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Defining biological and biophysical properties of SARS-CoV-2 genetic material in wastewater

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HIGHLIGHTS
• Raw wastewater contains intact, enveloped SARS-CoV-2 particles.
• Pasteurization and freeze-thaw of samples can greatly decrease signal.
• Raw wastewater does not contain infectious SARS-CoV-2

GRAPHICAL ABSTRACT

ABSTRACT

SARS-CoV-2 genetic material has been detected in raw wastewater around the world throughout the COVID-19 pandemic and has served as a useful tool for monitoring community levels of SARS-CoV-2 infections. SARS-CoV-2 genetic material is highly detectable in a patient’s feces and the household wastewater for several days before and after a positive COVID-19 qPCR test from throat or sputum samples. Here, we characterize genetic material collected from raw wastewater samples and determine recovery efficiency during a concentration process. We find that pasteurization of raw wastewater samples did not reduce SARS-CoV-2 signal if RNA is extracted immediately after pasteurization. On the contrary, we find that signal decreased by approximately half when RNA was extracted 24–36 h post-pasteurization and ~90% when freeze-thawed prior to concentration. As a matrix control, we use an engineered enveloped RNA virus. Surprisingly, after concentration, the recovery of SARS-CoV-2 signal is consistently higher than the recovery of the control virus leading us to question the nature of the SARS-CoV-2 genetic material detected in wastewater. We see no significant difference in signal after different 24-hour temperature changes; however, treatment with detergent decreases signal ~100-fold. Furthermore, the density of the samples is comparable to enveloped retrovirus particles, yet, interestingly, when raw wastewater samples were used to inoculate cells, no cytopathic effects were seen indicating that wastewater samples do not contain infectious SARS-CoV-2. Together, this suggests that wastewater contains fully intact enveloped particles.

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1. Introduction

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), the causative agent of coronavirus disease 2019 (COVID-19), was first identified in Wuhan, China in December 2019 and was declared a global pandemic by the World Health Organization (WHO) in March 2020. To date, SARS-CoV-2 has produced >201 million cases and >4.2 million COVID-19 related deaths worldwide (WHO, August 5th 2021). SARS-CoV-2 has been shown to be spread primarily by respiratory droplets and occasionally by aerosols (Liu et al., 2020; Port et al., 2021; Crawford et al., 2021; Wang and Du, 2020).

SARS-CoV-2 has 75–80% nucleotide similarity to severe acute respiratory syndrome coronavirus (SARS-CoV) that was responsible for outbreaks of severe acute respiratory syndrome in 2002 and 2003 in Guangdong Province, China (Zhu et al., 2020; Zhong et al., 2003; Ksiazek et al., 2003; Drosten et al., 2003). Both SARS-CoV and SARS-CoV-2 use the cellular receptor Angiotensin-converting enzyme 2 (ACE2) which is highly expressed in the lung and oral mucosa and expressed at lower levels in the digestive tract (Yan et al., 2020; Hammig et al., 2004; Xu et al., 2020). Due partially to population sizes, material shortages, inaccessibility to laboratory equipment, a vast array of disease severity, and healthcare coverage concerns, it has not been possible to test every individual regularly for a SARS-CoV-2 infection. Even as testing becomes more and more available, asymptomatic individuals may not get tested and can unknowingly continue spread. These limits cause difficulty in monitoring community spread.

It has been reported that SARS-CoV, SARS-CoV-2, and other coronavirus RNA is detectable in feces of infected patients up to 6 to 10 days before symptom onset (Corman et al., 2016; Leung et al., 2003; Gu et al., 2020; Holshue et al., 2020; Song et al., 2020; Chen et al., 2020a). Additionally, screening of sewage, both community and hospital, for the detection and prevalence of viruses including SARS-CoV-2, Poliovirus, noroviruses, adenoviruses, rotaviruses, polyomaviruses, Hepatitis A virus, and gastroenteritis viruses have been documented and can correlate closely with the occurrence of cases in the community (Katayama et al., 2008; Fumian et al., 2010a; Wang et al., 2005a; Kroiss et al., 2018; Pecia et al., 2020; Asghar et al., 2014; Bofill-Mas et al., 2000; Fumian et al., 2010b; Pintó et al., 2007; Rodríguez-Díaz et al., 2009; Victoria et al., 2010; Villena et al., 2003). This supports wastewater surveillance as a useful method for monitoring community levels of SARS-CoV-2 infections (Pecia et al., 2020; Baldovin et al., 2021; Agrawal et al., 2021; Cao and Francis, 2021; Bivins et al., 2020; Venugopal et al., 2021).

