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Human viruses in the environment activated by excessive use of COVID-19 prevention supplies

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Abstract

Due to extensive COVID-19 prevention measures, millions of tons of chemicals penetrated into natural environments. Alterations of human viruses in the environment, the neglected perceiver of environmental fluctuations, remain obscure. To decipher the interaction between human viruses and COVID-19 related chemicals, environmental samples were collected on March 2020 from surroundings of designated hospitals and receivers of wastewater treatment plant effluent in Wuhan. The virus community and chemical concentration were respectively unveiled in virome and high-performance liquid chromatography-tandem mass spectrometry. The complex relationship between virus and chemical was further elaborated by random forest model. As an indicator, environmental viruses were corroborated to sensitively reflect the ecological disturbance originated from pandemic prevention supplies. Chemicals especially trihalomethanes restrained the virus community diversity. Confronting this adverse scenario, Human gammaherpesvirus 4 and Orf virus with resistance to trihalomethanes flourished while replication potential of Macacine alphaherpesvirus 1 and SARS-CoV-2 RNA were detected near designated hospitals, suggesting environment as a missing link in the transmission route. This research innovatively underlined the human health risk of pandemic prevention supplies from the virus - environment interaction, appealing for monitoring of environmental viruses in long term.

1. Introduction

Human viruses in the environment pose considerable effects to public health and microbial ecology. Environment matrices were gradually recognized as huge reservoirs of human viruses \cite{Labadie2019} with the assistance of rapidly developed non-culture methods such as metagenomics and virome \cite{Paez-Espino2016,Schulz2020}. In aquatic environments, gastrointestinal-related Adenovirus, Astrovirus and Rotaviruses were detected in rivers, reservoirs \cite{Wang2020}, and even drinking water \cite{Mehle2018}. In terrestrial ecosystems, the prevalence of Influenza A virus was over 80% in soil matrix associated with poultry \cite{Lau2019}. Apart from causing disease, Influenza A virus threatened public health for indirectly initiating the expression of antibiotic resistance in human microbes \cite{Zhang2020}. With the raging of COVID-19 pandemic, the footprint of waterborne SARS-CoV-2 was tracked in waste water treatment plant (WWTP) systems all over the world \cite{Carducci2020}. Besides, the ubiquitous human viruses in the environment affect the microbial ecology through antagonistic or mutualistic interaction. As a consequence of antagonistic relationship (Paul et al., 2021), bacteria that grazed on Echovirus via proteolytic enzymes could outcompete their counterparts (Olive et al., 2020). Norovirus in the environment attached to pathogenic bacteria via histo-blood group antigens (Amarasiri and Sano, 2019) and heightened the bacterial adherence as well as their co-
infection to eukaryotic cells (Neu and Mainou, 2020). In addition, human viruses, as the proxy of human-originated contaminants (Farkas et al., 2020), would be sensitive perceivers of environmental turbulence. During COVID-19 when anthropogenic activities to prevent the spread of pandemic tremendously altered environment, human viruses lying in the intersection of “One Health” perspective (Shaheen et al., 2022) should be elaborated to suppress the potential side effect.

Human viruses in the environment were regulated by various parameters such as temperature, relative humidity and pH (Carratalá et al., 2020). Nevertheless, the interaction between human viruses and chemicals originated from pandemic prevention supplies that were excessively used during COVID-19 remain misty. For the sake of disinfection, chlorine and quaternary ammonium surfactant containing biocidal agents were exorbitantly used in hospitals and wastewater treatment plants (WWTPs) (Elsaid et al., 2021). In Wuhan, the average usage of disinfectants in 26 WWTPs was roughly 40 t day$^{-1}$ normally (Text S1) but it ascended to 84.64 t day$^{-1}$ during pandemic (Wuhan Water Affairs Bureau, 2020). As a recommended adjuvant therapy for treating COVID-19 patients (Sterne et al., 2020), glucocorticoids were clinically used with a proportion of 44.5% (Gu et al., 2020). The consumption rate of glucocorticoids in Wuhan was reckoned at 44,168 mg day$^{-1}$ on March 2020 (Text S2). Environment, the reservoir of human viruses, was inevitably confronted with the lash from residues and by-products derived in disinfectants and glucocorticoids. To what extent chemicals related with COVID-19 prevention i.e., chlorine, trihalomethanes, quaternary ammonium surfactants, and glucocorticoids shaped human viruses in the environment was obscure. To delve into the interplay between chemicals from COVID-19 prevention supplies and human virus community in the environment, soil and water samples were collected from designated hospital surroundings and downstream of WWTPs in Wuhan on March 2020. Human virus community in this special period was manifested by means of metagenomics and virome. Random forest models on various taxonomic levels were executed to investigate the environment - virus interaction with total chlorine, trihalomethanes, quaternary ammonium surfactants, and glucocorticoids taken into consideration. Furthermore, mechanism underlying the bilateral relationship was explored from the dimension of virus viability represented by replication, recombination and repair genes.

