that of Icn2Y2H_GFP as \(S_{GFP} = S_{lsnGFP} × 1.33 = 0.33 × 1.33 = 12\%\). Given that Y2H_GFP covered 70\% (\(T_{GFP} = 70\%\)) of all possible virus–human protein combinations, the completion level of the Y2H_GFP dataset is \(C_{GFP} = T_{GFP} × S_{GFP} = 0.70 × 0.12 = 8.4\%\). Only 4 out of 28 (14.2\%) hsPRS-v1 pairs detected by the union of Y2H_BIS and Icn2Y2H_GFP were detected with both methods, indicating a high degree of orthogonality (that is, different detection profiles of the methods used). In addition, Y2H_GFP implemented in this study includes further differences such as high-copy and C-terminal fusion constructs for human proteins. Therefore, we conservatively estimate 90% orthogonality between Y2H_BIS and Y2H_GFP (that is, ~90% of detected interactions are different: \(O_{his-GFP} = 90\%\)). Thus, we estimate that the fraction of all true interactions captured by our merged interactome maps is \(C_{his-GFP} = (C_{his} + C_{GFP}) × O_{his-GFP} ≅ (0.108 + 0.084) × 0.9 = 17.3\%\). Given the uncertainties associated with derivation of screening sensitivity, we estimate lower and higher bounds to be 15% (\(S_{GFP} = 9\%\), excluding inferred gain in sensitivity due to high-copy C-terminal fusions) and 22% \(S_{GFP} = 13.5\%\, S_{his-GFP} = 70\%\) and \(O_{his-GFP} = 100\%\), respectively.

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Pairwise Y2H testing of previously identified SARS-CoV-1 interactions

We identified 97 unique curated binary interactions with SARS-CoV-1 and human interaction partners (Supplementary Table 2). For 77 of these, reagents to test interactions with SARS-CoV-2 orthologues were available in the barcoded human ORFeome. These involved 63 human proteins, 60 of which were covered by two barcode sets and three by a single barcode set. These were tested according to the ‘pairwise retesting’ protocol (above). Successful interactions were indicated by colony growth of both replicates in either condition.

Pairwise Y2H testing with SARS-CoV-2 variants

Lineage-defining mutations for the SARS-CoV-2 ‘variants of concern’ as defined by the Centers for Disease Control and Prevention (Alpha, Beta, Gamma and Delta) were obtained from CoV Spectrum2,3 and mapped to the SARS-CoV-2 reference genome (NCBI accession number NC_045512.2). To generate variant ORFs, Y2Hhuman plasmids were used as template for mutation PCR (primers in Supplementary Table 12). Mutation PCR reaction products were transformed and sequenced verified. Plasmids containing the desired mutation were directly transformed into yeast and processed in pairwise mating as described above. A complete list of mutations generated is shown in Supplementary Table 12. SARS-CoV-2 proteins for which interactions were identified in AD-fusions (N and E) were tested only against the identified interactors. All other variant proteins were tested against all HuSCI interactors. In total, 19 individual mutations in 14 unique variant proteins from 9 different viral proteins were tested. Four proteins with 8 cloned variants had interactors in HuSCI and 1 protein with a single cloned variant had interactors in HuSCI and 4 proteins with 5 variants had no HuSCI interactors.

 γN2H validation

Using Gateway cloning, ORFs from the indicated subsets (Supplementary Table 3) were transferred into pDEST-N2H plasmids (pDEST-N2H-N1, -N2, -C1, and -C2) containing a LEU2 (N1/C1 vectors) or a TRP1 (N2/C2 vectors) auxotrophy marker and transformed into haploid Saccharomyces cerevisiae Y8800 (MATa) and Y8930 (MATa) strains. For cross-plate calibration, two protein pairs from the hsPRS-v2, with different N2H signal intensities, were included in duplicate on every plate (NCBP1/NCBP2 and SKP1/SKP2). Virus–human protein pairs were randomly distributed across the plates and tested together with hsPRS-v2/hsRRS-v2, which were in separate plates.

Overnight-grown haploid cultures were mated by mixing 5 µl of each haploid strain in 160 µl YEPD medium followed by overnight incubation. To measure background, all interactor ORFs were also mated to the viral protein and human domain of interest, and V and D correspond to the viral protein and human domain of interest, and IV and ID to the rest of viral proteins and domains in the HuSCI network, respectively. We identified as enriched associations those with at least 121 vRICs and 294 cRICs were interrogated by our experiment; 5 HuSCI proteins were vRICs, and 13 HuSCI proteins were cRICs. All the observations were tested for enrichment using Fisher’s exact tests and by permutation tests with 10,000 permutations.

