Virologic Features of Severe Acute Respiratory Syndrome Coronavirus 2 Infection in Children

Lael M. Yonker,1,2,3,4 Julie Boucau,4,5 James Regan,5 Manish C. Choudhary,3 Madeleine D. Burns,1 Nicola Young,1 Eva J. Farkas,1 Jameson P. Davis,1 Peter P. Moschovis,2,3 T. Bernard Kinane,5,6 Alessio Fasano,1,2,3 Anne M. Neilan,2,3,6 Jonathan Z. Li,3,5 and Amy K. Barczak1,2,3,6

1Massachusetts General Hospital, Mucosal Immunology and Biology Research Center, Boston, Massachusetts, USA, 2Massachusetts General Hospital, Department of Pediatrics, Boston, Massachusetts, USA, 3Harvard Medical School, Boston, Massachusetts, USA, 4Ragon Institute of Massachusetts General Hospital, Massachusetts Institute of Technology and Harvard, Cambridge, Massachusetts, USA, 5Brigham and Women’s Hospital, Department of Medicine, Boston, Massachusetts, USA, and 6Massachusetts General Hospital, Department of Medicine, Boston, Massachusetts, USA


Methods. Respiratory swabs were collected from children with COVID-19. Viral load was quantified by reverse-transcription polymerase chain reaction (RT-PCR); viral culture was assessed by direct observation of cytopathic effects and semiquantitative viral titers. Correlations with age, symptom duration, and disease severity were analyzed. SARS-CoV-2 whole genome sequences were compared with contemporaneous sequences.

Results. One hundred ten children with COVID-19 (median age, 10 years [range, 2 weeks–21 years]) were included in this study. Age did not impact SARS-CoV-2 viral load. Children were most infectious within the first 5 days of illness, and severe disease did not correlate with increased viral loads. Pediatric SARS-CoV-2 sequences were representative of those in the community and novel variants were identified.

Conclusions. Symptomatic and asymptomatic children can carry high quantities of live, replicating SARS-CoV-2, creating a potential reservoir for transmission and evolution of genetic variants. As guidance around social distancing and masking evolves following vaccine uptake in older populations, a clear understanding of SARS-CoV-2 infection dynamics in children is critical for rational development of public health policies and vaccination strategies to mitigate the impact of COVID-19.

Keywords. SARS-CoV-2; pediatric COVID-19; viral dynamics.
adults and adolescents in many places in the world and our understanding of transmission dynamics have evolved, masking and distancing policies are being relaxed [12]. Policy changes have necessarily been made despite the paucity of data providing insight into the role that pediatric disease might play in ongoing transmission. As viral variants that enhance the potential for transmission and/or reduce vaccine efficacy emerge [13–15], the importance of identifying potential reservoirs of viral replication and transmission has been brought into the spotlight. Defining the virologic features of SARS-CoV-2 infection in children and the potential for children to transmit virus will facilitate rational public health decision-making for pediatric populations.

In this work, we sought to define fundamental virologic features of SARS-CoV-2 in a pediatric population across a range of disease severity. We analyzed respiratory swabs from children presenting to urgent care clinics or the hospital with symptomatic and asymptomatic COVID-19 infection. Clinical factors, such as age, COVID-19 risk factors, and disease severity were compared with viral features including SARS-CoV-2 viral load, isolation of replication-competent virus, and whole viral sequencing. Our data indicate that age, from infancy through adulthood, is not a predictor of viral infection dynamics, and that children of all ages can have high SARS-CoV-2 viral loads of replication-competent virus, including variants, displaying comparable dynamics to those seen in adults.

MATERIALS AND METHODS

Sample Collection

Infants, children, and adolescents <21 years of age presenting to Massachusetts General Hospital (MGH) urgent care clinics or the hospital with either symptoms concerning for or known exposure to COVID-19 (April 2020–April 2021) were prospectively enrolled in the Institutional Review Board (IRB)–approved MGH Pediatric COVID-19 Biorepository (IRB number 2020P000955) [16]. After informed consent, and assent when appropriate, a research-designated swab of the nasopharynx, oropharynx, and/or anterior nares was obtained and placed in phosphate-buffered saline. Samples were aliquoted and stored at –80°C. Samples from patients who tested positive for COVID-19 on clinical SARS-CoV-2 reverse-transcription polymerase chain reaction (RT-PCR) testing were analyzed. Nasal samples from adults hospitalized with acute COVID-19 [10] (April 2020–August 2020; enrolled in Institutional Review Board–approved MGH COVID-19 Biorepository, IRB number 2020P000804) with duration of symptoms equal to the hospitalized pediatric cohort were selected for comparative studies.

