SARS-CoV-2 targets neurons of 3D human brain organoids

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Abstract

COVID-19 pandemic caused by SARS-CoV-2 infection is a public health emergency. COVID-19 typically exhibits respiratory illness. Unexpectedly, emerging clinical reports indicate that neurological symptoms continue to rise, suggesting detrimental effects of SARS-CoV-2 on the central nervous system (CNS). Here, we show that a Düsseldorf isolate of SARS-CoV-2 enters 3D human brain organoids within 2 days of exposure. We identified that SARS-CoV-2 preferably targets neurons of brain organoids. Imaging neurons of organoids reveal that SARS-CoV-2 exposure is associated with altered distribution of Tau from axons to soma, hyper-phosphorylation, and apparent neuronal death. Our studies, therefore, provide initial insights into the potential neurotoxic effect of SARS-CoV-2 and emphasize that brain organoids could model CNS pathologies of COVID-19.

Keywords brain organoids; cell death; neurons; SARS-CoV-2; Tau pathology
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Introduction

The novel coronavirus disease 2019 (COVID-19) caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) is spreading worldwide, and the outbreak continues to rise, posing a severe emergency (Worl Health Organization, 2020). Understanding the biology of the current COVID-19 pandemic is a high priority for combating it efficiently. Thus, it is essential to gain initial insights into the infection mechanisms of SARS-CoV-2, including its target cell types and tropism, to contain its short- and long-term effects on human health. Furthermore, it is vital to establish an experimental system that could allow designing measures on how to stop viral replication and protect human health rapidly. However, practical problems associated with the isolation and handling of highly infective viral strains and lack of reliable in vitro human model systems that can efficiently model COVID-19 hamper these efforts.

Clinical symptoms of COVID-19 patients include upper respiratory tract infection with fever, dry cough, and dyspnea, indicating that the respiratory tract is the first target (Yang et al, 2020b). However, emerging case reports showed that patients infected with SARS-CoV-2 suffered a sudden and complete loss of the olfactory function, stroke, and other severe neurological symptoms (Chen et al, 2020; Helms et al, 2020; Poyiadji et al, 2020; Sedaghat & Karimi, 2020; Virani et al, 2020). All of these indicate that SARS-CoV-2 could infect the central nervous system (CNS) and is therefore neurotropic (Baig et al, 2020; Conde Cardona et al, 2020; De Felice et al, 2020). Earlier studies show that SARS-CoV target the brains of mice, and since the different coronaviruses share a similar structure, it is likely that SARS-CoV-2 exhibits the same infection mechanism and possibly invades into the brain (McCray et al, 2007). Indeed, a clinical report detected the presence of viral RNA in autopsy of brain samples (Puelles et al, 2020). Furthermore, a postmortem brain MRI analysis has identified the presence of hemorrhagic and encephalopathy syndromes suggesting that
SARS-CoV-2 infection could cause neuronal stress and inflammations (Coenen et al, 2020). Thus, at this point, it is of utmost priority to test whether SARS-CoV-2 directly infects human neurons and productively replicates in the CNS.

To investigate the potential neurotropism of SARS-CoV-2, it is essential to employ a suitable in vitro human model system that recapitulates the physiological effects of SARS-CoV-2 infection. In this regard, the recently emerged human brain organoids that closely parallel the complex neural epithelium exhibiting a wide diversity of cell types could serve as a suitable model system to test the neurotoxic effects of SARS-CoV-2. Induced pluripotent stem cells (iPSCs)-derived human brain organoids have revealed useful insights into human brain development and helped to model a variety of neurological disorders (Lancaster et al, 2013; Gabriel et al, 2016; Birey et al, 2017; Gabriel & Gopalakrishnan, 2017; Xiang et al, 2017; Goranci-Buzhala et al, 2020). Notably, others and our work using brain organoids have revealed unprecedented insights into infection mechanisms, target cell types, and the toxicity effects of the Zika virus (ZIKV) during the recent ZIKV epidemic (Cugola et al, 2016; Qian et al, 2016; Gabriel et al, 2017). These studies validate organoids as a tool for studying not only genetic but also environmental hazards to the human brain.

Here, we report that SARS-CoV-2 readily targets neurons of 3D human brain organoids. Neurons invaded with SARS-CoV-2 at the cortical area display altered distribution of Tau, Tau hyperphosphorylation, and apparent neuronal death. Moreover, we show that although SARS-CoV-2 can readily target brain organoids, SARS-CoV-2 does not appear to efficiently replicate, suggesting that the CNS may not support the active replication of SARS-CoV-2.

Results

Isolation of an infectious SARS-CoV-2 virus

We isolated SARS-CoV-2 (SARS-CoV-2 NRW-42) from a nasopharyngeal and oropharyngeal swab specimen of an infected patient admitted to our university hospital, University of Düsseldorf (see Materials and Methods section for culturing and propagation). To investigate whether SARS-CoV-2 replicates in inoculated African green monkey kidney cells (Vero CCL-81), we performed real-time quantitative polymerase chain reaction (qPCR) analysis with cell culture supernatant. The amount of SARS-CoV-2 RNA drastically increased from 0-dpi until 3-dpi (Appendix Fig S1A). Next, we analyzed the infectivity of generated SARS-CoV-2 particles by propagating virus-containing supernatant to yet uninfected Vero cells. We confirmed the infection of new Vero cells by the emergence of virus-induced cytopathic effects (CPEs) and an increase in SARS-CoV-2 RNA over 4-dpi. The sequence (access number PRJNA627229 at the European Nucleotide Archive and the Sample accession number for NRW-42 which is SRR6522606) showed only eight nucleotide exchanges compared to SARS-CoV-2 Wuhan-Hu-1 isolate.