GO enrichment analysis

gProfiler4 (database versions: Ensembl104, Ensembl Genomes S1 and Wormbase ParaSite 15) was applied to identify enriched functional categories in HuSCI, AP-MS9,12–15 and BioID studies16–18. The hORFeome9.1, which was used for contactome mapping, served as the background for HuSCI, otherwise the universal annotated human genes. ‘Inferred from electronic annotations’ annotations were excluded. Adjusted Pvalues were calculated using the Benjamini–Hochberg procedure. Functional terms with a hypergeometric P < 0.05 and term size between 5 and 1,000 were collected and enrichment calculated as the ratio between observed and expected gene counts. To categorize HuSCI host proteins, five meta categories inspired by the functional enrichment analysis results were used, namely ‘immune response’ (GO:0006959), ‘viral process’ (GO:0016032), ‘protein ubiquitination’ (GO:0016567), ‘cytoskeleton’ (GO:0005886) and ‘vesicle-mediated transport’ (GO:0016192). Human proteins related to these categories were obtained from the AmiGO 2 (ref. 19) (July 2021), and HuSCI host proteins were categorized based on their annotation to these meta categories.

Domain enrichment of host interacting proteins

Structural domains in human targets were identified from Pfam release 34.0 (ref. 20) (March 2021). Interactions of viral proteins with human interactors that have common domains were defined as shared-domain interactions and counted for HuSCI. The procedure was repeated for 1,000 randomized HuSCI networks (degree-preserved random rewiring). The significance of every viral–human domain was assessed by Fisher’s exact tests (Supplementary Table 6) using the number of V-D, V-ID, IV-D, and IV-D interacting pairs, in which V and D correspond to the viral protein and human domain of interest, and IV and ID to the rest of viral proteins and domains in the HuSCI network, respectively. We identified as enriched associations those with at least two V-D interactions and P < 0.05. We repeated the process for 1,000 randomized HuSCI networks (see above). Multiple domain copies in a given human protein were counted once.

NF-κB reporter assays

HEK293 (RRID: CVCL_0045, DSMZ) were cultured in complete DMEM (high glucose) supplemented with 10% fetal calf serum, 100 U/ml penicillin and 100 µg/ml streptomycin and maintained in humidified atmosphere at 5% CO2 at 37°C. For the reporter assay, 1 × 10^5 HEK293 cells were seeded in a 60-mm cell culture dish one day before transfection. Transfection was done using the calcium phosphate protocol using 10 ng NF-κB reporter plasmid (6 × NF-κB firefly luciferase pGL2), 50 ng pTK reporter (Renilla luciferase) and expression vectors (Flag-IKKβ (pRK5), Flag-A20 (pEF4) and SARS-CoV-2 constructs (pMH)) using a total of up to 6 µg DNA. Briefly, the DNA was diluted in 200 µl 250 mM CaCl2 solution (Carl Roth, 5239.1), vortexed and added dropwise to cell culture dishes. Transfection media was replaced after 6-h incubation with complete DMEM. Then, 24 h after transfection cells were stimulated with 20 ng ml⁻¹ TNF-α for 4 hours. Luciferase activity was measured.
incubated for 4 hours at 37°C and 5% CO2. The lentiviral inoculum was
containing 4.5 × 10⁷ transduction units hACE2 lentivirus per well and
ard media). After 24 h culture medium was replaced by fresh medium
shuffling.

by enriched GO terms in this study were selected: G3BP1, G3BP2, TRAF2,
and anti-HA (Sigma-Aldrich, F3165) were used
3′-CTCCCTTTGTGTTGTGTT-5′) (n = 9, three technical replicates per
biological replicate). GAPDH was used for normalization. Viral RNA
quantified according to the ∆∆Ct standard method⁸¹. The effect of gene
KO on viral replication was determined using the wild-type ORF1a RNA
level as a control as shown in the following equation: 2⁻¹²⁽∆DDCt⁾ = 2⁻¹²⁽∆DCt KO - ∆DCt

Functional effects on viral replication
Selection of host-target candidates. To evaluate if identified host
targets are involved in viral replication, the following HsSCI proteins
involved in host immune regulation and viral life cycle regulation by
enriched GO terms in this study were selected: G3BP1, G3BP2, TRAF2,
USP25, EIF2AK2, REL, IKKBG and KLC1.

Engineering of hACE2-expressing cells. A549 cells were seeded at 5 × 10⁵ cells per well in six-well cell culture plates and cultured in DMEM with 10% FCS and 1 penicillin/streptomycin at 37°C and 5% CO₂ (standard media). After 24 h culture medium was replaced by fresh medium containing 4.5 × 10⁴ transduction units hACE2 lentivirus per well and incubated for 4 hours at 37°C and 5% CO₂. The lentiviral inoculum was then replaced with 2 ml DMEM 10% FCS and 1 penicillin/streptomycin. After 24 h, the transduction was repeated with the same steps as above. Cell surface expression of hACE2 was monitored by FACS using the Attune NxT Flow Cytometer (Thermo Fisher Scientific) and results were analyzed with FlowJo v10 Software (BD Life Sciences). The resulting cells are referred to as A549-hACE2.