Unlike from sputum, samples collected from feces of infected patients do not generally appear to contain infectious viral particles despite high levels of detectable viral RNA (Wolfel et al., 2020a; Chen et al., 2020b; Ling et al., 2020; Wolfel et al., 2020b); however, infectious particles cultured from feces has been reported before (Xiao et al., 2020). To date and to the best of our knowledge, there have been no confirmed cases of COVID-19 linked directly to wastewater treatment plants. Several groups have examined the survival rate of various other coronaviruses in raw, unpasteurized wastewater, and generally conclude that after a maximum of 3 days there is a 99.9% decrease in infectivity (Gundy et al., 2009; Wang et al., 2005b; Ye et al., 2016; La Rosa et al., 2020; Carducci et al., 2020).

During periods of lower community infection rates, it was necessary to concentrate raw wastewater samples to reliably detect SARS-CoV-2 genetic material by quantitative reverse transcription polymerase chain reaction PCR (qPCR). Viral concentration from wastewater has not been possible to test every individual regularly for a SARS-CoV-2 infection. Even as testing becomes more and more available, asymptomatic individuals may not get tested and can unknowingly continue spread. These limits cause difficulty in monitoring community spread.

To create the control sequence for qPCR standard curve, the plasmid described above was digested with EcoRI. The primer pair (COVID19-N 5p: 5′ ATCTCGTATAATGGACCCCAATATCCGGC 3; COVID19-N 3p: 5′ TATAGGCTTGAAGTTAGTGACTCACGTG 3′) was used to amplify the N ORF fragment from IDT’s 2019-nCoV_N_Positive Control plasmid and cloned into pSC-A of StrataClone PCR cloning kit of Agilent for sequence check. The primer pair vectors linking the unique puromycin resistance gene sequence not found in nature. Interestingly, we noticed a disparity in recovery rates between our control virus and SARS-CoV-2 signal, leading us to question the difference in make-up of the genetic material detectable in wastewater in comparison to our control virus. In this innovative manuscript, we examine recovery, temperature resistance, density, detergent resistance, and infectivity and conclude, for the first time, that the genomic material detected in wastewater is enveloped and non-infectious.

2. Methods

2.1. Plasmids and Puro Virus production

The NL4-3 derived HIV containing the CMV driven Puromycin resistance gene and lacking the accessory genes Vif, Vpr, Nef, and Env was engineered using InFusion Cloning (TaKaRa). To make this construct, we used a previously described NL4-3 derived HIV-CMV-GFP provided by Vineet Kewal Rammani (National Cancer Institute (NCI) – Frederick) (Zufferey et al., 1997; Lucas et al., 2010). This proviral vector lacks the accessory genes vif, vpr, nef, and env and contains a CMV promoter driven GFP in the place of nef. The NL4-3 derived HIV-CMV-GFP was digested using Stu1 and Xma1 (New England Biolabs (NEB)) to remove the GFP gene, and a gBlock fragment (Integrated DNA Technologies (IDT)) of a uniquely codon optimized Puromycin resistance gene put in its place. The unique puromycin resistance gene sequence is as follows:

ATGACAGAGTATAAGACCAAGCTCGGCTCAGACAGACAGATG
TCCCGAGGCGACATGCGGACACCCCGCCGCGCTGGCTTTGGACTAC
CAGAAGAACACACTGGACATCCGACCACAGACAGCCTCAGACTCGT
GAAGGCCGAAAGAAGGTGACGCGAGGCTGGACTGCAAGTTGTA
GAGGCCAATTGCTGGATGCCGCTGCTCGTACGCAGCATGGAAG
GCCGTCTGCACCTACTACAGACAAAAAGGCGAGCGGCTTCGAGCA
GCTGACTACCTGCTGATGGTGAGACGGCAAGAGGGCGGGGCGG
GCCCTTCGAGAACACTCGCACAGCCACAACTGGCCTTCACAGGAG
CTGAGTAGCCAGACCTCAGCTTCGGCGAGACAGGGTCGGAGCCG
CAGCTTGCTGACATGACGAGGAAAGCGGGGGGCGTCTAGA

To create the control sequence for qPCR standard curve, the plasmid described above was digested with EcoRI. The primer pair (COVID19-N 5p: 5′ ATCTCGTATAATGGACCCCAATATCCGGC 3; COVID19-N 3p: 5′ TATAGGCTTGAAGTTAGTGACTCACGTG 3′) was used to amplify the N ORF fragment from IDT’s 2019-nCoV_N_Positive Control plasmid and cloned in a pSC-A of StratataClone PCR cloning kit of Agilent for sequence check. Later, the N ORF fragments were infused using an InFusion kit (TaKaRa) as described above.