2. Materials and methods

This study aimed to unravel the effect of excessively used pandemic prevention supplies on human viruses in the environment and the response of human viruses to the lash from pandemic prevention supplies. To this end, environmental samples were collected from surroundings of hotspots in Wuhan city i.e., designated hospitals and WWTPs on March 2020. Consumption of glucocorticoids and disinfectants were calculated to measure the environmental burden from overused pandemic prevention supplies. Virome was conducted to illustrate the virus community; and UPLC-MS/MS was adopted to detect chemicals generated by pandemic prevention supplies. Random forest model was constructed to identify the interaction between human viruses and chemicals. Genes related with viability were picked out to clarify the mechanism of human viruses - pandemic prevention supplies interplay.

2.1. Sample collection

As one of the first cities that took measures to suppress COVID-19 (Zhou et al., 2020), Wuhan was typical in evaluating the ecological impact of pandemic prevention supplies. Designated hospitals for novel coronavirus pneumonia, the biggest consumer of drugs, were considered as source of glucocorticoids in the environment during COVID-19. WWTPs, the major consumer of disinfectants, linked chemicals and environment with effluent (Larsen and Wigginton, 2020). Medical wastewater from JinYinTan Hospital, HuoShenShan Hospital, and LeiShenShan Hospital was ultimately discharged to FuHe River, Hanjiang River, and XunSi River after disposal of SanJinTan WWTP, CaiDian WWTP, and HuangJiaHu WWTP (Fig. 1). Hence, water and soil samples were collected at 11 sites from surroundings of designated hospitals and receivers of WWTPs (Table S1). To meet the demand of following virome analysis, 500 g of topsoil (0–10 cm) and 1.0 L surface water (~0.5 m) were simultaneously gathered (Liu et al., 2013; Hu et al., 2021). The concentration of Fe$^{3+}$ was adjusted to 1 mg L$^{-1}$ through adding FeCl$_3$ solution for flocculating virus particles (John et al., 2011). Virus particles were then enriched by ultracentrifugation and 0.22 μm filtration. Soil samples and 0.22 μm membranes were stored at −80 °C for subsequent treatment.

2.2. Virome sequencing and processing

DNA and RNA were respectively extracted by PowerSoil DNA Isolation Kit (Qiagen, Germany) and RNeasy PowerSoil Total RNA Kit (Qiagen, Germany) (Wang et al., 2019). Reverse transcription reaction was conducted with PrimeScript RT Master Mix (Takara, Japan). Library construction and sequencing were carried out on Illumina HiSeq 2500 platform. To diminish ribosomal and host sequences, raw data were first processed by Trimmomatic (Bolger et al., 2014) and BWA alignment (Li and Durbin, 2009). For downstream virome analyses, clean data were processed with Megahit v1.1.2 (Li et al., 2016) and clustered to unique contigs through CDHIT v4.7 (Fu et al., 2012). Two strategies i.e., database alignment (blast v2.9.0 +) and hidden Markov models (Paez-Espino et al., 2016) were concurrently adopted to identify virus contigs. The RPKM (Reads Per Kilobase per Million mapped reads) value of each contig (Liu et al., 2020) was calculated to determine virus abundance. Information about virus functional genes was acquired through blastp (v2.9.0 +) with UniProtKB/Swiss-Prot database as blueprint. Especially, zoonotic viruses and viruses that exclusively host human were categorized as human viruses in this research. A catalogue comprising sequences annotated as human viruses was constructed as human viral genomes.