Generation of KO cell lines. KO cells were generated using the target-specific CRISPR-Cas9-HDR (homology-directed recombination) KO directed technology developed by Santa Cruz Biotechnology, which enables selection of KO cells with puromycin and red fluorescent protein (Supplementary Table 15). Briefly, A549-hACE2 cells were seeded at 2.5 × 10⁶ cells in T25 flasks and standard media. After 24 h, cells were cotransfected with 7.5 µg each of KO and HDR plasmids for

using the dual luciferase reporter kit (Promega, E1980) according to
the manufacturer’s protocol. The firefly and Renilla lucinescence was
determined with a luminometer (Berthold Centro LB960 microplate
reader, software MikroWin 2010) and quantified in relative light units
(RLU). NF-κB induction was specified as the ratio of firefly luci-
essence (RLU) to Renilla lucinescence (RLU). Significance of relative
NF-κB transcriptional activity was assessed via one-way ANOVA with
Dunnnet’s multiple comparisons. Data evaluation was performed in
GraphPad Prism v7.04.

Protein expression was verified by western blot of lysates. Briefly, proteins were separated by SDS-PAGE and transferred on polyvi-
nylidene fluoride membranes. Membranes were blocked with 5% milk in 1 × PBS + 0.1% Tween-20 (PBS-T) for 1 h at room temperature. Primary antibodies in 2.5% milk in PBS-T were incubated overnight at 4°C, the membranes were washed three times with PBS-T and secondary antibodies were incubated (1.25% milk/PBS-T) for 1 h at room temperature. Anti-actin β (SCBT, sc-47778), anti-FLAG M2 (Sigma-Aldrich, F3165) and anti-HA (Sigma-Aldrich, 11583816001, RRID:AB_2124846) were used.

For generation of IKKBG KO HEK293 cells, oligonucleotides cod-
sing sgRNAs targeting exon 3 (5′-TGCATTTCGAAGCCAGCA-3′) or exon 2 (5′-GCTGCACCATCTCACACAGT-3′) were cloned into pX458 (Addgene, 48138). HEK293 were transfected with 5 µg plas-
mid by standard calcium phosphate transfection. After one day, GFP-positive cells were sorted with a MoFlo cell sorter (Beckman Coulter, Cytomation) and seeded in 96-well plates at dilutions of 0.5–5.0 cells per well. Single-cell clones were expanded and screened for loss of IKKBG expression by western blot (RRID:AB_2128448).

IKKBG-negative clones were verified by amplifying and sequenc-
ing a region of genomic DNA encompassing the sites targeted by
PCR (exon 3: forward primer 5′-CTGGCCCAACAGCTATTTTA-3′, reverse primer 5′-GGTACCGTGAGCAGGCTC-3′; exon 2: for-
ward primer 5′-CTGACATCCTGCTCACAAC-3′ and reverse primer 5′-GGAGCTGAACTGAACCTTCC-3′).

Assessment of SARS-CoV-2 infection in A549-hACE2 KO versus
wild-type cells. Wild-type and KO A549-hACE2 cells were seeded at 1 × 10³ cells per well in 12-well plates and standard media. After 24 h, cells were infected at a multiplicity of infection (MOI) of 10⁻¹, with SARS-CoV-2 isolate hCoV/France/GE1973/2020 (n = 3, biological replicates). Total RNA was extracted from infected cells at 72 h after infection, and SARS-CoV-2 replication was assessed by RT-qPCR using ORF1a primers (5′-ATGACCTTAGTCTGGTCTG-3′; 3′-CTCCCTTTGTGGTTGTGTT-5′) (n = 9, three technical replicates per biological replicate). GAPDH was used for normalization. Viral RNA was quantified according to the ∆∆Ct standard method.⁸¹ The effect of gene KO on viral replication was determined using the wild-type ORF1a RNA level as a control as shown in the following equation: 2⁻¹²⁽∆DDCt⁾ = 2⁻¹²⁽∆DCt KO - ∆DCt

Assessment of the viability of the KO cell lines. A total of 8.0 × 10⁵ cells of each KO cell line were seeded in a white 96-well plate and incubated at 37°C and 5% CO₂ for 24 h. Cell media was replaced with DMEM and incubated at 37°C and 5% CO₂ for 72 h. Cell viability was measured using Cell Titer-Glo Luminescent Cell Viability Assay kit (Promega, G7750). Luminescence was measured on a Centro XS luminometer (Berthold; integration time, 0.5 s). Wild-type cells served as the reference and significance of cell viability was calculated against the mock KO using an ordinary one-way nonparametric ANOVA Kruskal-Wallis with Dunn’s multiple comparisons test using GraphPad Prism v9.