The control, Puro Virus, was made via stable cell line. To make the cell line, the above plasmid and Vesicular Stomatitis Virus protein G (VSV-G) expression plasmids were transfected into HEK293FT cells (Invitrogen, Carlsbad, CA, USA). After 48 h, viral supernatant was collected and used to transduce fresh HEK293FT cells which were selected with puromycin 48 h post-transduction. HEK293FT cells were maintained in Dulbecco’s Modified Eagle Medium (DMEM) supplemented with 7.5% fetal bovine serum, 2 mM l-glutamine, 1 mM sodium pyruvate, and 10 mM nonessential amino acids. Virus was quantified using qPCR (described below).

2.2. Quantitative RT-qPCR assay

The TaqMan probe (VIC-5′ CGGTAAGGTTTGACTCCGCGAC 3′-QSY) and the primer pair (puro Forward: 5′ CCCATGCACACATAGACG G3′; puro Reverse: 5′ CCACTTACGAGACTTTTGACG 3′) were designed and used to target the puro RNA sequence described above. Primers and probe containing an artificial gene sequence not found in nature.
specifity was tested by BLAST analysis (NCBI) to prevent known non-
specific binding targets that could be obtained in a human specimen.
The choice of VIC fluorescent dye for the puro TaqMan probe is for the
application in the multiplex reactions with the SARS-CoV-2 N gene
TaqMan probe utilizing the FAM reporter. The TaqMan probe (FAM-5'
ACCCCGCATACATTGGTGAC3′ BHQ1) and the primer pair (2019-
ncov-N1-F: 5′ GACCCCCAATTCACGGAAT 3′; 2019-ncov-N1-R: 5′
TCTGGTACTCAGCAATCGT 3′) for N1 detection, and The TaqMan
probe (FAM 5′ ACAATTTGGCCCGATTCG 3′ BHQ1) and the primer pair
(2019-ncov-N2-F: 5′ TTACAAACATTGGCGCAGAAA 3′; 2019-
ncov-N2-R: 5′ GGCGGACATATTGGAAAGA 3′) for N2 detection were pur-
chased from Integrated DNA Technologies (IDT), based on the CDC
2019-ncov Real-Time RT-PCR Diagnostic Panel (Acceptable Alternative
Primer and Probe Sets) https://www.cdc.gov/coronavirus/2019-ncov/
downloads/List-of-Acceptable-Commercial-Primers-Probes.pdf.
A plasmid (described above) carrying a unique puro resistance gene
fragment along with a N gene fragment was constructed, purified from
Escherichia coli, and used as standards for the RT-qPCR assay to ensure
an equal molar ratio of puro and N gene detection. A standard curve
was constructed at concentrations of 200,000 through 2 gene copies/μL
and utilized to determine the copy number of the target puro gene in the
wastewater samples that had the puro control virus added prior to con-
centration as an internal control of RNA extraction rate.

Final RT-qPCR one step mixtures consisted of 5 μL TaqPath 1-step
RT-qPCR Master Mix (Thermo Fisher cat# A15299), 500 nM each of
puro or SARS-CoV-2 primers, 125 nM of TaqMan probe, 5 μL of waste-
water RNA extract and RNase/DNase-free water to reach a
final concentration of 20 μL. All RT-qPCR assays were performed using a 7500
Fast real-time qPCR machine (Applied Biosystems). The reactions were initiated with
1 cycle of Uracil N-glycosylase (UNG) incubation at 25 °C for 2 min to
eliminate carryover and then 1 cycle of reverse transcription at 50 °C
for 15 min, followed by 1 cycle of activation of DNA polymerase at
95 °C for 2 min and then 45 cycles of 95 °C for 3 s for DNA denaturation
and 55 °C for 30 s for anneal and extension. The data is collected at the
55 °C extension step.