2.3. Determination of chemicals from COVID-19 prevention supplies

Four types of chemicals derived from pandemic prevention supplies i.e., total chlorine, trihalomethanes represented by trichloromethane, quaternary ammonium surfactants represented by benzalkyl dimethylammonium compounds, and glucocorticoids represented by methylprednisolone were taken into consideration in this study. Prior to chemical detection, water samples and soil samples treated with ultrasonic were enriched by solid-phase extraction (SPE) (Xu et al., 2019). Subsequently, concentrations of 4 trihalomethanes, 5 quaternary ammonium surfactants, and 40 glucocorticoids were determined by ultra-high-performance liquid chromatography-tandem mass spectrometry (UPLC-MS/MS) (Chen et al., 2021). Detailed information referring to concentration of above chemicals were exhibited in Table S2.

2.4 Estimation of COVID-19 prevention supply.

Consumptions of glucocorticoids and disinfectants were estimated in designated hospitals and WWTPs during COVID-19. According to the COVID-19 designated hospitals in Wuhan, the whole city could be divided to three regions (Fig. 1). In every region, designated hospital and WWTP would respectively be the major consumer of glucocorticoids and disinfectants. Thus, the estimated amount of glucocorticoids and disinfectants could stand for consumption of pandemic prevention supplies in each region.

$$N_{\text{glucocorticoids}} = N_{\text{diagnosed}} \times p \times W \times C_{\text{glucocorticoids}}$$

$N_{\text{glucocorticoids}}$: amount of glucocorticoids used in COVID-19 designated hospital

$N_{\text{diagnosed}}$: number of COVID-19 cases
3. Results

3.1. Amount of pandemic prevention supplies used during COVID-19

To clarify the environmental pressure imposed by excessively used COVID-19 prevention supplies, a rough calculation was drawn to specify the amount of glucocorticoids and disinfectants. The sampling date (2020/3/24) was between the inflection point and zero point of the case curve, representing scenario under impact from prevention supplies during COVID-19 (Fig. 2A). The usage of glucocorticoids form 2020/2/7 to 2020/3/12 exceeded the conventional value before COVID-19, leading to a 1.83 times burden from glucocorticoids (Fig. 2B). The consumption of glucocorticoids in JinYinTan Hospital, HuoShenShan, and LeiShenShan Hospital on sampling date reached up to 6673.68 mg day\(^{-1}\), 11654.76 mg day\(^{-1}\), and 12597.78 mg day\(^{-1}\). To cater to the strict sterilizing demand during COVID-19, the daily amount of disinfectant consumed in Wuhan city rose by 2.12 times. Specifically, the usage of disinfectants in SanJinTan WWTP, CaiDian WWTP, and HuangJiaHu WWTP respectively climbed to 6.36 t day\(^{-1}\), 4.24 t day\(^{-1}\), and 2.12 t day\(^{-1}\).

3.2. Virus community during COVID-19

To comprehensively illustrate the composition of viruses during COVID-19, read-based and denovo methods were simultaneously adopted to identify viral genomes. A final catalog comprising 62,356 DNA contigs and 4,530 RNA contigs were screened under sequencing depth at 30 G per sample. Compared to aquatic viruses (bray-curtis dissimilarity: 0.90), viruses in soil samples were significantly more homogenized (bray-curtis dissimilarity: 0.82; \(p < 0.01\)) with minor families taking up > 90% sequences (Fig. S1). Consequently, aquatic samples were assumed to be more dynamic during COVID-19 and were targeted in following analyses. DNA virus community in water was dominantly occupied by phages i.e., Myoviridae (19.04%), Siphoviridae (18.52%), and, Podoviridae (17.31%) affiliated to Caudovirales (Fig. 3A). Besides, dsDNA Herpesviridae (3.46%) whose members could infect human was ubiquitous in all samples. Aquatic RNA viruses were represented by invertebrate ssRNA(+) (Iflaviridae (21.87%)) and animal ssRNA(+) Arenaviridae (10.37%) (Fig. 3A). Notably, vertebrate ssRNA (+) Coronaviridae was detected in water samples with average

2.4. Data analyses

Relationships between chemicals and viruses were expatiated based on random forest model. Random forest regression models were constructed between chemicals and virus composition on order, family, genus, and contig levels. To a considerable extent, chemicals with high model accuracy were supposed to robustly affect virus community. Response of viruses to chemicals was calibrated as importance i.e., the descent of model accuracy after eliminating this virus. An aggregate of viruses with high importance (increase in mean squared error, IncMSE > 1%) in models (Wright et al., 2020) was prescribed as group that sensitively responded to chemicals. Phylogenetic relationship of viruses was demonstrated by Cytoscape v3.8.0. Principal components analysis was performed to distinguish geographical distribution of virus community regulated by chemicals. Mantel test and Pearson correlation was used to examine the relationship between chemical and virus community (Hu et al., 2020).
abundance at 3.36%.