Clinical Data Collection

Demographic and clinical factors were recorded through a combination of manual chart reviews and data extraction from electronic health records, then collected in a REDCap database [17] through the Partners Electronic Health Record Registry of Pediatric COVID-19 Disease (IRB number 2020P003588). Trained reviewers collected demographics, SARS-CoV-2 risk factors, comorbid conditions, medications, COVID-19–related symptoms, and laboratory tests. Outcome of initial presentation to care, admission status, and complications of COVID-19 disease were also extracted by manual review.

SARS-CoV-2 Viral Load Quantification

Virions were pelleted from anterior nasal, oropharyngeal, and nasopharyngeal swab fluids by centrifugation at approximately 21 000g for 2 hours at 4°C. RNA was extracted using Trizol-LS (Thermo Fisher) according to the manufacturer’s instructions. RNA was then concentrated by isopropanol precipitation, and SARS-CoV-2 RNA was quantified using the CDC N1 primers and probe (https://www.cdc.gov/coronavirus/2019-ncov/lab/rt-pcr-panel-primer-probes.html) as previously described [10]. Standards for the quantitative assay were created by transcribing SARS-CoV-2 RNA containing the nucleocapsid gene and making serial dilutions from 10^9 copies to 3 copies. Standards were run with all samples and the resulting standard curve was used to quantify the amount of SARS-CoV-2 RNA in each sample well. Each sample was tested in triplicate. As there was no significant difference in viral load from respiratory secretions obtained from the anterior nares, nasopharynx or oropharynx of participants (Supplementary Figure 1), samples were analyzed together, regardless of collection site.

Viral Culture

Vero-E6 cells (ATCC) were maintained in D10+ media (Dulbecco’s modified Eagle’s medium [DMEM, Corning] supplemented with HEPES [Corning], 1X penicillin/streptomycin [Corning], 1X glutamine [Glutamax, ThermoFisher Scientific], and 10% fetal bovine serum [FBS, Sigma]) in a humidified incubator at 37°C in 5% carbon dioxide (CO₂). Vero-E6 cells were passaged every 3–4 days, detached using Trypsin–EDTA solution (Fisher Scientific), and seeded at 150 000 cells per well in 24-well plates for culture experiments and 20 000 cells per well in 96-well plates the day before inoculation for median tissue culture infectious dose (TCID₅₀) experiments.

After thawing, each specimen was filtered through a Spin-X 0.45µm filter (Corning) at 10 000g for 5 minutes. Fifty microliters of the supernatant was then diluted in 450 µL of D₇ media (DMEM supplemented with HEPES, 1X penicillin/streptomycin, and 1X glutamine). The viral culture experiments were performed as previously reported [18] with the following modifications: 100 µL of the solution was used to inoculate wells in a 24-well plate and 1 mL of D₇ media (D + media with 2% FBS) was added to each well after 1 hour of incubation. The plates were then placed in a 5% CO₂ incubator at 37°C. For TCID₅₀ measurements conducted in parallel, 25 µL of the Spin-X
flow-through was used to inoculate Vero-E6 cells in a 96-well plate in the presence of 5 µg/mL of polybrene (Santa Cruz Biotechnology) using 5-fold dilutions (5:1 to 5:6) and 4 repeats for each sample. The plates were centrifuged for 1 hour at 2000g at 37°C before being placed in a 5% CO₂ incubator at 37°C. The SARS-CoV-2 isolate USA-WA1/2020 strain (BEI Resources) was used as a positive control for cytopathic effect (CPE) in both culture and TCID₅₀ experiments.

Viral culture and TCID₅₀ plates were observed at 3 and 6 days postinfection with a light microscope and wells showing CPE were counted. The TCID₅₀ titers were calculated using the Spearman–Karber method. For the culture plates, the supernatant of the wells displaying CPE was harvested 10–14 days postinfection and RNA was isolated using a QIAamp Viral RNA Mini kit (Qiagen) for confirmation of the viral sequence.