2.3. Sample acquisition

Wastewater treatment plants from around Missouri collected at
least 1 L of 24-hour composite sample of raw sewage from influent
wastewater intake. Data such as influent flow rate, pH, water tempera-
ture, chemical oxygen demand (COD), and total suspended solids (TSS)
were collected from the 1 L composite sample, 3 × 50 mL aliquots of
sample were packed in coolers containing ice packs and delivered to
the University of Missouri within 24 h from collection.

2.4. Concentration and recovery

Nine samples in duplicate containing 50 mL of raw wastewater were
stored at 4 °C (18 total 50 mL samples). Raw samples were spun at
2000 × g for 5 min to remove large particulates, then vacuum filtered
through a 0.22 μm filter (Millipore cat# SCGPO0525), mixed with a
50% (wt/v) Polyethylene glycol (PEG) (Research Products International
RPI) cat# P48080) and 1.2 M NaCl solution for a

50% (wt/v) Polyethylene glycol (PEG) (Research Products International
RPI) cat# P48080) and 1.2 M NaCl solution for a

final concentration of 12% PEG and 0.3 M NaCl. NL4-3 derived HIV containing the CMV
driven Puromycin resistance gene and lacking the accessory genes Vif,
Vpr, Nef, and Env were added to the filtered sample/PEG/NaCl mixture at a concentration of
5.1 × 107 viral particles per sample (50 mL). Samples were mixed thoroughly and kept at 4 °C for 1 h, then spun at
12,000 × g at 4 °C for 2 h. All but 140 μL of supernatant was removed, and the entire volume was extracted for RNA purification. RNA was
extracted from the samples using the Qiagen QIAamp Viral RNA mini kit
(cat# 52906) in a QIAcube Connect (Qiagen cat# 9002864). Addition-
ally, 140 μL of wastewater was collected prior to filtering the sample,
after filtration (before addition of PEG solution); these samples were mul-
tiplied to be comparable to the full 50 mL volume. After concentration of
virus. Viral recovery was determined by qPCR as described above. Impor-
tantly, for samples containing high copy numbers of SARS-CoV-2 genetic
material (>1.5 million copies/L), it was not necessary to concentrate sam-
ple, so for some experiments, unconcentrated wastewater was used to remove variables that may be introduced in concentration.

2.5. Stability assessment

Six samples containing 50 mL of raw wastewater were stored at 4 °C. Samples were mixed gently and split into 3 × 16.7 mL aliquots. Aliquots
were stored at either 4 °C, Room Temperature (RT) (~22 °C), or 37 °C for
24 h. After 24 h, 140 μL of sample was collected, RNA was extracted, and
viral recovery was determined as previously described. As the samples
had a concentration of SARS-CoV-2 signal over 1.5 million copies/L,
the concentration step was not necessary and removed any further var-
iables that would be introduced from this step. Statistics are a paired,
two tailed Student’s t-test run on Microsoft Excel. The function used was =t-test, array1, array2, tails, type where array1 and array2 are
the values of each sample at each condition, respectively, and options
for two tails and paired tests were selected.

2.6. Pasteurization

Six duplicate samples containing 50 mL of raw wastewater were
stored at 4 °C (12 total samples). One 50 mL tube of each duplicate sample
was kept at 4 °C, and the other 50 mL tube of sample was incubated at
60 °C for 2 h then the entire sample was concentrated as described
above. RNA extraction from the entire pellet, RT-qPCR, and statistis
were performed as described above in Sections 2.4, 2.2 and 2.5, respecti-
vely.

For pasteurization effect on signal 24 h later, six duplicate samples
were pasteurized and RNA extraction was done either immediately
after the 2-hour incubation at 60 °C or 24–36 h after pasteurization
(12 total samples). RNA extraction, RT-qPCR, and statistics were performed as described above in Sections 2.4, 2.2 and 2.5, respecti-
vely.

2.7. Freeze-thaw sensitivity

Six duplicate samples each containing 50 mL of raw wastewater col-
lected from a wastewater facility as described above, the same week,
were stored at 4 °C. At the start of the study, one 50 mL tube of each du-
plicate sample was kept at 4 °C, and the other 50 mL tube of sample was
stored at −80 °C for 36 h then thawed and concentrated as described
above. RNA extraction, RT-qPCR, and statistis were performed as de-
scribed above in Sections 2.4, 2.2 and 2.5, respectively.