Isolation and validation of COVID-19 convalescent serum to detect SARS-CoV-2 infection

As of April 1, 2020, we could not procure commercial antibodies that can specifically determine SARS-CoV-2 infection. Therefore, we isolated COVID-19 convalescent serum and tested if they can specifically recognize SARS-CoV-2 infections in our experiments. We obtained blood samples of four independent individuals who recently recovered from COVID-19 (AB1, AB2, AB3, and AB4). Testing them in an enzyme-linked immunosorbent assay (ELISA) that used the SARS-CoV-2 S1 domain of the spike protein as an antigen revealed that, except for AB2, the rest of the convalescent serum contained SARS-CoV-2-specific IgG (Appendix Fig S1B). We then affinity-purified the convalescent serum against the full-length ORF of SARS-CoV-2-N (see Materials and Methods section). In Western blots, which used extracts of brain organoids and Vero cells exposed to SARS-CoV-2, the antibodies affinity-purified from convalescent serum specifically recognized a signal similar to the size of the nucleoprotein of SARS-CoV-2. The recombinant SARS-CoV-2-N serves as a positive control in this experiment (Appendix Fig S1C).

The convalescent serum AB4 also specifically recognized SARS-CoV-2-infected Vero cells. To further validate the specificity of the AB4, we performed co-immunostaining with a mouse monoclonal anti-SARS-CoV-2 S and a polyclonal anti-SARS-CoV-2 NP. As expected, all of these antibodies recognized only the SARS-CoV-2-infected Vero cells (Appendix Fig S2A). Similarly, AB4 could specifically recognize somas of SARS-CoV-2-positive cells in SARS-CoV-2 exposed brain organoids which were further labeled by the monoclonal anti-SARS-CoV-2 S antibody (Appendix Fig S2B). In Western blots that used SARS-CoV-2-exposed organoid extracts, both AB4 and mouse monoclonal antibodies recognized protein bands around 50 and 180 kDs, sizes similar to the nucleoprotein and uncleaved spike proteins. Together, these experiments validate that AB4 detects SARS-CoV-2 infection (Appendix Fig S2C).

SARS-CoV-2 targets neurons of human brain organoids

Before we infected our 3D human brain organoids with the new SARS-CoV-2 NRW-42 isolate, we first tested if our experimental conditions are suitable to infect the well-studied ciliated human respiratory epithelial cells (hRECs), an apparent target for the SARS-CoV-2 (Lamers et al, 2020). We noticed that SARS-CoV-2 readily targets hRECs within 2 days of virus exposure (Fig I A). We then tested if SARS-CoV-2 could infect 3D human brain organoids. To do this, we adapted our previously described protocol and differentiated brain organoids from two different iPSC lines (Donor 1, IMR90 and Donor 2, Crx-iPS; Gabriel et al, 2017). In brief, we started with 10,000 iPSCs and induced differentiation into neural epithelium directly using SB431542 and dorsomorphin, the TGF beta and BMP4 inhibitors, respectively. Our differentiation condition did not also include an exogenous addition of retinoic acid, which could activate retinoic acid receptors (RAR) and induce an aberrant neuronal differentiation (Janesick et al, 2015; Gabriel et al, 2016, 2017; Gabriel & Gopalakrishnan, 2017). As this method skips embryoid bodies formation, it reduces the heterogeneity in organoid sizes simultaneously avoiding the formation of mesoderm and endoderm, which are not required for ectodermal differentiation at early stages of differentiation (Streit et al, 2000). As described before, organoids exhibit their specific neuronal cell types, which are spatially restricted. The ventricular zone (VZ) harbors proliferating neural progenitors cells (NPCs) that display typically elongated nuclei which align to form a lumen, a neural tube-like structure. Cortical neurons are positioned basally to the VZ, forming a cortical plate.
Figure 1.
A positive control experiment. SARS-CoV-2 readily targets ciliated human respiratory epithelial cells (hREC). Acetylated α-tubulin labels cilia. Arrows point SARS-CoV-2-positive cells labeled by AB4 (green). Figures display scale bars. Bar diagram at right quantifies frequencies of SARS-CoV-2-positive cells in hRECs. At least six hREC sections from three (n = 3) independent samples were examined. Data presented as mean ± SEM.

B MOCK organoids of two age groups: Day-15 (i) and -60 (ii) display typical citoarchitecture of brain organoids. L, lumen, VZ, ventricular Zone is containing compact and palisade-like elongated nuclei of neural progenitor cells (NPCs, blue) and CP, a cortical plate containing TuJ-1-positive neurons (magenta). Note a distinct difference TuJ-1 labeling pattern between younger (Day-15) and older (Day-60) brain organoid. Figures display scale bars. Representative images from eight organoids cultured in at least three independent batches (n = 3) derived from donor-1 (MR90) iPS line.

