The presence of SARS-CoV-2 in raw and treated wastewater in 3 cities of Iran: Tehran, Qom and Anzali during coronavirus disease 2019 (COVID-19) outbreak

Simin Nasseri 1,2 · Jila Yavarian 3 · Abbas Norouzian Baghapi 1 · Talat Mokhtari Azad 3 · Ahmad Nejati 3 · Ramin Nabizadeh 1 · Mahdi Hadi 2 · Nazanin Zahra Shafiei Jandaghi 3 · Behnam Vakili 4 · Seyed Koushyar Azam Vaghefi 5 · Mahtab Baghban 6 · Somayeh Yousefi 1 · Shahrokh Nazmara 1 · Mahmood Alimohammadi 1,2,7,8

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Abstract
This study aimed to identifying the presence of SARS-CoV-2 RNA in raw and treated wastewater during the COVID-19 outbreak in Tehran, Qom and Anzali cities (Iran). From three wastewater treatment plants (WWTPs), 28 treated and untreated wastewater composite samples were collected from April 4 to May 2, 2020. In this study, polyethylene glycol 6000 (PEG 6000) was used through one-step real-time quantitative reverse transcriptase-polymerase chain reaction (qRT-PCR) for identification of RNA viruses. SARS-CoV-2 RNA was elicited from wastewater composite samples in all inlet samples taken from the three above mentioned cities. The results of outlet samples were as follows: 1) Results from Qom and East Anzali outlets showed no trace of SARS-CoV-2 RNA despite the difference in treatment disinfection method used (chlorine vs. ultraviolet (UV) disinfection). 2. In Tehran, SARS-CoV-2 RNA was not detected in any of the outlet samples taken from the modules disinfected by UV. Out of the four samples taken from the modules disinfected by chlorine, two were positive for the SARS-CoV-2 RNA which could have been caused by deficiencies in operation and maintenance. It can be concluded that meeting the standards of operation and maintenance (O&M) in WWTPs can considerably ensure that wastewater does not act as one of the roots of transmission for the disease.

Keywords Wastewater treatment · COVID-19 · SARS-CoV-2 · Disinfection · Pandemic

Introduction
The rapid outbreak of coronavirus disease 2019 (COVID-19) caused by the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) that is spread via airborne, droplet, contact, and fecal-oral transmission modes constitutes a worldwide and disruptive challenge to societies [1, 2]. Both symptomatic and asymptomatic carriers have been recognized to transmit the SARS-CoV-2 virus via various modes of transmission [3–6]. Causing severe respiratory infections among patients receiving medical care in hospitals [7–12]. Hence, the COVID-19 outbreak has been declared a global public health

1 Department of Environmental Health Engineering, School of Public Health, Tehran University of Medical Sciences, Tehran, Iran
2 Center for Water Quality Research (CWQR), Institute for Environmental Research (IER), Tehran University of Medical Sciences, Tehran, Iran
3 Department of Virology, School of Public Health, Tehran University of Medical Sciences, Tehran, Iran
4 Office of Improvement on Wastewater Operation Procedures, National Water and Wastewater Engineering Company, Tehran, Iran
5 Manager of Water Quality Control Bureau, National Water and Wastewater Eng. Co., Tehran, Iran
6 Reference Laboratory of Water and Wastewater, Tehran Province of Water and Wastewater Company, Tehran, Iran
7 Health Equity Research Center (HERC), Tehran University of Medical Sciences, Tehran, Iran
8 Department of Environmental Health Engineering, Health Faculty, Tehran University of Medical Sciences, Tehran, Iran
emergency of international concern by the World Health Organization (WHO) [13–15]. The original source and the way of transmission to humans is the subject of current studies [6, 14, 16–18].

Usual human coronaviruses, containing kinds alpha coronavirus and beta coronavirus, commonly create mild to moderate and upper respiratory tract infections (URTI), similar to the common cold disease [19–22], while coronaviruses, containing the SARS-CoV-2 (a new coronavirus) has created severe respiratory illness and even death in some patients [16–18, 20, 23–25]. In addition, it should be noted that SARS-CoV-2 belongs to beta-CoVs and its size is between 60 and 140 nm [26].