2.8. Detergent sensitivity

Six duplicate samples containing 50 mL of raw wastewater col-
lected as described above and stored at 4 °C (12 total samples). One of
each duplicate sample was treated with either 1% Triton X 100 or PBS
for 2 h at 37 °C. From unconcentrated, raw wastewater samples, RNA
extraction, RT-qPCR, and statistis were performed as described above in Sections 2.4, 2.2 and 2.5, respectively.

2.9. Density

Three samples containing 50 mL of raw wastewater were stored at
4 °C. The Puro Virus (NL4-3 derived HIV containing CMV driven
Puromycin resistance and lacking the accessory genes Vif, Vpr, Nef,
and Env) was added to raw samples at a concentration of 5.1 × 107
viral particles per sample. Samples were concentrated as described
above. Concentrated samples were then added to a density gradient
ranging from 0% to 28% iodixanol in a 0.23 M sucrose diluant according
to the Optiprep protocol (Sigma cat# 92339-11-2). Gradients were spun
in a Sorvall Discovery 100SE ultracentrifuge at 31,000 RPM for 3 h at
4 °C. After centrifugation, the gradient was fractioned. RNA extraction and RT-qPCR were performed as described above in Sections 2.4 and 2.2, respectively. Density of each fraction was confirmed using a densitometer (Abbe, model: C10).

2.10. Infectivity

Ten raw wastewater samples from the week of June 28th 2021 were collected and brought to University of Missouri as described in Section 2.3. Fresh samples were filtered through a 0.22 mm filter (Millipore cat# SCG00525). For the cell maintenance, the Vero E6 cells (CRL1569™, ATCC) were maintained in Dulbecco’s Modified Eagle Medium (GIBCO, Grand Island, NY) containing 10% fetal bovine serum (FBS) (Atlanta Biologicals, Lawrenceville, GA) at 37 °C with 5% CO2. For the first recovery, 200 ml of each sample was inoculated to Vero E6 cells in 6-well plates at a confluence of ~90%. After 1 h of adsorption, the inoculum was removed, and the cells were washed with PBS and covered with fresh optiMEM (Gibco, Thermo Fisher Scientific) containing 1× Antibiotic-Antimycotic (Gibco, Thermo Fisher Scientific). Three days post inoculation, 1 mL of the supernatant from the last virus recovery was centrifuged and inoculated to fresh Vero E6 cells for the second and third virus recovery and cytopathic effect was observed. A clinical isolate from Missouri, SARS-CoV-2/human/USA/20 × 1003/2020, (Full genome available at GenBank Accession ID: MW521470.1) was used as the positive control at a multiplicity of infection (MOI) of 0.001. This isolate has a D614G mutation.

3. Results

3.1. SARS-CoV-2 signal recovery through concentration

Generally, it has been necessary to concentrate wastewater samples for reliable detection of SARS-CoV-2 by qPCR. Because concentration is sometimes necessary, it is important to know the rate of recovery throughout the process. During a COVID-19 outbreak at a Missouri prison, we were able to reliably detect signal in raw, unconcentrated wastewater as signal was high in this wastewater collection facility at a prison, we were able to reliably detect signal in raw, unconcentrated wastewater. Filtration, and after concentration of virus. Signal from unconcentrated samples was multiplied based on the total volume of sample to be concentrated to allow for equal comparison at each step. Error bars represent standard deviation.

Although wastewater is not thought to contain infectious SARS-CoV-2, raw wastewater contains a variety of other pathogens. Filtration through a 0.22 μm filter should remove many of these pathogens, but some have suggested pasteurizing samples at 60 °C for 1 to 2 h to inactivate potential pathogens (Pecson et al., 2021). To test the effect on signal due to pasteurization, duplicate samples were kept at either 4 °C or 60 °C for 2 h prior to RNA extraction (Fig. 2A). We found no significant difference between pasteurized and non-pasteurized samples (P value = 0.23) when RNA was extracted immediately after pasteurization; however, it is important to note that signal dropped significantly if samples were returned to 4 °C after pasteurization and RNA was collected 24–36 h after pasteurization was completed (Fig. 2B). A 24–36-hour period between pasteurization resulted in a 44% reduction in signal (P value = 2.05 × 10−5). Additionally, in a separate experiment, when samples were frozen at −80 °C for 36 h and thawed prior to concentration, over 90% of the SARS-CoV-2 signal was lost (Fig. 2C).