Compared to mock organoids, SARS-CoV-2-exposed Day-15 organoids display SARS-CoV-2-positive cells (AB4, green) in their outer periphery, a region of the cortical plate (i) that is specified by TuJ-1-positive neurons (magenta). L, the lumen of a VZ, the inner area of an organoid where NPCs are located, is free from SARS-CoV-2-positive cells. Magnified region (dotted white box) is given below. At least 10 organoids from five different batches (n = 5) are tested. Figures display scale bars.

Compared to Day 15 organoids and mock (i), Day-60 organoids display an increased number of SARS-CoV-2-positive cells (AB4, green) in their cortical plate that is specified by TuJ-1-positive neurons (magenta) (ii). Magnified region (dotted white box) is given below, showing the perinuclear location of SARS-CoV-2 in cortical neurons. At least 10 organoids from five different batches (n = 5) are tested. Figures display scale bars.

The bar diagram quantifies frequencies of SARS-CoV-2-positive cells in different brain organoid sections derived from two donor iPS lines (MR90 and Cnx-iPS, see Materials and Methods). Please note that each point represents one organoid section. SARS-CoV-2 shows an enhanced tropism for Day-60 organoids. Note, comparative statistics are shown between different age groups and respective days post-infection (dpi) of organoids, and the significance is given as Asterisks in Figure 1A. There is no significant difference in SARS-CoV-2-positive cells between 2- and 4-dpi within each age groups. At least twelve organoids sections from four (n = 4) independent batches, from each donor and day post-infections (dpi), were analyzed. One-way ANOVA, followed by Tukey’s multiple comparisons test,

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Figure 2.
SARS-CoV-2-positive neurons reveal aberrant Tau localization

Next, we identified that the SARS-CoV-2-positive region of the cortical plate is further substantiated by Tau, a microtubule-associated protein that stabilizes neuronal microtubules and promotes axonal growth (Fig 2) (Wang & Mandelkow, 2016). Tau dysfunction is implicated in Alzheimer’s disease (AD) and other Tauopathies. Post-translational modifications in Tau, in particular, phosphorylations, modulate the ability of Tau to bind and assemble microtubules. In Tauopathies, Tau is aberrantly phosphorylated (hyperphosphorylation; Cho & Johnson, 2004; Cohen et al, 2011; Castellani & Perry, 2019). A recent report showed that herpes simplex virus type 1 can induce AD-like effects, including hyperphosphorylation of Tau in 3D human brain-like tissue model (Cairns et al, 2020). This prompted us to investigate if SARS-CoV-2 has a consequence upon its entry into neurons.

Under physiological conditions, Tau is mainly an axonal protein that localizes at the axons of mature neurons (Fig 2A–iv). Applying high-resolution imaging followed by deconvolution, we could visualize Tau’s localization (as probed by a Pan-Tau antibody Tau5A6) exclusively in axons of the cortical neurons (Fig 2Av). The term Tau “missorting” is used when Tau protein is mislocalized into a cell soma and is observed at the early stages of Tau pathology (Zempel & Mandelkow, 2014).

Compared to control organoids where Tau normally localizes in axons, SARS-CoV-2-positive neurons exhibited an altered Tau localization pattern, although it was challenging to visualize mislocalization of Tau in 3D tissues. Nevertheless, using selected confocal sections, we could image an altered Tau localization in SARS-CoV-2-positive neurons. In particular, we identified an enhanced level of Tau into the somas of the SARS-CoV-2-positive neurons. Importantly, we could visualize fractions of these neurons still contained Tau and TUU-1 in their axons, indicating that these neurons are still viable (Fig 2B and C and Appendix Fig S6A).

During the pathogenesis of AD and other Tauopathies, Tau also gets hyperphosphorylated at multiple sites. Sequential phosphorylation at different sites ultimately leads to hyperphosphorylation of Tau (Castellani & Perry, 2019). Phosphorylation of Threonine 231 (T231) is one of the first events in the cascade of phosphorylation, and it regulates the microtubule binding. Still, it is also implicated in disease progression such as detachment of Tau from axonal microtubules (Sengupta et al, 1998; Augustinack et al, 2002a,b; Luna-Munoz et al, 2007; Alonso et al, 2010; Frost et al, 2015). More precisely, we found that compared to control organoids, early Tau phosphorylation marker AT180 recognizes pT231Tau at the soma of the SARS-CoV-2-positive neurons (Fig 2D–F). Imaging the neurons for additional phosphorylated Tau using AT8 antibodies (specific for S202 and T205 of Tau) and p396 (specific for S396 of Tau) revealed that unlike pT231Tau, these phospho-species were restricted to the axons and did not mislocalize to the soma of SARS-CoV-2-positive neurons (Appendix Fig S6B–E). In summary, these results demonstrate the aberrant localization of Tau pT231Tau in SARS-CoV-2-positive neurons suggesting the potential neuronal stress reactions upon virus entry.