SARS-CoV-2 Sequencing
cDNA synthesis was performed using Superscript IV reverse transcriptase (Invitrogen). Whole viral amplification was performed with the Artic protocol using multiplexed primer pools designed with Primal Scheme generating 400-bp tiling amplicons [19, 20]. PCR products were pooled and Illumina library construction was performed using the Nextera XT Library Prep Kit (Illumina). The comparison dataset included 183 representative contemporaneous SARS-CoV-2 genomes from Massachusetts present in GISAID to assess for local clustering. Nucleotide sequence alignment was performed with MAFFT (multiple alignment using fast Fourier transform) [21]. Best-fit nucleotide substitution GTR + G + I was used for the datasets using model selection in IQ-Tree followed by maximum likelihood phylogenetic tree construction using IQ-Treeweb server with 1000-bootstrap replicates [22].

Analysis
All statistical analyses were performed using parametric comparisons in GraphPad Prism (version 9.1.1), including Pearson correlation, analysis of variance (ANOVA) with multiple comparisons, and unpaired t test.

RESULTS
Clinical Cohort
One hundred ten children diagnosed with COVID-19 with a mean age of 10 years (range, 0–21 years) were included in the study (Table 1). There were slightly more boys (56%) than girls (44%) with SARS-CoV-2 infection included in our analyses. One-third of the participants were white (33%), 10% black, and 4% Asian; one-third (38%) reported their ethnicity as Hispanic. Past medical history in children is reported in Supplementary Table 1. Thirty children were asymptomatic but were identified as having COVID-19: 26 children (27%) presented to urgent care/COVID-19 testing sites because of a COVID-19 exposure, while 4 children (4%) were identified on routine screening during hospital admission. Eight (7%) presented with COVID-19 symptoms but had no known COVID-19 contact. The majority of participants with COVID-19 did not require hospitalization (75 children [68%]). Thirty-six children (33%) were hospitalized with COVID-19, although only 18 children (16%) required supplemental oxygen and/or invasive or non-invasive respiratory support (referred to as moderate/severe COVID-19).

Age Did Not Impact SARS-CoV-2 Viral Load or Recovery of Replication-Competent Virus
Age is a well-established risk factor for developing severe COVID-19. Accordingly, asymptomatic patients were significantly younger than patients with mild disease, and pediatric patients who were hospitalized with hypoxemia were significantly older than asymptomatic children or children with mild disease (ANOVA, P = .0002) (Figure 1A). However, viral load was not increased in more severe disease: asymptomatic children and children with mild disease displayed significantly higher viral loads than adults hospitalized with COVID-19 with comparable duration of symptoms (≤10 days) (ANOVA, P < .0001) (Figure 1B). However, there were no differences in viral load between pediatric patients hospitalized with moderate/severe disease and hospitalized adults of similar duration.
of illness (ANOVA, pediatric hospitalized vs adult hospitalized, \( P = \) not significant [ns]) (Figure 1B) (Adult demographics are detailed in Supplementary Table 2).

We then sought to assess whether COVID-19 severity impacted the relationship between viral load and age in pediatric cohorts of varying severity: asymptomatic, mildly symptomatic, and moderate/severe pediatric COVID-19 patients. There was no significant correlation of age with viral load in any of the COVID-19 severity groups (Pearson correlation: \( r = -0.1, 0.01, \) and \(-0.2, \) respectively; \( P = \) ns) (Figure 1C). Thus, a child’s age did not appear to impact viral load and viral load did not drive disease severity: All children, from 2 weeks through 21 years of age, regardless of disease severity, were equally capable of carrying a high viral load.

Furthermore, there were no differences in viral clearance over the duration of illness, not only when comparing mild pediatric COVID-19 with moderate/severe COVID-19, but also when comparing pediatric COVID-19 with adults hospitalized with COVID-19 (comparison of simple linear regression, \( P = \) ns) (Figure 1D). Duration of symptoms did not affect the finding that viral load does not correlate with age of the infected child (Supplementary Figure 2).

As SARS-CoV-2 RNA detection by RT-PCR does not specify whether replication-competent virus is being shed, we next sought to ascertain risk factors for shedding live virus by performing viral culture assays for recoverable SARS-CoV-2 in parallel with viral load testing. From the 110 participants, we collected 126 samples; live virus was cultured from 33 samples coming from 32 participants. Of note, 8 of these children with culturable SARS-CoV-2 were asymptomatic. Higher viral load was significantly predictive of shedding of live virus (t test, \( P < .0001 \)) (Figure 2A). Consistent with the results for