Genes ranked by number of publications
Publication counts are derived from the gene2pubmed file from NCBI, downloaded on 16 November 2021. Only protein-coding genes were considered. For visualization, but not statistical assessment, of genes with equal numbers of publications, order was determined by random

Tissue specificity analysis
The Tissue Atlas dataset was obtained from the HPA database (version 2021.04.09). The HPA categories ‘tissue enriched’, ‘group enriched’ and ‘tissue enriched’ were combined with ‘tissue-specific’, ‘low tissue speci-
city’ was denoted as ‘common’ and the ‘not detected’ category was not
included in this analysis. A total of 11,069 of 19,670 genes (56.3%) in the HPA dataset were defined as tissue specific, and 8,385 of 19,670 genes (42.6%) showed common expression profiles. Tissue distribution differences were determined using Fisher’s exact test with Bonferroni correction.

SARS-CoV-2 organotropism data were obtained from post mor-
tem examinations. The RNA tissue-specific NX value (normalized transcripts per million) was extracted and used to denote whether the gene is specifically expressed in a given tissue. Tissues from the

https://doi.org/10.1038/s41587-022-01475-z
Tissue Atlas were combined into organ systems and used to assess host-target tissues. Significance was evaluated by Fisher’s exact test with Bonferroni correction.

Identification of genetic variation in host targets and network communities

Host network communities were identified using the OCG hierarchical community clustering algorithm on the Human Reference Interactome26, as implemented in the linkcomm R package (V1.0-13) using ‘centered cliques’ as initial class system83. A total of 3,603 communities with a minimum size of 4 were found, of which 204 contained a significant number of virus interactors (that is, were significantly targeted) (nominal P < 0.05, Fisher’s exact test; Supplementary Table 8). A community was annotated to a function if a GO term was enriched (FDR < 0.05) or if ≥20% or ≥30% of the annotated constituent proteins shared an annotation41. From AP-MS-based association studies9–13, 57, 43, 18 and 17 significantly targeted communities were found, respectively (nominal P < 0.05, Fisher’s exact test; Supplementary Table 8).

Uniformly processed GWAS summary statistics were downloaded for 114 traits from the GTEx GWAS analysis41. MAGMA analysis was implemented in R 3.6.1 and consists of three steps: first, GWAS summary statistics across all single-nucleotide polymorphisms (SNPs) within a gene region are aggregated into a gene-level association P value. Next, the gene-level P value is transformed to a z-score (using the inverse normal cumulative distribution function). Finally, z-scores across all genes are modeled as a function of gene set membership and the default gene-level covariates (gene size in number of SNPs, the gene density (a measure of within-gene linkage disequilibrium), the inverse mean minor allele count) using a linear model. Association between gene set membership and GWAS z-scores is tested based on the null hypothesis beta = 0 for the coefficient associated with the gene set membership variable. All targets, and the targeted network communities, were considered gene sets. Entrez gene IDs were used on the human genome assembly 38. Individual MAGMA analyses were performed for each trait based on summary statistics and linkage disequilibrium structure from the 1,000 genomes European reference panel always conditioning on default gene-level covariates (for example, gene length). For each gene set, standard error normalized beta coefficients constituted the association score, with larger values indicating greater chance of getting significant association. Following Benjamin–Hochberg multiple hypothesis correction, gene set–trait associations with FDR < 0.05 were selected. These pairs were subjected to follow-up analysis. SNPs localizing within genes of enriched gene sets were selected, and genes containing SNPs with GWAS P < 5.0 × 10^-8 were selected for the enriched traits, which were considered ‘GWAS hits’. As control the analysis was repeated for the 3,399 network communities that were not significantly targeted (Supplementary Table 8). For both targeted and non-targeted communities the probability of observing traits that are linked to COVID-19 outcomes was assessed. A literature survey identified 33 traits clinically linked to COVID-19 (score 2 in Supplementary Table 8), 18 related to immune function and 61 without connection. For the enrichment analysis we focused on the ‘COVID-linked’ traits; traits related to immune function are also indicated in Fig. 3. Finally, Fisher’s exact test was used to assess the significance traits being linked to COVID-19 (score 2) vs not (scores 0 and 1) in traits that are associated with not-virus-targeted communities (P > 0.5) vs virally targeted communities (P ≤ 0.01). For the control analysis of AP-MS targeted communities, only genetic variation related to COVID-19 severity was evaluated. The contactome-targeted communities with significant GWAS trait associations were numbered 1–31.

Small-molecule inhibition

Remdesivir (Bio-Techne, 7226/10) and USP25/28 inhibitor AZ1 (Bio-Connect, HY-117370-Smg) were dissolved in DMSO. HEK293-ACE2 and Vero E6 (3 × 10^4 cells per well) were plated in white 96-well plates. After 24 h, cells were infected with SARS-CoV-2 (ref. 49) (0.01 MOI) containing a nanoluciferase reporter and treated with the compounds in a 12-point twofold dilution series with 0–10 μM concentration. Each condition was done in triplicate, except for AZ1, which was done in quadruplicate for HEK293-ACE2 and one replicate for Vero E6. Cells were cultured for 24 h, and luminescence was quantified66. Cell viability was measured using the Cell Titer-Glo Luminescent Cell Viability Assay kit (Promega, G7750). EC_50 values were calculated via the variable slope model in GraphPad Prism v9.