SARS-CoV-2 induces neuronal cell death

Phosphorylation of Tau at T231 allows for isomerization into distinct cis- and trans-conformations by the propyl-isomerase PIN1 (Lu et al, 1999). Cis-pT231Tau is acutely produced by neurons after traumatic brain injury, leading to disruption of the axonal microtubule network and apoptosis (Nakamura et al, 2012; Kondo et al, 2015). Analyzing the nuclei of SARS-CoV-2-positive cells (Fig 3A), we realized that they are highly condensed or fragmented exhibiting a strong reaction to 4’,6-diamidino-2-phenylindole (DAPI) that labels nuclei, a feature quite frequently observed in dead cells. To test neuronal cell death as a consequence of SARS-CoV-2 infection, we stained the SARS-CoV-2-exposed samples with terminal deoxynucleotidyl transferase dUTP nick end labeling (TUNEL) that detects fragmented DNA in dead cells (Darzynkiewicz et al, 2008). Compared to un-exposed control organoids, we identified an overall increase in TUNEL-positive cells in SARS-CoV-2-exposed organoids suggesting that virus exposure has caused cell death within 2-dpi (Fig 3B). Staining for SARS-CoV-2-positive cells revealed that most of the virus-positive cells were TUNEL-positive. Besides, we also noticed that some SARS-CoV-2-positive cells were also positive for caspase-3, a protease that specifies programmed cell death (Fig 3C). Interestingly, a fraction of...
Figure 3.
caspase-positive cells displayed pT231Tau localization at the cell soma. Furthermore, TUNEL-positive cells in un-exposed control organoids (which could be after programmed cell death) did not contain pT231Tau suggesting that this different Tau phosphorylation pattern is associated with SARS-CoV-2 entry (Fig 3iii). Thus, it appears that Tau is aberrantly phosphorylated in response to the viral-induced stress, which may elicit further cell death programs that remains to be elucidated.

Discussion

So far, the possible direct effect of SARS-CoV-2 on the CNS has been debated but not experimentally demonstrated (Baig et al., 2020; Conde Cardona et al., 2020:718; Coolen et al., 2020; Helms et al., 2020; Poyiadji et al., 2020). Thus, it was essential to examine whether SARS-CoV-2 can directly target human neurons and whether this leads to productive infection. In contrast to vascular, kidney, and intestinal organoids (Lamers et al., 2020; Monteil et al., 2020; Zhou et al., 2020), brain organoids do not appear to strongly support the active replication of SARS-CoV-2 at least until 6-dpi. There are several reasons for this. Firstly, the developmental stages of brain organoids used in this work may not contain the full complement of SARS-CoV-2’s host cell replication factors. As an example, efficient replication of SARS-CoV requires ACE-2 (Li et al., 2003) whose expression appears to be relatively low in brain organoids (Appendix Fig S4B). Next, in brain organoids, post-mitotic neurons seem to be susceptible for SARS-CoV-2 that may not be permissive (Fig 1, Appendix Figs S3 and S5). Finally, brain organoids are simplified reductionist models and lack an additional cell type that can influence viral replication such as blood–brain barrier, vasculature, and mature glial cells, including microglia. To our surprise, we did notice the appearance of fewer Iba-1 and S100β-positive cells in our organoids pointing toward the need of further engineering of our differentiation conditions which could lead to the differentiation of mature microglia and astrocytes (Appendix Fig S3C and D). Thus, future experiments using aged organoids and bioengineered organoids with SARS-CoV-2 replication factors are required to conclude if brain organoids can support productive infection of SARS-CoV-2.

ACE-2 is an entry receptor for SARS-CoV and efficient replication of SARS-CoV (SARS outbreak in the year 2003) and also depends on the expression level of ACE-2 (Li et al., 2003; Hoffmann et al., 2020). Curiously, SARS-CoV could only infect the brain of transgenic mice expressing an elevated level of human ACE-2 but not non-transgenic mice. This key finding suggests that the neurotropism of SARS-CoV, to some extent, depends on the expression level of human ACE-2 in the brain (McCray et al., 2007). Using our 3D human brain organoid system, we unexpectedly find that although these organoids express low level of ACE-2, the human neurons are indeed a target for SARS-CoV-2. This finding offers a couple of possibilities. First, even a basal level of ACE2 expression is sufficient for viral entry into the neurons. Second, the presence of yet unknown neuron-specific viral entry factors has to be elucidated. It is indeed intriguing that even a low level of ACE-2 is sufficient for the viral entry, and this could explain why SARS-CoV-2 has a broad spectrum of target organs and cell types (Puelles et al., 2020).

Detection of Tau phosphorylation at T231 in SARS-CoV-2-positive neurons is remarkable as it could trigger a cascade of downstream effects that finally could initiate neuronal stress and toxicity. Intriguingly, there is growing evidence that viral infections, particularly herpes simplex virus type 1 (HSV-1), is a potential causative agent leading to Alzheimer’s disease (AD). Indeed a recent work demonstrated infection of 3D human brain-like tissue model with HSV-1 and showed that the HSV-1 infection is sufficient to elicit AD-like effects, including hyperphosphorylation of Tau. Early Tau phosphorylation at T231 could be reversible (Castellani & Perry, 2019). However, phosphorylation events observed in conjunction with apparent neuronal cell death suggest that SARS-CoV-2 has potential detrimental effects on neurons at least in our organoid test system (Fig 3). Future biochemical experiments dissecting the ratio of soluble and sarkosyl-stable Tau extracted from SARS-CoV-2-positive neurons are required to obtain insights into the cause and effect of potential Tau pathology and neuronal death. Although we observe Tau abnormalities in SARS-CoV-2-positive neurons, we could not conclude whether the observed effect is directly caused by the virus or an effect due to neuronal stress, which warrants future investigations.