![Figure 1. Coronavirus disease 2019 (COVID-19) disease severity and severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) viral load across age groups. A, Age of pediatric patients with SARS-CoV-2 infection, stratified by disease severity: asymptomatic (n = 27), mild disease, outpatient (n = 48), or moderate/severe COVID-19, hospitalized (n = 18). Analyzed by ordinary 1-way analysis of variance (ANOVA). B, SARS-CoV-2 viral load was quantified across a range of disease severities. Patients presenting with <10 days of symptoms were compared, including asymptomatic pediatric outpatients (n = 27), mildly symptomatic pediatric outpatients (n = 44), moderate/severe pediatric hospitalized patients with oxygen requirement (n = 18), and moderate/severe adult hospitalized patients (n = 29). Analyzed by ordinary 1-way ANOVA. C, Correlation of viral load and age, stratified by asymptomatic (n = 30), mild outpatient (n = 48), and moderate/severe hospitalized (n = 18) cohorts. Analyzed by Pearson correlation. D, Viral load of hospitalized adults (n = 29), hospitalized pediatric participants requiring respiratory support (n = 18), and pediatric outpatients with mild disease (n = 48), plotted against duration of symptoms. Dotted lines depict limit of detection. Analyzed by simple linear regression, comparison of slopes. ∗\( P < .05, \) ∗∗\( P < .01; \) ∗∗∗\( P < .001, \) ∗∗∗∗\( P < .0001; \) ns = not significant.](https://academic.oup.com/jid/article/224/11/1821/6396772)
viral load, age was not associated with viral culture results; live virus was recovered from participants aged 1 month through 21 years \((t\) test, \(P = ns\)) (Figure 2B). Semi-quantitative assessment of the amount of virus shed by an individual participant was assessed by TCID\(_{50}\). The TCID\(_{50}\) for culture-positive specimens correlated strongly with viral load (Pearson correlation: \(r = 0.7, P < .0001\)) (Figure 2C) but did not correlate with age across all pediatric participants (Pearson correlation: \(r = -0.1, P = ns\)) (Figure 2D).

**Children With COVID-19 Were Most Infectious Within First 5 Days of Illness**

To define the likely period of infectiousness in our pediatric population, we analyzed viral load, culturability, and TCID\(_{50}\) in comparison with duration of symptoms. Of note, duration of symptoms does not necessarily indicate duration of infection, as time infection was acquired cannot be confirmed. Consistent with prior reports in adults [9], viral loads in children were the highest earliest in the course of illness and declined over time after symptom onset (Pearson correlation, \(r = -0.4, P < .001\)) (Figure 3A). Viral load was highest in the first 2 days of symptoms, with significant decrease after 5 days of symptoms and further decline after 10 days of symptoms \((P < .0001\)) (Figure 3B). Analysis of pediatric viral culture results demonstrated that children tested early after symptom onset were more likely to shed replication-competent virus \((P = .004\)) (Figure 3C). Correspondingly, semi-quantitative assessment of infectious viral shedding in children showed that the TCID\(_{50}\) was higher early after symptom onset and decreased over time. When grouped by days of symptoms, children in days 0–2 of their symptoms had the highest infectivity, while children with >6 days of illness shed less virus \((P = .004\)) (Figure 3D).

**Pediatric SARS-CoV-2 Sequences Were Representative of Those Found in the Community**

We successfully performed whole viral sequencing of 57 respiratory samples from 54 children. Phylogenetic analysis of these pediatric sequences with contemporaneous Massachusetts sequences from GISAID showed that they were representative of the spectrum of sequences found in the community (Figure 4). Notable variants identified in the pediatric samples included 4 Alpha (B.1.1.7) and 3 Iota (B.1.526.2) variants. To validate our culture results on a subset of culture-positive samples, we sequenced virus isolated from the supernatant from 8 positive samples. Sequences from the supernatant and respiratory specimens were identical in 7 cases and demonstrated only 1 nucleotide change in the last case.

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**Figure 2.** Severe acute respiratory syndrome coronavirus 2 culture results across age groups and viral load. A and B, Samples with observable cytopathic effect (CPE) (culture positive [+], \(n = 31\)) or without observable CPE (culture negative [−], \(n = 95\)) plotted against viral load (A) and participant age (B) and compared using \(t\) test. C and D, Semiquantitative viral titer expressed as median tissue culture infectious dose (TCID\(_{50}\))/mL for culture-positive samples plotted against corresponding viral load (C) or participant age (D). Analyzed using Pearson correlation. Dotted line depicts limit of detection. ****\(P < .0001\).
DISCUSSION

As the global COVID-19 pandemic took hold, infected older adults suffered high rates of hospitalization and death while infected children typically experienced paucisymptomatic or asymptomatic infection. While it is now clear that children can become infected with and transmit SARS-CoV-2, viral dynamics in children have been understudied, and a full understanding of the dynamics of infection in children is needed to inform public health policies specific to the pediatric population. Here, we show that pediatric patients of all ages, from infancy to young adulthood, can carry a high SARS-CoV-2 viral load in their upper airways, particularly early in the course of infection, and an elevated viral load corresponds with high levels of viable, replicating virus. Pediatric sequences were largely reflective of those found in the general community and the presence of novel variants was identified.