Focusing on human viral genomes in the environment, an aggregate consisted of 189 contigs subject to 26 species were extracted (Fig. 3B). An overwhelming proportion of human viruses was taken up by 20 species of DNA viruses that incorporated 179 contigs. Among DNA human viruses, Herpesviridae and Poxviridae that embraced 17 species stood out as representatives (Fig. 3B). Specifically, Human gamma-herpesvirus 4 (0.06%) related to various neurological diseases and Mollicute contagiosum virus (0.04%) connected with infectious papules ranked top 2 by percentage. By contrast, Only 6 species with average abundance at 4.47% were recognized in human RNA virus community (Fig. 3B). Guanarito mammarnavirus (10.37%), the hemorrhagic pathogen, preponderated in human disease related RNA virome. To be vigilant, gene segments of SARS-CoV-2 were assembled in samples nearby designated hospitals i.e., HouGuan Lake (0.51%) and HuangJia Lake (36.38%) (Fig. S2) (Table S3).

3.3. Effect of pandemic prevention chemicals on virus community

The variation of virus community was demonstrated under the massive input of COVID-19 related chemicals i.e., total chlorine, trihalomethanes, quaternary ammonium surfactants, and glucocorticoids. Samples from receivers of WWTP effluent and surroundings of designated hospitals were respectively featured by high concentration of trihalomethanes and glucocorticoids (Fig. 4A–D). Correspondingly, the virus community was significantly divergent (adonis: $r = 0.49, p < 0.05$).
with trihalomethanes as one of the vital influencing factors (Fig. 4E). Similar pattern was observed when it comes to human virus community. The geological variation of human viruses was critically shaped by trihalomethanes (adonis: $r = 0.56$, $p < 0.05$) (Fig. 4F). Moreover, significant negative relationships were seen between community diversity and trihalomethanes (richness: $r = -0.70$, $p < 0.05$; Shannon index: $r = -0.87$, $p < 0.001$). Human virus community also became homogeneous with concentration of trihalomethanes elevating (richness: $r$...
= -0.73, \( p < 0.05 \); Shannon index: \( r = -0.85, p < 0.001 \).)

To further verify the impact of chemicals generated from pandemic prevention supplies on virus community, random forest model was conducted on order, family, genus, and contig levels. Concentrations of chemicals were determined by virus composition via regression model in which the accuracy indicated the extent of influence. Models built on contig and phylum levels averagely held the highest (96.7%) and lowest (86.8%) accuracy (Fig. 5A). Regression was most successful when ascertaining the concentration of trihalomethanes (96.4%) (Fig. 5A).

3.4. Response of viruses to pandemic prevention chemicals

To delineate the bidirectional associations, viruses that sensitively responded to chemicals were evaluated. As the most decisive factor (Fig. 5A), trihalomethanes were identified as the major environmental stress to virus community. Thence, 11 contigs affiliated to 9 families were picked out as sensitive viruses to trihalomethanes (Fig. 5B). Among the 11 contigs, 4 types were phages from dsDNA viurs Myoviridae and Siphoviridae. Confronting the disturbance from trihalomethanes, contig_1880145 (%IncMSE = 4.24), a member of algal dsDNA virus Phycooviridae, was the first to bear the brunt, followed by contig_601 (% IncMSE = 4.20) belonged to invertebrate ssRNA(+) virus Iflaviridae.
Notably, 3 human viruses including Human gammaherpesvirus 4 ($p < 0.001$), Orf virus, and Semliki Forest virus ($p < 0.001$) entered the list.