In conclusion, COVID-19 research has taken center stage in biomedical research. It is noteworthy that three coronavirus epidemics have occurred within the last two decades, and thus, the future zoonotic coronavirus outbreak is not unexpected. With the advent of emerging human organoid research, which did not exist 20 years ago, we should be able to model the current SARS-CoV-2
infections and sufficiently prepare us for the future. Recent works utilizing kidney, gut, and liver organoids have already revealed insights into the infection mechanisms (Lamers et al., 2020; Monteil et al., 2020; Yang et al., 2020a; Zhou et al., 2020). Adding to them is the current work that establishes brain organoids as a test system for SARS-CoV-2 infection and provides indications for potential neurotoxic effects of SARS-CoV-2. Since organoids are an experimentally tractable human in vitro system and convenient to culture as well as to infect, organoid systems may serve well as a test-bed to screen for anti-SARS-CoV-2 agents. The presented work only provides initial insights into primitive brain-like tissues and requires further experiments to dissect viral replication mechanisms and whether there are ACE2 independent pathways for viral entry. It is important to note that although the virus seems to preferably target neurons, future experiments are required to test if the virus can have extended access across the entire organoids. Advanced experiments utilizing a mature state of brain organoids, bioengineered organoids, and orthogonal experiments with complementary in vivo experimental models are assured to dissect the neuropathology of SARS-CoV-2.

Materials and Methods

Clinical specimens

For the isolation of infectious SARS-CoV-2 particles, nasopharyngeal and oropharyngeal swab specimens from one individual with positive qRT–PCR results for SARS-CoV-2 infection were used. The swab specimen was transported in a viral cultivation medium and stored at 4°C overnight. Freezing at −20°C was found to interfere with the infectivity of viral particles. Before the inoculation of susceptible cells, 500 μl maintenance medium (Dulbecco’s Modified Eagle Medium (Thermo Fisher), 2% fetal calf serum (PAN Biotech), 100 U/ml penicillin, and 100 μg/ml streptomycin (Gibco) were added to the swab specimen. To get rid of major impurities, samples were briefly centrifuged (3,000 × g; 60 s) and the supernatant was transferred to new vials.

Human respiratory epithelial cells and culturing

To obtain respiratory epithelia, a MedScand Cytobrush Plus GT (Cooper Surgical, Trumbull, USA) with a gentle-touch tip was rinsed with isotonic saline before use. Afterward, the brush was inserted into the inferior nasal meatus followed by rotatory and linear motions against the medial and superior side. Isolated cells were transferred into a 15-ml centrifuge tube (Corning Incorporated, New York, USA) and 2% Antibiotic-Antimycotic 100× and 2% Antibiotic-Antimycotic 1000x and seeded on T-25 or T-75 rat-tail collagen-coated tissue flasks (Greiner Bio-One, Kremsmünster, Austria), according to the pellet size, respectively, and incubated at 37°C, 5% CO₂. To reduce the risk of contamination, the medium was replaced after 24 h, and the flasks were then integrated into the regular feeding procedure (exchange of medium every 48–72 h). After 1 week, the concentration of Antibiotic-Antimycotic was reduced to 1%. Reaching confluency of 90%, the collagen layer was digested by incubating with 200 U/ml collagenase type IV (Worthington Biochemical Company, New Jersey, USA) for 30–60 min, followed by several washing steps with DMEM/F-12 supplemented with 1% Antibiotic-Antimycotic. To reduce the number of fibroblasts, the pellet was re-suspended in 7 ml DMEM/F12 supplemented with 2% UltroserTM G, seeded on tissue culture treated T-25 flasks (Corning Incorporated, New York, USA) and incubated for 1 h at 37°C, 5% CO₂. The cells were then separated by incubating with Trypsin-EDTA 0.05% for 5 min before the reaction was stopped with FBS followed by centrifugation at 900 rpm for 5 min at room temperature.

After re-suspending in PneumaCultTM-Ex Medium (STEMCELL Technologies, Vancouver, Canada), 4 × 10⁵ cells/ml were seeded on collagen-coated 6.5 mm Transwell®, 0.4 μm pore Polyester membrane inserts (Corning Incorporated, New York, USA) with 250 μl medium on the apical side and 500 μl on the basolateral side, respectively. Before airlift, after 3–5 days, depending on cell confluency, PneumaCultTM-Ex Medium was replaced every day at the apical and basolateral side. To perform airlift, the medium on the apical side was carefully removed, whereas the basolateral medium was exchanged with PneumaCultTM-ALI Medium. The airlifted inserts were then integrated into the regular feeding procedure and incubated at 37°C, 5% CO₂. A fully differentiated pseud stratified epithelium is expected 15–30 days after airlift and resembles human airway epithelium (in vivo) with respect to function and morphology.

Inoculation of Vero cells

In compliance with the German committee’s decision on biological agents (ABAS) of the Federal Institute for Occupational Safety and Health, all experimental studies involving infectious SARS-CoV-2 were performed within the biosafety level 3 (P3) facility at the University Hospital Düsseldorf. To isolate SARS-CoV-2 from a clinical specimen, 2.5 × 10⁵ Vero cells (ATCC-CCL-81, obtained from LGC Standards) were seeded into T-25 cell culture flasks in maintenance medium and cultured at 37°C in a humidified cell culture incubator. The following day, SARS-CoV-2 inoculum was prepared by diluting 200 μl of a clinical specimen with 800 μl maintenance medium. The medium was removed from Vero cells, and 1 ml inoculum (1 ml of maintenance medium for control Vero cells) was added onto the Vero cell monolayer. Vero cells were incubated for 1 h on a laboratory shaker at 37°C in a humidified incubator. Afterward, 4 ml of maintenance medium was added. To monitor viral replication, 100 μl of supernatant was directly harvested as the first sample (0 h post-inoculation) and every 24 h for 4 days post-inoculation. Additionally, cells were imaged by light microscopy.