Reporting summary

Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The protein–protein interaction (PPI) data from this publication have been submitted to the IMEx (http://www.imexconsortium.org) consortium through IntAct and assigned the identifier IM-28880 (ref. 89). All data from the study are included in the article and associated files. Source data are provided with this paper.

The following data were obtained from the respective original publications: phosphorylation changes upon SARS-CoV-2 infection65; RNA-binding changes upon SARS-CoV-2 infection66; AP-MS virus–host association data: Gordon et al.13, Stukalov et al.41, Li et al.36, Nabeel-Shah et al.41; BioD virus–host proximity data: Laurent et al.41, St-Germain et al.41; Samavarchi-Tehrani et al.41; human expression data: Gordon et al.12, Li et al.14, Nabeel-Shah et al.9; AP-MS virus–host association data: Gordon et al.13, Stukalov et al.41, Li et al.36, Nabeel-Shah et al.41; human expression data: HuRI63; GWAS data for severe COVID-19 illness22,13; and GWAS summary statistics for 114 traits: doi:10.5281/ZENODO.3518299. Interaction data for other viruses were downloaded from IntAct (version: 28 April 2020). Publication counts were downloaded from gene2pubmed (NCBI) on 16 November 2021. Source data are provided with this paper.

Code availability

All source code related to this paper is available via GitHub (https://github.com/INET-HMGU/SARS-CoV-2-contactome)49.

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Competing interests
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Additional information
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Extended Data Fig. 1 | Screening space, orthogonal validation and comparison of HuSCI and IntraSCI to previous SARS-CoV-2 related datasets. 

a, Schematic of the experimental contactome mapping pipelines (left) and screening space for each of the two parallel Y2H$_{\text{His3}}$ and Y2H$_{\text{GFP}}$ screens (right). Proportional overlap is given relative to the union of protein pairs tested by both methods. 

b, Rate at which interactions are detected by yN2H for HuSCI and IntraSCI, as well as positive (hsPRS-v2 and vhLit-BM) and negative (hsRRS-v2 and vhRRS) benchmark sets, across stringency thresholds, error band: standard error of proportion. 

c, Left: overlap of previously identified intraviral SARS-CoV-2 interactions and IntraSCI; right: actual overlap (arrow) compared to n = 10,000 randomized control networks. One-sided, empirical $P = 0.0046$. 

d, Left: overlap of host targets identified in HuSCI and differentially phosphorylated proteins following infection by SARS-CoV-2; right: actual overlap (arrow) compared to n = 10,000 randomized control networks. One-sided, empirical $P < 0.0001$. 

e, Left: overlap of host targets identified in HuSCI and RNA Binding Proteins (RBPs) demonstrating differential RNA binding upon SARS-CoV-2 infection; right: actual overlap (arrow) compared to n = 10,000 randomized control networks. One-sided, empirical $P = 0.022$. 

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Extended Data Fig. 2 | Comparison of HuSCI with SARS-CoV-2 association and proximity datasets. a, Overlap of viral-human protein pairs between HuSCI, four AP-MS and three BioID based datasets (Gordon et al.12,13, Stukalov et al.9, Li et al.14, Nabeel-Shah et al.15, Laurent et al.16, St-Germain et al.17, Samavarchi-Tehrani et al.18). b, Statistical analysis of representation of host targets in common and specific expression groups from datasets in (a), compared to the Human Protein Atlas22 (HPA) (Fisher’s exact test with Bonferroni correction). c, Organotropism analysis across SARS-CoV-2 infected organs from datasets in (b). The percentage of genes within each dataset with specific organotropism (‘tissue-specific’ expression in tissues grouped into organ systems). b and c, Full analysis is shown in Supplementary Table 4. d, Proportion of HuSCI host interactors per SARS-CoV-2 protein in which the human protein has: domains present in other interactors of the viral protein (shared); domains not present in other interactors of the viral protein (unique); no structural domains. Full analysis is shown in Supplementary Table 6.
Extended Data Fig. 3 | See next page for caption.
Extended Data Fig. 3 | Traits associated with COVID-19 severity. a, Table showing COVID-19 critical illness associated loci from two GWAS meta-analyses. Locus-associated proteins present in HuRI are marked in bold. b, Genes in indicated COVID-19 datasets ranked across the human genome by number of publications. Error bars are 95% confidence intervals of the mean, calculated by 1,000 bootstrap samples (from top to bottom n = 45, 170, 383, 876, 29, 75, 25, 71, 233, 46, 49, 45, 15, 33, 10, 71, 97, 47, 58, 9, 46, 22, 20, 23, 39). c, Virus-interactor enrichment in contactome: number of direct SARS-CoV-2 protein interacting HuSCI proteins in a HuRI subnetwork formed by proteins encoded by COVID-19 critical illness associated loci (marked in bold in table (a)) and their first level interactors (arrow) compared to n = 10,000 randomized control networks (gray distribution). One-sided, empirical P = 0.012. d, Virus-interactor enrichment in co-complex associations: number of SARS-CoV-2 associated human proteins in two AP-MS based studies in a subnetwork formed by proteins encoded by COVID-19 critical illness associated loci (marked in bold in table (a)) and their first level interactors (arrow) either in HuRI or BioPlex 3.0. The comparisons are against n = 10,000 randomized control networks (gray distribution). One-sided, empirical P values are shown for each dataset. e, Upset plots showing number of communities targeted by SARS-CoV-2 (left) and associated with severe COVID-19 (right) in HuSCI and AP-MS based datasets. f, Table showing 15 traits for genetic variation identified within targeted network communities. An association with severe COVID-19 comorbidities is indicated, as well as trait references: T2D_UKBS, BMIA, FAT_UKBS, HRET, RET, HC_UKBS, ADPN, HYPOTHY_UKBS, SCZ_UKBS, GIANT_HIP, IBD_UKBS, OST_UKBS, EGG_PHF, GIANT_HEIGHT, NEUROT_UKB. g, Grouping of 31 network communities with significantly associated traits shown in Fig. 3d by protein membership measured by Jaccard similarity according to legend.
a) Summary statistics for unstimulated NF-κB reporter induction.