3.3. SARS-CoV-2 signal is higher than enveloped virus control

As a control throughout wastewater screening, an NL4-3 derived HIV virus containing a CMV driven Puromycin resistance gene and lacking the accessory genes Vif, Vpr, Nef, and Env (Henceforth called ‘Puro Virus’) was added to raw wastewater samples at a concentration of 5.1 × 10^7 viral particles per sample (Fig. 3A). Importantly, the Puro Virus contains a uniquely codon optimized puromycin resistance gene. This vector was chosen because the unique sequence present in the Puro Virus ensures that any signal detected throughout our experiments with this probe is from our internal matrix control and not the environment ensuring that nothing environmental will be amplified in our control. Additionally, Both SARS-CoV-2 and the Puro Virus are positive-sense RNA contained in an envelope at a similar size. Interestingly, upon comparison of signal detected in highly potent samples (>1.5 million copies/L of SARS-CoV-2 signal) before and after concentration, SARS-CoV-2 recovery was consistently higher than the Puro Virus recovery (Fig. 3B). On average, SARS-CoV-2 recovery was 2.7-fold higher than Puro Virus recovery (N = 14). The consistent disparity between the SARS-CoV-2 recovery and the Puro Virus recovery led us to hypothesize that the SARS-CoV-2 signal was coming from a different source than enveloped RNA, such as non-enveloped ribonuclear complexes, as we would expect that most enveloped particles would interact similarly with PEG during concentration.

Using the Puro Virus control, we also examined parameters at time of wastewater collection including Temperature, pH value, Chemical Oxygen Demand (COD) value, and Total Suspended Solids (TSS) value and the effect these parameters had on the control puro virus recovery (Fig. 3C). This data was analyzed two ways; first, R2 values were calculated for each dataset as shown in Fig. 3C-F. In addition, to determine if chemical or physical parameters in combination could affect the viral recovery, multiple linear regression analysis was performed using SAS 9.4 (SAS Institute Inc., Cary, NC, USA). Two separate full-model analyses were performed as not every sample had all four parameters collected. One analysis included TSS, COD, and pH (n = 55) and as independent variables and one which included temperature, COD, and pH as independent variables (n = 29); puro recovery (%) served as the dependent variable in both tests. Neither the model including TSS nor the model including temperature were significant (P = 0.061, r2 = 0.133 and P = 0.458, r2 = 0.097, respectively) (Data not shown). Following the finding that the Puro Virus had a consistently lower signal than SARS-CoV-2 signal and that chemical parameters at collection do not impact recovery, we aimed to characterize the SARS-CoV-2 genetic material detected in wastewater.
3.4. SARS-CoV-2 signal stability

After collection from a wastewater treatment plant as described above, samples are stored at 4 °C until they are processed; however, we wanted to test the stability of samples at a variety of temperatures to determine if temperature control made a large impact on SARS-CoV-2 signal. A portion of each sample was kept at either 4 °C, RT, or 37 °C for 24 h as described in methods. RNA was extracted from each sample, and SARS-CoV-2 signal was quantified using qPCR (Fig. 4). Although samples that were maintained at 4 °C had the least variability, there was no significant difference in signal from samples kept at any temperature (P-values range 0.097 to 0.363). This finding is interesting as an increase in temperature from 4 °C to either RT or 37 °C could impact the activity of various enzymes that could be present in raw wastewater and impact rates of degradation of genetic material.

3.5. Detergent removes SARS-CoV-2 signal

Because SARS-CoV-2 is an enveloped virus, it is very likely to be sensitive to detergent because of a disruption of the lipids composing the envelope. Presence of an envelope may protect genomic material from enzymes that may quickly degrade vulnerable genetic material. We were curious as to whether the SARS-CoV-2 signal detected in wastewater was sensitive to detergent. To answer this question, duplicate samples of raw wastewater were treated with either 1% TritonX-100 or PBS and kept at 37 °C for 2 h (Fig. 5). RNA was extracted from 140 μL of unconcentrated wastewater, and samples were quantified using qPCR. Treatment with TritonX-100 reduced signal about 100-fold in comparison to samples treated with PBS alone. Contrary to our original hypothesis, this indicates that SARS-CoV-2 signal detected in wastewater is likely protected by a lipid bilayer, but this finding could also have other causes.
3.6. Density of SARS-CoV-2