Real-time qPCR analysis for quantification of SARS-CoV-2 RNA copies per ml

For extraction, 100 μl cell culture supernatant was incubated with 400 μl AVL buffer (viral lysis buffer used for purifying viral nucleic
acids; cat No. 19073, Qiagen, Hilden Germany) for 10 min at RT and mixed with 400 µl 100% ethanol. RNA extraction was performed with 200 µl cell culture mix using the EZ1 Virus Mini Kit v2. (cat. no. 955134, Qiagen, Hilden, Germany) following the manufacturer’s instructions. A total of 60 µl were eluted from the 200 µl starting material. 5 µl of the eluate was tested in qRT–PCR using the real-time TaqMan®-technique. A 113 base pair amplicon in the E-gene of SARS-CoV-2 was amplified and detected, as described by Corman et al (2020) with minor modifications. The thermal protocol described has been shortened to 40 cycles of 95 °C. We used the LightMix® Modular SARS and Wuhan CoV E-gene (Cat.-No. 53-0776-96) and the LightMix® Modular EAV RNA Extraction Control. We used the AgPath-ID® One-Step RT–PCR Kit (Applied Biosystems, Cat. No. 4387391). RT–PCR was performed with an ABI 7500 FAST sequence detector system (PE Applied Biosystems, Weiterstadt, Germany). As a DNA-standard, a plasmid (pEX-A128-nCoV2019-E-gene) that encompasses the amplified region was created and serially diluted after purification. The software constructed a standard graph of the CT values obtained from serial dilutions of the standard. The CT values of the unknown samples are plotted on the standard curves, and the number of SARS-CoV-2 RNA copies was calculated.

For gene expression analysis of ACE2, quantitative RT–PCR analysis was performed by using qPCR MasterMix (PrimerDesign Ltd) and fluorescence emission was monitored by LightCycler 1.5 (Roche). For normalization, primers #5163 (5’CCA CTG CTC CAC CTT TGA 3’) and #5164 (5’ACC CTG TTG TAG CCA 3’) were used monitoring cellular GAPDH expression. Expression was then calculated as 2⁻ΔΔCT.

**Propagation of infectious SARS-CoV-2 particles**

For propagation of infectious SARS-CoV-2 particles from Vero cell culture supernatant, 2.5 × 10⁵ Vero cells were seeded into T-25 cell culture flasks in maintenance medium and incubated at 37°C in a humidified cell culture incubator. The next day, the supernatant of inoculated Vero cells at day four post-inoculation (see above) was diluted with maintenance medium (1:2, 1:10, 1:100, 1:1,000) in a total volume of 5 ml and added to the cells, which were incubated for 4 days at 37°C.

**Determining SARS-CoV-2 viral titer by TCID50 assay in 96-well plates with Vero cells**

For determination of viral titer in TCID50/ml, 5 × 10⁵ Vero cells were seeded in the first 10 columns of 96-well plate in 100 µl maintenance medium and incubated at 37°C in a humidified cell culture incubator for 24 h. In a new 96-well plate, 180 µl maintenance medium was added to all wells of the first 10 columns. For serial dilutions of the virus stock, 20 µl of the stock solution was added to the wells of the first column. Then, 20 µl of the first dilution was transferred to the wells of the next column to obtain 10-fold serial dilutions up to 10⁻⁹. The tenth column of the 96-well plate serves as a control. After exchanging the medium of the previously prepared Vero cell plate with 100 µl fresh maintenance medium, 100 µl of each virus dilution was transferred to the Vero cell plate. After incubation at 37°C for 4 days, microscopic inspection of the plate was used to monitor cytopathic effects (CPEs) in the form of detached cells. TCID50/ml was determined as:

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\text{TCID}_{50}/\text{ml} = \frac{D_{s} \times 1,000}{D_{s} \times V}
\]

\(D_{s}\) = dilution factor of consecutive dilutions (10); \(N\) = total number of wells showing CPE; \(R\) = replicates per dilution (8); \(D_{s}\) = dilution factor of the first dilution (10); \(V\) = volume per well in µl (200 µl).

To estimate MOI, we first calculated the viral titer as TCID50/ml of our generated SARS-CoV-2 by an end-point dilution assay as previously described (Flint et al., 2015). In brief, based on induced cytopathic effects, we calculated the TCID50/ml using the above formula based on the Spearman-Karber method (Ramakrishnan, 2016). To further confirm this calculation with respect to the novelty of this formula, we also applied the commonly used Reed and Muench method (Lei et al., 2020). Both of these methods resulted in a TCID50/ml of 5,000 that we then used to calculate the PFU/ml. Applying poisson distribution, we estimated that the amount of infectious viral particles per ml (PFU/ml) in our stock is 3,500 PFU/ml.

In the context of our infection experiments, we provided 5 µl virus stock per organoid. According to our calculation, the 5 µl volume of SARS-CoV-2 stock contains approximately 17.5 PFUs. Having then estimated the number of viable cells after disintegrating organoids (an average of 100,000 for Day 15 and 200,000 for day 60), we could determine the multiplicity of infection (MOI). Considering 17.5 PFUs, our estimated MOI is 1.8 × 10⁻⁴ and 8.8 × 10⁻⁵ for day 15 and day 60, respectively. Importantly, we found that such a low viral load is sufficient for our studies.