| Dunnett's multiple comparisons test       | Mean Diff. | 95.00% CI of diff. | Below threshold? | Summary | Adjusted P Value |
|------------------------------------------|------------|-------------------|-----------------|---------|-----------------|
| EV (pMH-P) vs. KO#9                     | 0.8437     | -0.1134 to 1.795  | No              | no      | 0.855           |
| EV (pMH-P) vs. KO#10                    | -0.1776    | -0.5947 to 0.2494 | No              | no      | 0.233           |
| EV (pMH-P) vs. KO#11                    | -0.2871    | -0.7035 to 0.1393 | No              | no      | 0.0206          |
| EV (pMH-P) vs. KO#12                    | 0.0507     | -0.5300 to 0.6314 | No              | no      | 0.9272          |
| EV (pMH-P) vs. KO#13                    | -0.1368    | -0.6870 to 0.4140 | No              | no      | 0.7417          |
| EV (pMH-P) vs. KO#14                    | 0.0636     | -0.5564 to 0.6836 | No              | no      | 0.9319          |
| EV (pMH-P) vs. KO#15                    | -0.1463    | -0.6645 to 0.3720 | No              | no      | 0.9997          |

b) Summary statistics for unstimulated and TNFα-stimulated NF-κB reporter induction.

| Dunnett's multiple comparisons test       | Mean Diff. | 95.00% CI of diff. | Below threshold? | Summary | Adjusted P Value |
|------------------------------------------|------------|-------------------|-----------------|---------|-----------------|
| EV (pMH-P) vs. KO#9                     | 0.8437     | -0.1134 to 1.795  | No              | no      | 0.855           |
| EV (pMH-P) vs. KO#10                    | -0.1776    | -0.5947 to 0.2494 | No              | no      | 0.233           |
| EV (pMH-P) vs. KO#11                    | -0.2871    | -0.7035 to 0.1393 | No              | no      | 0.0206          |
| EV (pMH-P) vs. KO#12                    | 0.0507     | -0.5300 to 0.6314 | No              | no      | 0.9272          |
| EV (pMH-P) vs. KO#13                    | -0.1368    | -0.6870 to 0.4140 | No              | no      | 0.7417          |
| EV (pMH-P) vs. KO#14                    | 0.0636     | -0.5564 to 0.6836 | No              | no      | 0.9319          |
| EV (pMH-P) vs. KO#15                    | -0.1463    | -0.6645 to 0.3720 | No              | no      | 0.9997          |