While several other studies have reported viral load, here we characterize additional features of virus culturability, which corresponds with infectivity by live virus, and genomic sequencing, which corresponds with the capacity of viral SARS-CoV-2 to generate variants. These features add important insight into SARS-CoV-2 infection in children. Similar to prior studies, we show that children can carry large quantities of SARS-CoV-2 [2], although reports on which clinical factors correspond with the greatest viral load vary [2, 23, 24]. These differences can be driven by time from exposure to time of testing, and behavioral or environmental factors of the cohort tested. Importantly, consistent with prior reports, we found that asymptomatic children can make up a significant proportion of an infected pediatric cohort [23]. We found that asymptomatic children can carry high viral loads and shed culturable virus, emphasizing the importance of considering this subgroup when characterizing pediatric COVID-19.

Our findings have significant implications for both public health policy and the potential role of universal vaccination of pediatric populations in fully curbing the COVID-19 pandemic. As vaccination has rolled out in adult populations, public health policies are being adjusted to account for changes in risk that result from vaccination. Our results emphasize the importance of considering and clarifying how these policy implications differ between pediatric and adult populations.
changes relate to children. As adult populations have been vaccinated, pediatric cases have represented a growing proportion of infections, currently accounting for up to 25% of all COVID-19 cases across different regions of the United States [3]. Our results suggest that the low rates of transmission in settings such as schools and daycares cannot be attributed to low viral loads, low rates of viral shedding, or rapid clearance of virus in younger patient populations. As changes in masking and distancing policies are implemented for vaccinated adults, consideration of how and whether policy changes will be applied for children will be critical for ongoing reduction of new COVID-19 cases.

Our results additionally suggest that pediatric populations have the potential to serve as a community reservoir of actively replicating virus, with implications for both new waves of infection and the evolution of viral variants. The durations of natural and vaccine-induced immunity for each vaccine in clinical use are not yet known. If a community reservoir of actively replicating virus is maintained and transmitted within unvaccinated pediatric populations, that population could then serve as a source of new infections as vaccine-induced immunity wanes in vaccinated adult populations. In addition, viral genomic variants were readily identified in the pediatric samples and these variants have the potential to impact viral transmission [25–27], disease severity [28, 29], and vaccine efficacy [30]. Ongoing viral replication within pediatric populations has the potential to serve as a source of existing and new viral variants that interfere with eradication efforts.

Figure 4. Phylogenetic analysis of pediatric and community severe acute respiratory syndrome coronavirus 2 sequences. Maximum likelihood tree generated from pediatric sequences (red) and 183 contemporaneous Massachusetts sequences from GISAID.

2.0E-4
Our study has several limitations. First, the data collected here represent a single medical center and affiliated pediatric urgent care/COVID-19 screening clinics. However, these were among the few pediatric testing centers encompassing a large catchment area during the duration of this study, and patients enrolled spanned a wide range of symptoms. There may have been an enrollment bias that was not assessed with this study. Additionally, many of these samples were collected early in the pandemic and SARS-CoV-2 variants of interest have shifted over time. Ongoing studies analyzing shifts in virologic features of SARS-CoV-2 infection in children alongside studies of infection in adults are needed to better understand the full reach of the COVID-19 pandemic.

Ultimately, our data suggest that although age is generally protective against severe disease, children, especially early in the infection course, carry high viral loads of SARS-CoV-2, which can include viral variants. Our results underline the importance of defining public health policy with viral dynamics in children in mind and of including pediatric populations in vaccine efforts aimed at eradication.

Supplementary Data
Supplementary materials are available at The Journal of Infectious Diseases online. Supplementary materials consist of data provided by the author that are published to benefit the reader. The posted materials are not copyedited. The contents of all supplementary data are the sole responsibility of the authors. Questions or messages regarding errors should be addressed to the author.

Notes

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