To further elucidate the response of viruses in the environment to COVID-19 related chemicals, relationships between sensitive taxa and trihalomethanes were dissected. Sharp declines of contig_383240 (Mimiviridae) (pearson correlation: $r = -0.79, p < 0.01$) and contig_1061672 (Myoviridae) ($r = -0.67, p < 0.05$) were seen under high concentration of trihalomethanes. Contrarily, a boost in contig_192507 (Siphoviridae) corresponded with the increase of trihalomethanes ($r = 0.67, p < 0.05$). It was noteworthy that trihalomethanes were considerably favorable for the existence of sensitive human viruses i.e., Human gammaherpesvirus 4 ($r = 0.75, p < 0.01$) (Fig. 5C) and Orf virus ($r = 0.83, p < 0.01$) (Fig. 5D).

3.5. Potential human health risk resulted from viruses - pandemic prevention chemicals interactions

The latent hazard to human health from pandemic prevention supplies was distinguished by human viruses - trihalomethanes interaction. Samples in receiving water of WWTP effluent were marked by high concentration of trihalomethanes and relatively barren virus diversity (Fig. 4A–D) (Fig. S3). Whereas, human pathogens tremendously governed virus community in these sites. In HanJiang River that accepted treated medical wastewater from HuoShenShan Hospital, Crimean-Congo hemorrhagic fever virus, a member of Orthonairovirus causing human febrile illness, surprisingly reached the proportion of 98.65%. Receiving wastewater form JinYinTan hospital, FuHe River was notable...
with the prevalence of a human arbovirus - Aroa virus (41.39%) (Table S3). At XunSi River in which LeiShenShan hospital discharged wastewater, the abundance of Orf virus, a zoonotic virus causing contagious erythema, increased by at least 30 times compared to other sites (Table S3). Additionally, human viruses i.e., Human gammaherpesvirus 4 (pearson correlation: $r = 0.75$, $p < 0.01$) (Fig. 5C) and Orf virus ($r = 0.83$, $p < 0.01$) (Fig. 5D), that sensitively responded to trihalomethanes displayed a significant ascend in abundance. The abundance of Human Gammaherpesvirus 4 and Orf Virus multiplicatively increased by 9.90 and 32.06 times per 1 $\mu$g L$^{-1}$ rise in trihalomethanes.

To take a deeper insight into the health threat from human viruses - pandemic prevention supplies connection, genes concerned with replication, recombination and repair were analyzed. Among the chemicals, glucocorticoids were conspicuous for links with several genetic material related genes (Fig. 6A) although their influence on virus abundance was not as strong as other chemicals (Fig. 5A). As a marker gene of virus viability, RNA polymerase gene positively correlated with glucocorticoids ($r = 0.71$, $p < 0.005$). The abundance of genes regulating DNA primase activity (essential genes in replication) ($r = 0.91$, $p < 0.001$), $3'–5'$ exonuclease activity (ensuring the high-fidelity in replication) ($r = 0.98$, $p < 0.001$), and HNH endonuclease (required for genome packaging reactions) ($r = 0.91$, $p < 0.001$) was raised under glucocorticoids. The viability of human viruses represented by the abundance of these genes ascended by 13.34 times with 1 ng L$^{-1}$ rise in glucocorticoids. Additionally, positive correlation between virus gene that encoded DNA lipase, the most fundamental element in duplication, and usage of glucocorticoids ($r = 0.72$, $p < 0.05$) was observed (Fig. 6C). Human viruses i.e., Maccacine alphaherpesvirus 1, Sapporo virus, and SARS-CoV-2 were found to be objects of effect from glucocorticoids because they simultaneously related with above genes as well as glucocorticoids (Fig. 6B). Not coincidently, the concentration of glucocorticoids in Huangjia Lake (500 m from LeiShenShan Hospital) was at least 9.8 times higher than that in other samples while the proportion of SARS-CoV-2 reached peak at 36.38% in this site. Especially, the abundance of Maccacine alpha-
herpesvirus 1 escalated with the expanded usage of glucocorticoids ($r = 0.95$, $p < 0.05$) (Fig. 6D).