Extended Data Fig. 4 | See next page for caption.
Extended Data Fig. 4 | Effect of viral proteins on NF-κB reporter activity and of viral interactors on viral replication. a, Tables showing statistical details of NF-κB transcriptional reporter activity in the absence and presence of selected viral proteins under unstimulated (top) and TNFα stimulated (bottom) conditions. One-way ANOVA with Dunnett’s multiple comparisons test, n = 3, adjusted P values are shown. b, Table showing statistical details of NF-κB transcriptional reporter activity at different amounts of transfected viral protein-encoded plasmid under unstimulated (left) and TNFα stimulated conditions (right). One-way ANOVA with Dunnett’s multiple comparisons test, n = 3 and n = 6, respectively, adjusted P values are shown. a and b, Raw data and full analysis is shown in Supplementary Table 9. c, Table showing statistical details of NF-κB transcriptional reporter activity under unstimulated (left), TNFα-stimulated (middle) and NSP14-induced conditions in WT and IKBKG KO HEK293 cells (two-way ANOVA with Dunnett’s multiple comparisons test, n = 3), adjusted P values are shown. d, Representative anti-IKBKG (top) western blot demonstrating levels of IKBKG in WT and three independent IKBKG knockout clones of HEK293 cells relative to actin beta (ACTB) loading controls (bottom). e, Representative anti-hemagglutinin (HA) western blot demonstrating levels of tagged NSP14 protein in NF-κB induction experiments relative to actin beta (ACTB) loading controls (bottom). f, Table showing statistical details of viral replication in wild-type, mock KO and CRISPR KOs of the indicated HuSCI host proteins. Kruskal-Wallis with Dunn’s multiple comparisons test, n = 9. Adjusted P values are shown. g, Cell viability of mock KO and CRISPR KOs of the indicated HuSCI host proteins relative to WT cells. Kruskal-Wallis with Dunn’s multiple comparisons test, n = 3. Adjusted, Fisher’s exact P values are shown. f and g, Raw data, Fisher’s exact P values, and full analysis is shown in Supplementary Table 10. h, Cell viability and relative replication of icSARS-CoV-2-nanoluciferase in HEK293 cells (left) and Vero E6 cells (right) at different concentrations of remdesivir. The EC50 values shown for each cell line were calculated with a variable slope model. Error bars: standard deviation of the mean, n = 3 biological repeats, full analysis in Supplementary Table 11.
Extended Data Fig. 5 | See next page for caption.
Extended Data Fig. 5 | Mutations of SARS-CoV-2 variants affect specific interactions with uSCI host targets. a, Y2H_{HuSCI} yeast growth on selective plates of HuSCI interaction partners as DB-fusion proteins tested against AD-fusion of the SARS-CoV-2 Nucleocapsid (AD-N) protein (Wuhan-Hu1, original screen) and AD-N containing ‘lineage defining’ amino acid substitutions: D3L and S235F (α-strain), T205I (β-strain), or P80R (γ-strain). Shown is one representative result of 5 repeats. b, Y2H_{HuSCI} yeast growth on selective plates of HuSCI interaction partners as DB-fusion proteins tested against AD-fusion of the SARS-CoV-2 Envelope (AD-E) protein (Wuhan-Hu1, original screen) or AD-E containing ‘lineage defining’ substitution P71L (β-strain). Shown is one representative Y2H_{HuSCI} result on selective media, out of 2 repeats. a–c. Black circles indicate changes in yeast colony growth between human proteins tested against viral variant ORFs or the originally screened Wuhan strain ORFs observed consistently across all repeats. c, AD-empty control plate for a, b indicates lack of autoactivation. d, Layout of DB-fusion HuSCI interactors (purple) tested with AD fusion SARS-CoV-2 proteins or AD-empty control, respectively in a–c. N/A indicates human interactors.
Reporting Summary

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Statistics

For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.

☐ The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement

☐ A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly

☐ The statistical test(s) used AND whether they are one- or two-sided

☐ Only common tests should be described solely by name; describe more complex techniques in the Methods section.

☐ A description of all covariates tested

☐ A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons

☐ A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals)

☐ For null hypothesis testing, the test statistic (e.g. F, t, r) with confidence intervals, effect sizes, degrees of freedom and P value noted

☐ Give P values as exact values whenever suitable.

☐ For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings

☐ For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes

☐ Estimates of effect sizes (e.g. Cohen’s d, Pearson’s r), indicating how they were calculated

Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection

| B箭| Bowie2 2.2.3 |
|----|-----------|
| Illumina bc12/fastq 2.20.0.422 |
| Python 2.7.10 |
| Custom code for Y2H screening pipeline |
| Tissue Atlas dataset from the Human Protein Atlas database (version: 2021.04.09) |

Data analysis

| C箭| Cytoscape 3.8.2 |
|----|---------------|
| Matlab 2015a |
| gProfiler (Database versions: Ensembl 104, Ensembl Genomes 51 and Wormbase ParaSite 15) |
| R 3.6.1 |
| linkcomm R package |
| Metascape web-platform |
| Pfam release 34.0 |
| Flowjo Software 10 |
| MASMA v1.09e |
| MikroWin 2010 |
| Graphpad Prism v. 7.04, Graphpad Prism v. 9, Amigo2 |

For manuscripts utilizing custom algorithms or software that are not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Portfolio guidelines for submitting code & software for further information.
Policy information about availability of data
All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:
- Accession codes, unique identifiers, or web links for publicly available datasets
- A description of any restrictions on data availability
- For clinical datasets or third party data, please ensure that the statement adheres to our policy.