**SARS-CoV-2 infection**

All experiments including SARS-CoV-2 infections were performed in a P3 safety laboratory (see above). Neurons and brain organoids were tested and found free from mycoplasma contamination using the mycoplasma kit (Minevera, Cat. No. 11-1050). For viral exposure, 15- and 60-day-old organoids were transferred from spinner flasks into low-adherent 12 well plates. Each well contained one organoid in 2 ml differentiation medium and added with SARS-CoV-2 and was incubated as stationary suspension culture. To exclude that the observed effects were not induced by SARS-CoV-2, the control organoids (control, uninfected) were treated with supernatants of non-infected Vero cells.

**Generation of convalescent serum, ELISA validation, and affinity purification of SARS-CoV-2-N specific antibodies**

AB1 and AB2 were obtained 23 and 16 days after the diagnosis of SARS-CoV-2 infection. AB3 and AB4 were obtained 27 and 28 days after the diagnosis of SARS-CoV-2 infection (by PCR). Blood samples were drawn directly into serum collection tubes and spun for 15 min at 1,450 g. After centrifugation, the clear supernatant was aliquoted and stored at −80°C. ELISA was performed using semi-quantitative SARS-CoV-2-1gA and SARS-CoV-2-1gC ELISAs that detect binding against the recombinant S1 domain of the SARS-CoV-2 spike protein (Euroimmun, Lübeck, Germany).

The full length ORF of SARS-CoV-2-N was amplified from the vector pUC57-2019-nCoV-N (GeneScript) with the primers:
We differentiated iPSCs into NPCs using STEMdiff Neural Induction Medium (Stem cell technologies, USA). Five days later, the formed neurospheres were collected and cultured on poly-l-ornithine (PLO)-/laminin coated dishes. Seven days later, using a neural rosette selection medium (Stem cell technologies, USA), we re-cultured neural rosettes to generate NPCs. NPCs were differentiated into cortical neurons as described previously (37). Briefly, NPCs were seeded on poly-l-ornithine (PLO)-/laminin coated coverslips. Forty-eight hours later, NPCs were switched to cortical neuronal differentiation medium consisting of BrainPhys basal medium (38) supplemented with 1× B27 (without vitamin A, Thermo Scientific, USA), 1× N2 (Thermo Scientific, USA), 20 ng/ml BDNF (PeproTech, USA), 20 ng/ml GDNF (PeproTech, USA), 20 ng/ml NT3, 1 μM cAMP (Sigma, USA), and 0.2 μM ascorbic acid (Sigma, USA). Fresh medium was added every 2–3 days.

**Generation of iPSCs-derived brain organoids and outgrowths**

Organoids were generated from two different iPS cell lines, namely IM980 (Donor 1, Miltenyi, 130-096-726) and Crx-iPS (Donor 2) as described previously (Gagliardi et al., 2018). We adapted previously described protocol to differentiate iPSCs into brain organoids described earlier (Lancaster et al., 2013; Gabriel et al., 2016). Five-day-old neurospheres were harvested and embedded in matrigel (Corning, USA) drops. Differentiation medium mixture of DMEM/F12 and Neural Basal Medium (in 1:1 ratio), supplemented with 1:200 N2, 1:100 l-glutamine, 1:100 B27 w/o vitamin A, 100 U/ml penicillin, 100 μg/ml streptomycin, 23 μM insulin (Sigma-Aldrich), 0.05 mM MEM non-essential amino acids (NAA), and 0.05 mM β-mercaptoethanol (Life Technologies) was used to differentiate the matrigel embedded droplets in suspension culture. After 4 days of culturing, embedded neurospheres were transferred to spinner flasks (IBS, Integra biosciences) containing the same differentiation medium supplemented with 0.5 μmol dorsomorphin (Sigma-Aldrich, USA).

Organoids with outgrowing neurons were generated as described previously using 60-day-old organoids (Gabriel et al., 2016). Organoids were sliced and plated onto coverslips previously coated with poly-l-ornithine and laminin. The slices were grown in organoid medium for 14 days until the extended neural structures were observed under a stereomicroscope.

**Western blot**

The gel electrophoretic separation of proteins was performed under denaturing conditions in the presence of SDS in a non-continuous gel system, which consisted of a 5% stacking gel and 10% resolving gel, which was then transferred to nitrocellulose membranes. Once the transfer was finished, the membrane was soaked into 5% milk in TRIS–HCl-based buffer (TBST) for a minimum of 30 min at RT. After incubating with primary antibodies overnight at 4°C, the blots were treated with secondary antibodies at 1 h. Super Signal West Pico or Femto Chemiluminescent substrates (Pierce) were used for detection. Antibody dilutions for Western blots: human convalescent IgG antibodies conjugated to HRP (1:5,000, 31430, Thermo Fisher Scientific), goat anti-rabbit IgG (1:5,000, 31466, Invitrogen), anti-human secondary antibodies conjugated to HRP (1:5,000, Thermo Scientific).
Immunofluorescence and confocal microscopy

For light microscopy analysis, monolayer cells (Vero and aspc-derived neurons) were fixed for 10 min. Brain organoids were fixed for 30 min. We used 4% paraformaldehyde/PBS as a fixative (Gabriel et al., 2016). Organoids were incubated in 30% sucrose overnight at 4°C, embedded in Tissue-Tek O.C.T. compound (Sakura, Netherlands). Organoids were cryofrozen at −80°C before sectioning into 10–15 μm thin slices using Cryostat Leica CM3050 S. Thin sections and cells were permeabilized with a buffer containing 0.5% Triton X100 for 10 min. Specimens were blocked with 0.5% fish gelatin/PBS for 1 hr, both at room temperature. For SOX2 staining, antigen retrieval was required. For this, sections were treated with repeated heating (microwave) in owium citrate buffer (10 mM Sodium citrate, 0.05% Tween 20, pH 6.0) and were applied before permeabilization and blocking.