4. Discussion

4.1. Trace of SARS-CoV-2 RNA in the environment

RNA segments of novel coronavirus in the water implied the involving of environmental mediums on the transmission route. Unpredictably, RNA fragments of novel coronaviruses distinguished to SARS-CoV-2 were discovered in water near designated hospitals. Distinct from researches that quantified SARS-CoV-2 in raw sewage (Randazzo et al., 2020), wastewater (Trottier et al., 2020), or receiving water of WWTP effluent (Guerrero-Latorre et al., 2020) through RT-qPCR, this work was one of the first researches determining that gene segments of SARS-CoV-2 existed in natural water irrelevant with wastewater discharging. To be vigilant, RNA fragments of SARS-CoV-2 in natural water implied transmission pathway in which aquatic environment served as intermediate host. Presumably, the genetic material of novel coronavirus in water nearby designated hospital might originate from COVID-19 patients through direct contact or bioaerosol (Setti et al., 2020). Considered to be a falciform transmitted virus (Ding and Liang, 2020), SARS-CoV-2 was supposed to persist in water environment (Kumar et al., 2020). Although SARS-CoV-2 could not survive the reinforced disinfection process in WWTPs (WHO, 2020), integrated enveloped particles of SARS-CoV-2 were found to be stable in natural water for 12 days (Wurtzer et al., 2021). Despite the existence, whether SARS-CoV-2 was viable in genuine aquatic condition was vital to evaluate the possibility of dissemination through environment. Attempts were made to assess the infectivity of SARS-CoV-2 in water from aspect of cytopathic effect (Rimoldi et al., 2020). Further endeavor were imperative to evaluate the menace of SARS-CoV-2 in natural environment.

4.2. Succession of virus community initiated by COVID-19 prevention supplies

Chemicals generated from pandemic prevention supplies led to ecological impact through re-constructing virus community. It was deemed that the environment pollution level descended sharply due to the sweeping and strict lockdown (Muhammad et al., 2020), but the secondary pollution from medical treatment and disinfection substances imperiled ecological balance (Bhat et al., 2021). In particular, glucocorticoids from treatment and total chlorine, trihalomethanes, quaternary ammonium surfactants from disinfectants burst into environment through medical and household waste. Microbes in the environment undoubtedly were confronted with the interference of chemicals above, which has been ignored so far. This research shed light on the short-term dynamic of viruses, underlining the ecological damage of pandemic prevention supplies from dimension of environment microbial especially viral community. Generally, the succession of microbial community followed the cycle of collapse, reorganization, exploitation, and conservation (Shabarova et al., 2021). Pathogens i.e., human viruses tended to dominate the community during succession, which was similar in other researches (Rovetta et al., 2013). Apart from the above chemicals, antibiotics were consumed in an inordinate scale with the clinical prevalence at 74.6% (Langford et al., 2021). The residue of antibiotics would inevitably induce antimicrobial resistance (AMR) in the environment even at a trace concentration (Sun et al., 2019). Viruses that mediated horizon gene transfer were proved to participate in the acquisition and dissemination of AMR (Debroas and Siguret, 2019). Therefore, excessively used antibiotics during COVID-19 were likely to interfere with the succession of human viruses via bacteria-virus interaction. Distinguished with the chemical pollution, biological contaminants including human viruses to some extent were more persistent for their ability to proliferate. Although the consumption of non-antiviral drugs i.e., glucocorticoids gradually returned to normal with the clearance of COVID-19 cases, disinfectants were still massively used in the ongoing pandemic (Chu et al., 2021). As a strategy to resist the trihalomethanes that continuously entered environment, it was assumed that the virus community would be dominated by Human gammaherpesvirus 4 (Fig. 5C) and Orf virus (Fig. 5D) in the long time scale. To comprehensively depict the succession process, subsequent monitoring of human viruses in the environment should be attached great importance.

4.3. Excessive disinfection in favor of human virus resistance

Viruses especially specific human viruses whose abundance soared with trihalomethanes withstood side effect of exaggerated disinfection through resistance. High concentration of trihalomethanes in receiving water of WWTP effluent corresponded with excessive usage of disinfectants during COVID-19 (Fig. 4A). Nevertheless, extravagant disinfection was found to be the villain of microbial resistance (Lu and Guo, 2021). Under disinfection, transforms of physiological state, e.g., increasing membrane permeability, SOS response and stress response offered an easier access for microorganisms to resistance (Cai et al., 2021). Equipped with horizontal transfer, viruses served as antibiotic resistance gene pool (Debroas and Siguret, 2019). Gene that specifically methylated the adenine in position 37 of tRNA (1) favored resistance under oxidative stress (Golovina et al., 2009). In this research, this gene was detected in virus genome and enriched in receiving water of WWTP effluent. Also, the long-term exposure of disinfection byproduct was in favor of gene mutation (Kurasam et al., 2018). Under the induction of trihalomethanes during COVID-19, it was possible for Human gammaherpesvirus 4 and Orf virus that strongly correlated with trihalomethanes (Fig. 5 C-D) in this study to obtain resistance through gene mutation or horizontal transfer. Moreover, herpesvirus (Bello-Morales et al., 2020) as
well as Orf virus (Lan et al., 2016) were discovered to unite as vesicles. Depending on this morphology, viruses could efficiently combat with adversaries e.g., disinfection (Zhang et al., 2021). Despite the prevalence of Human gammaherpesvirus 4 (Young and Rickinson, 2004) and Orf virus (Zhang et al., 2014), few researches focused on their distributions in natural environment. This pattern needs to be inspected in a broader geographical range. The mechanism lied behind the boosting relationship between trihalomethanes and human viruses remains to be further elaborated through microcosm simulation experiment.