The PPI data from this publication have been submitted to the IMEX [http://www.imexconsortium.org] consortium through intAct and assigned the identifier IM-28880104. All data from the study are included in the manuscript and associated files. Source data are provided with this paper. The following data were obtained from the respective original publications: phosphorylation changes upon SARS-CoV-2 infection10,11; RNA-binding changes upon SARS-CoV-2 infection12; AP-MS virus-host association data: Gordon et al.14,15, Stukalov et al.10, Li et al.13, Nabeel-Shah et al.16, BioID virus-host proximity data: Laurent et al.17, St-Germain et al.19 Samavarchi-Tehrani et al.18; Human expression data: Human Proteome Atlas22, SARS-CoV-2 organotropism23,99; human host interactome: HuRI27; GWAS data for severe COVID-19 illness33,34; GWAS summary statistics for 114 traits: doi:10.5281/zenodo.3518299; Interaction data for other viruses were downloaded from IntAct9 (version: April 28, 2020). Publication counts downloaded from gene2pubmed (NCBI), on 2021-11-16.

Field-specific reporting
Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

☑ Life sciences ☐ Behavioural & social sciences ☐ Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see nature.com/documents/nr-reporting-summary-list.pdf

Life sciences study design
All studies must disclose on these points even when the disclosure is negative.

Sample size
Sample size for PPI validation assay experiments was chosen based on a 95% CI value of 5% inputting and an expected recovery rate of 20%, from previous similar experiments, using the standard error of proportion. For IntraSci, the full dataset was tested. For all cell biological assays 3 - 8 biological repeats were done. Many years of experience have convinced us that this produces robust results.

Data exclusions
No data were excluded.

Replication
All attempts at replication were successful. For Y2H, 3 fold-replication is part of the standard quality control pipeline. In all cell biology experiments, i.e. viral replication assays and NF-κB reporter activation assays, biological results were replicated at least twice (3 biological repeats).

Randomization
Randomization for network permutations was done while maintaining network topology (maintaining the number of interactions but shuffling the edges), yN2H validations were done with a fully randomized setup to avoid experimental bias and eliminate covariates, i.e. all pairs, except hsPRSV2/hs/RvRv2, were randomly distributed across the plates. For experimental reasons, and because extensive prior data are available for hsPRSV2/hs/RvRv2 benchmark pairs these were randomly mixed among each other, but separate from other benchmark and test pairs. For all cell culture and validation experiments, all samples were treated by the same researcher in parallel in random order to eliminate covariates. In all experiments, negative controls from the same experimental batch were used as initial reference.

Blinding
Y2H screens were done in a fully random and unbiased manner. Manual scoring of Y2H pairwise tests was blinded. Experimental design of yN2H validation was blinded during the experiment. For cell culture experiments with small sample sizes blinding was not possible, but all experiments are internally controlled to reduce researcher biased experimental differences.

Reporting for specific materials, systems and methods
We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.
Materials & experimental systems

| n/a | Involved in the study |
|-----|------------------------|
| ☐   | Antibodies             |
| ☒   | Eukaryotic cell lines  |
| ☒   | Palaeontology and archaeology |
| ☒   | Animals and other organisms |
| ☒   | Human research participants |
| ☒   | Clinical data          |
| ☒   | Dual use research of concern |

Methods

| n/a | Involved in the study |
|-----|------------------------|
| ☐   | ChiP-seq               |
| ☒   | Flow cytometry         |
| ☒   | MRI-based neuroimaging |

Antibodies

Antibodies used

1. Anti-b-Actin (SCBT, sc-47778, RRID: AB_2714189)
2. anti-FLAG M2 (Sigma Aldrich, F3165, RRID:AB_259529)
3. anti-HA (Sigma-Aldrich, 11583816001, RRID:AB_514505)
4. Secondary antibody (Jackson Immunoresearch, Jim-715-035-150)

Validation

The antibodies have been validated by the manufacturers and were not revalidated in this study.

Eukaryotic cell lines

Policy information about cell lines

Cell line source(s)

1. A549 (purchased from ATCC)
2. Hek293 (RRID: CVCL_0045, purchased from DSMZ)
3. Vero E6 (ATCC CRL-1586), kindly provided by Prof. Nicolas Gillet (University of Namur, Belgium)
4. Yeast strains Y8800 and Y8930: the same strains used in the many previous studies [Itolland, et al. Cell, 2014 Nov 20;159(5):1212-1226;]
5. Yeast strains - RY9011 and RY9031: Derived from Y8800 and Y8930 in Dr. Fritz Roth's lab at the University of Toronto

Authentication

Vero E6 cells were re-authenticated by RNA-Seq. Other cell lines were purchased from either ATCC or DSMZ and not re-authenticated.

Mycoplasma contamination

All cell lines were tested by PCR for mycoplasma contamination and found to be negative

Commonly misidentified lines (See ICCLAC register)

No commonly misidentified cell lines were used.