A loss of signal following treatment with detergent could be caused by a variety of things including breaking up of protein complexes. Because of this, we wanted to know if the density of SARS-CoV-2 particles was comparable to densities of other known enveloped viruses. To test density, concentrated wastewater samples also containing the Puro Virus at a concentration of $5.1 \times 10^7$ viral particles per sample were run through a density gradient containing 0% to 28% iodixanol in a 0.25 M sucrose dilutant and RNA was extracted from each fraction (Fig. 6). Each fraction was then probed by qPCR for SARS-CoV-2 signal and the Puro Virus control. The fraction containing the highest signal for both the Puro Virus and SARS-CoV-2 correlates with a density ($\rho$) between 1.16 and 1.18 g·mL$^{-1}$ as calculated according to the Optiprep protocol and confirmed by refractometer. This finding is in agreement with the expected density of retroviruses and further supports that the genetic material in wastewater is similar to an enveloped viral particle (Poiesz et al., 1980).

3.7. No cytopathic effects from wastewater samples

The similarities in density along with detergent sensitivity suggest that the genetic material is enveloped. We were curious to know if wastewater samples contained infectious particles. To examine this question, aliquots of 10 raw wastewater samples with SARS-CoV-2 signals ranging from 169,433 to 3,255,921 Copies/L were collected and used to inoculate Vero E6 cells within a week of collection. Seventy-two hours after inoculation, supernatant was collected and used to inoculate fresh Vero E6 cells. At each passage, cells were examined for cytopathic effects. No cytopathic effects were seen at any point during the experiment (Fig. 7). The absence of cytopathic effects suggests that wastewater samples do not contain infectious SARS-CoV-2 Particles. As a positive control, cells were infected with SARS-CoV-2/human/USA/20 × 1003/2020 (GenBank Accession ID: MW521470.1) at a MOI of 0.001. These cells showed significant cytopathic effects whereas the negative control cells had no cytopathic effects.

4. Discussion

COVID-19 causes a wide variety of symptoms and disease severity. This combined with inaccessibility to testing due partially to supply costs, availability, and varied access to healthcare makes accurately

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**Fig. 4. Temperature Stability.** N = 6. Samples were mixed gently and split into 3 aliquots. Aliquots were stored at either 4 °C, RT, or 37 °C for 24 h. After 24 h, RNA was extracted, and viral recovery was determined by qPCR. Error bars represent standard deviation. P-Values: 4 °C to RT = 0.097, 4 °C to 37 °C = 0.108, RT to 37 °C = 0.363.

**Fig. 5. Detergent Sensitivity.** N = 6. Duplicate samples were treated with either 1% Triton X 100 or PBS for 2 h at 37 °C. RNA was extracted from raw samples, and viral recovery was determined by qPCR. Error bars represent standard deviation. P-Value = 2.10E−20.

**Fig. 6. Density.** N = 3. Puro Virus was added to raw wastewater samples at a concentration of $5.1 \times 10^7$ viral particles per sample. Raw samples were concentrated then added to a density gradient ranging from 0% to 28% iodixanol in a 0.25 M sucrose dilutant and spun in a Sorvall Discovery 100SE ultracentrifuge at 31,000 RPM for 3 h at 4 °C. RNA was extracted from fractions, and viral recovery was determined by qPCR.
tracking cases of COVID-19 difficult. Like several other viruses, SARS-CoV-2 was shown to be present in the COVID-19 patient feces and therefore wastewater despite being primarily transmitted via respiratory droplets (Corman et al., 2016; Leung et al., 2003; Gu et al., 2020; Holshue et al., 2020; Song et al., 2020). To date, there have been no confirmed cases of SARS-CoV-2 infection from wastewater treatment plants, and infectious virus has not been able to be reliably cultured from wastewater (Gundy et al., 2009; Wang et al., 2005b; Ye et al., 2016). In early stages of the pandemic and during periods of lower community spread of SARS-CoV-2, it has been necessary to concentrate wastewater samples to reliably detect genetic material using qPCR. Using samples from wastewater treatment facilities around Missouri from areas with high community levels of COVID-19, we have shown that filtration and concentration of wastewater samples reliably reserves ~60% of signal from raw wastewater, supporting that this method of filtering and concentration is a consistent method for concentrating and comparing community viral loads (Fig. 1). While the method described here includes filtration through a 0.22 μm filter to remove other debris and bacterial pathogens, other groups have suggested pasteurizing wastewater samples to inactivate any pathogens present in wastewater (Wu et al., 2020; Weidhaas et al., 2021). We have shown here that pasteurizing samples for 2 h does not impact signal if RNA is extracted from samples immediately; however, this does not remain true for samples that have been pasteurized and returned to 4 °C for RNA extraction between 24 and 36 h post-pasteurization (Fig. 2). While the decrease in signal is approximately 2-fold, it remains an important note that pasteurization may make a consistent experimental timeline of higher importance for those who are quantifying viral loads after pasteurization. Some groups such as Wu et al. (2020) and Weidhaas et al. (2021) report no significant difference between pasteurized and unpasteurized samples; however, others, like Palmer et al. (2021) report a 50–55% decrease in SARS-CoV-2 following pasteurization (Wu et al., 2020; Weidhaas et al., 2021; Palmer et al., 2021). The studies done by Wu et al. (2020), Weidhaas et al. (2021), or Palmer et al. (2021) do not specifically discuss time after pasteurization, so this may be the cause of contradictory data.