4.4. Glucocorticoids advanced the proliferation of human viruses

Glucocorticoids were potential promoting factors to viability of human virus. Genomes recognized as SARS-Cov-2 RNA taken up a staggering proportion of 36.38% in HuangJia Lake (500 m from LeiShenShan Hospital). Meanwhile, concentration of glucocorticoids peaked 4.4. Glucocorticoids advanced the proliferation of human viruses elaborated through microcosm simulation experiment.

ship between trihalomethanes and human viruses remains to be further elaborated through microcosm simulation experiment. Meanwhile, concentration of glucocorticoids peaked 4.4. Glucocorticoids advanced the proliferation of human viruses elaborated through microcosm simulation experiment.

4.5. Threat from human viruses in the environment

To practically evaluate the human health risk from viruses promoted by COVID-19 prevention chemicals, several factors should be taken into consideration in the future research. 1) Source of contaminants. Besides WWTP system that were proved to be hotspot of diverse human viruses (Zhang et al., 2018), surroundings of hospitals where the viability (Fig. 4D) (Fig. 6A–B) of human viruses were high should also be monitored. 2) Persistence and viability of human viruses in the environment. It is necessary to figure out the decay pattern of human viruses in the environment when assessing related health risk. Taking SARS-CoV-2 as an example, the infection of mere RNA was arguable even though it could linger in environment for weeks (Ahmed et al., 2020). 3) Propagation of human viruses in environmental matrices. Exposure pathway of human viruses hinge on their dissemination and circulation in environmental matrix. Viruses were enriched (Adriaenssens et al., 2021) while the infectivity decreased (Fauvel et al., 2017) with water flow. Similarly, a more diverse human virus community was observed in river that received WWTP effluent than in upstream lake (Fig. S3). 4) Integration of epidemiological data. Epidemiological information is the communication might be advantageous for SARS-CoV-2 to retain (Ma et al., 2020). Thus far, rare researches had referred to glucocorticoids - virus relationship in aquatic environment. On the days when hormone residue in environment has become a widespread problem (Zhong et al., 2021), this research enlightened the environmental interaction between virus and glucocorticoids.

5. Conclusion

This research focused on characteristics of environmental viruses under high-intensity use of pandemic prevention supplies during COVID-19 based on virome. Combined with random forest model, mechanisms between viruses and pandemic prevention chemicals were systematically probed on community, species, and gene levels. The abundance and viability of specific human viruses were potentially promoted by trihalomethanes and glucocorticoids, leading to an overlooked threat to human health. Following up studies should also be launched to assess the subsequent dynamics of virus community and tracking the community succession process. With the vision of environmental virology, this research disclosed the environmental damage under unconventional anthropogenic activities i.e., excessive usage of pandemic prevention supplies. Scaling up to a boarder extent, this work serve as foundation for formulating ecological management policy in the ongoing pandemic and emphasized the necessity of regular human virus monitoring in the environment.

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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Author contributions

Z.C.H., L.H.Y. and J.H. conducted the sampling. Z.C.H. L.H.Y. and Y. H.J. carried out experiments. Z.C.H. and Z.S.L. analyzed study data. Y.X. Z. collected environmental data. Z.C.H. visualized research data. Z.C.H. and L.H. conceived of the study. Z.S.L. and B.L.H. reviewed and edited the original draft. L.Z.Z. offered fund support. B.L.H. supervised the research.

Appendix A. Supplementary material

Supplementary data to this article can be found online at https://doi.org/10.1016/j.envint.2022.107192.

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