We used different antibodies as follows: human convalescent serum (AB1 to AB5, 1:40), AB4, 1:40 or 1:50 or 1:100, rabbit anti-TUJ-1 (1:400, Sigma-Aldrich, T2200), monoclonal mouse anti-phospho-Tau AT180 (1:100, Thermo Fisher, MN1040), polyclonal rabbit anti-phospho-Tau (S396), (1:100, Thermo Fisher), monoclonal mouse anti-SARS-CoV-2 (1:200, Genetex, GTX635679), rabbit anti-SARS-CoV-2 NP (1:200, Biozol, GTX-GTX632604), rabbit anti-S-100 beta (1:100, Abcam, ab52642), rabbit anti-iba-1 (1:100, Abcam, ab178846), rabbit anti-MAP2 (1:100, Proteintech, Cat# 17490-1-AP), mouse anti-Pan-Tau (1:100, DSHB, 5A6). Specimens with primary antibodies were incubated overnight at 4°C. For secondary antibodies, donkey anti-rabbit IgG (H+L) secondary antibody, Alexa Fluor 594 (Thermo Scientific, Cat# A21207), donkey anti-rabbit IgG (H+L) secondary antibody, Alexa Fluor 647 (Thermo Scientific, Cat# A31573), donkey anti-rabbit IgG (H+L) secondary antibody, HRP (Thermo Scientific, Cat# A16023), goat anti-mouse IgG (H+L) secondary antibody, Alexa Fluor 488 (Thermo Scientific, Cat# A28175), Goat anti-mouse IgG (H+L) secondary antibody, Alexa Fluor 594 (Thermo Scientific, Cat# A-11032), Goat anti-mouse IgG (H+L) secondary antibody, Alexa Fluor 647 (Thermo Scientific, Cat# A-21236), Alexa Fluor Dyes conjugated either with goat/donkey anti-mouse, anti-human, or goat-anti-rabbit (1:500 or 1:100, molecular probes, Invitrogen) was used. For DNA staining, DAPI at a concentration of 1 μg/ml (Thermo Scientific, Cat# 32670) was used, and the coverslips were mounted using Mowiol (Carl Roth, Germany). The raw images were collected using a Leica SP8 confocal system (Leica microsystems, Germany) and processed with the help of Adobe Photoshop (Adobe Systems, USA). For deconvolution, the captured image files were processed using ZEN software (2.3, SP1, black, 64 bit, release version 14.0.0.0; ZEISS, Oberkochen, Germany) for 3D reconstruction and deconvolution. After deconvolution, files were imported into Fiji and further processed using Image J, Adobe Photoshop CC 2018, and Adobe Illustrator CC 2018. For 3D surface and volume rendering, raw image files were processed using Imaris (64× version 7.7.1).

TUNEL assay

Apoptotic cells were detected by using DeadEndFluorometric TUNEL System (Promega, G3250, USA) according to the manufacturer’s protocol.

Ethical approval and patient samples

Serum samples AB1 and AB2 were obtained under a protocol approved by the ethical committee, medical faculty, University Hospital Düsseldorf, Heinrich-Heine-University (study number 5350). Serum samples AB3 and AB4 were obtained under a protocol approved by the Institutional Review Board of the University of Cologne (protocol 16-054). Human respiratory epithelial cells (hREC) were obtained by nasal brush biopsy from healthy control individuals. The study was endorsed by the local ethical committee at the University of Münster, and each patient gave written informed consent (Study number, 2015-104-5-S, Flimmerepithel) and 2020-274-5-S (COVID-19). Trained physicians from the Department of General Pediatrics, University Hospital of Münster, performed biopsies.

Statistical analysis

The statistical analyses were performed using GraphPad Prism version 8. All experiments were performed at least in triplicates, and the statistical significance of each dataset was analyzed using Student’s t-test followed by Welch’s correction and non-parametric one-way ANOVA followed by Tukey’s post hoc test. For immunofluorescence-based experiments, we randomized the samples to avoid any bias. The values are expressed as mean ± SD or SEM. Independent experiments have been represented by “n”.

Data availability

No amenable data sets.

Expanded View for this article is available online.

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Author contributions

AR and JG conceptualize the project. AR designed, coordinated and conducted experiments. LM and PNO performed virology works. HS supervised virology. EG, PA-I, AM-S and AM provided technical support. AM-S and CK purified antibodies. OG provided iPS cells. HG, DW and FK provided convalescent serum. KW and HO provided airway cells. AW, MA, SH, TH, AD, OA and JT involved in sequencing and dent experiments have been represented by “n”.

Conflict of interest

The authors declare that they have no conflict of interest.

References


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