Interestingly, we noticed that recovery of the Puro Virus control was consistently about 3-fold lower than the recovery of SARS-CoV-2 through concentration, and that chemical properties of wastewater at time of collection did not impact Puro Virus Recovery (Fig. 3). Initially, this led us to believe that the SARS-CoV-2 signal present in wastewater was coming from a different source than an enveloped particle, but our data suggests the contrary and led us to investigate the nature of the genomic material producing signal.

From the time it is deposited into a sewer system to its arrival at a wastewater treatment plant, a fecal sample may go through a variety of temperature changes. Additionally, as temperatures get closer to 37 °C, enzymes that degrade genetic material may become more active. We show that in a 24-hour period, temperature changes are tolerated as there is no significant difference in signals from samples kept at 4 °C, room temperature, or 37 °C (Fig. 4). This finding is interesting as it suggests some sort of protection of genetic material from degradation and suggests that outdoor temperatures may not impact reliability of signal detection when levels of SARS-CoV-2 genetic material are monitored over time.

Because of genomic material found in feces, it was a high concern early in the pandemic that feces and wastewater may be a source infectious virus; however, efforts to culture infectious virus from fecal or wastewater samples have failed (Chen et al., 2020a; Wolfel et al., 2020a; Ling et al., 2020; Wolfel et al., 2020b). Here we demonstrate that genomic RNA collected from wastewater samples is sensitive to detergent (Fig. 5). This finding suggests that the genomic material is protected by a lipid bilayer as the inactivation of viral particles by detergents has been well documented (Welch et al., 2020; Hellstern and Solheim, 2011; Horowitz et al., 1998; Prince et al., 1986; Roberts, 2008). Additionally, the material concentrated from wastewater samples has a very similar density to non-infectious retroviral particles again supporting that the material concentrated from wastewater shares similarities to an enveloped viral particle (Fig. 6). In agreement with prior findings discussed above, wastewater samples did not contain any infectious SARS-CoV-2 particles (Fig. 7). It is feasible that enzymes present in the digestive tract, such as Trypsin, cleave much of the Spike glycoprotein from virus present in the digestive tract resulting in viral particles that are enveloped yet not infectious. Further studies need to be done to determine whether the genomic material is full length SARS-CoV-2 genomic material and to investigate why the genomic material found in wastewater may be non-infectious while also remaining to be protected by a lipid bilayer.

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draft. Hsin-Yeh Hsieh: Methodology, Investigation. Shu-Yu Hsu: Investigation. Yang Wang: Investigation. Braxton T. Saledo: Investigation. Writing – review & editing. Anthony Belenchia: Conceptualization, Data curation, Funding acquisition. Jessica Klutts: Conceptualization, Investigation. Sally Zemmer: Conceptualization, Investigation. Melissa Reynolds: Conceptualization, Data curation, Funding acquisition. Writing – review & editing. Elizabeth Semikw: Conceptualization, Data curation. Trevor Foley: Conceptualization, Investigation. Xiufeng Wan: Conceptualization, Funding acquisition, Writing – review & editing. Chris G. Wieber: Conceptualization, Project administration, Funding acquisition, Writing – review & editing. Jeff Wenzel: Conceptualization, Project administration, Funding acquisition, Writing – review & editing. Chung-Ho Lin: Conceptualization, Funding acquisition. Marc C. Johnson: Conceptualization, Project administration, Writing – review & editing.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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