In addition, the monitoring of viruses and bioaerosols in wastewater and water have been reported many years ago [27, 28]. For example, former studies have confirmed the presence of rotavirus, measles virus and poliovirus in water and wastewater [29, 30]. Besides, previous studies have shown that coronaviruses can exist and maintain viable in sewages originated from the fecal discharge of infected patients [31, 32]. In the case of SARS-CoV epidemic in March 2003, there had been a potential link to water and sewage, and more than 300 people in Hong Kong had been infected with the virus through a faulty sewage system [33], due to SARS-CoV which could multiply the intestinal tract with possibility of gastrointestinal transmission of the virus [34].

In the case of SARS, Chan et al. (2004) reported that the symptom of diarrhea was observed between 8 and 73% in patients, and they declared the raising concerns about the potential of environmental transmission of the virus [34]. Furthermore, Liu et al. (2004) and reported that the SARS-CoV virus was also cultured from the feces of SARS-CoV patients up to 3 weeks after infection [35]. In addition, the recent works declared that coronaviruses can also remain infectious for a few days in sewage [36, 37].

The transmission of SARS-CoV-2 is still unknown, while according to previous works, the main way of transmission of SARS-CoV-2 is human-to-human [14, 38]. Furthermore, it is not yet known whether the SARS-CoV-2 was first transmitted to humans through animals or through virus-infected surfaces; however, transmission of the virus from surfaces is also another way of transmitting the virus [39]. Besides, presence of SARS-CoV-2 RNA in the feces of COVID-19, demonstrates that transmission can potentially create by fecal-oral routes [40–42].

Hence, although direct transmission through respiratory aerosols, where the persons are in close contact and contact with contaminated surfaces, are considered as the most important routes of transmission [43–46], there is still generally limited information about the potential for transmission through the environment like water and wastewaters. However, some researchers reported that sewage/domestic wastewater surveillance could be a sensitive tool to monitor the circulation of the virus in the population [30, 47–49]; For example, Ahmed et al. (2020) studied on the wastewater in a catchment in Australia for identification of SARS-CoV-2 RNA [50]. The results of their work described that RNA copies were counted in two positive wastewater samples in a six-day period by reverse transcriptase quantitative polymerase chain reaction (RT-qPCR) [50].

In addition, whereas the new virus has already begun to spread worldwide, it is important for water engineers and professionals to understand the nature and fate of coronavirus after discharging into aquatic environments and the effective measures to protect public health. Due to the outbreak of coronavirus disease 2019 (COVID-19) in Iran and lack of sufficient knowledge and information about the environmental persistence of this the RNA viruses in wastewater in this country, it is important to conduct research on the identification of the RNA viruses in raw and treated wastewater in main epicenters of the country. Moreover, the use of raw and treated wastewater for agriculture in Iran is a common practice in many cities especially the COVID-19 epicenters cities, including Tehran, Anzali and Qom. Therefore the presence of this the RNA viruses in produced wastewater of such cities may be associated with the possibility of its presence in agricultural produces and make it possible for the virus to be transmitted to humans.

The basic novelty of this research is that of the presence of SARS-CoV-2 RNA in raw and treated municipal wastewater in the most densely populated cities of Iran (Tehran, Anzali and Qom) with high outbreak of COVID-19. Hence, the goal of this work is multi-fold: 1) study the presence of SARS-CoV-2 RNA in inlets (raw municipal wastewater) and outlets (treated municipal wastewater) of three wastewater treatment plants (WWTPs) in the three cities of Iran: Tehran, Qom and Anzali; 2) investigate the impact on the composition of wastewater on SARS-CoV-2 RNA; 3) study the impact on disinfection (UV and chlorine) on the SARS-CoV-2 RNA in treated municipal wastewater (outlets of wastewater treatment plants (WWTPs)). The results of this work can have broad implications for other regions owing to the pervasiveness of COVID-19.

Materials and methods

Chemicals

Sodium chloride (NaCl) (Merck: 7647-14-5), sodium hydroxide (NaOH) anhydrous pellets (CAS Number: 1310-73-2), dextran sulfate sodium salt (Sigma-Aldrich: Lot Number, 8C8R4068V), polyethylene glycol (PEG 6000) (Sigma-Aldrich: CAS Number: 81255) were purchased and used for identification of SARS-CoV-2 RNA from wastewater samples. In addition, SPL Centrifuge Tubes (falcon tubes
50 mL SPL Life Sciences Co-50,050) was applied for sample preparation.

**Study area**

In this study, sampling were performed in three WWTPs in the three cities of Iran: Tehran, Qom and Anzali (Fig. 1).

The southern Tehran wastewater treatment plant (southern Tehran WWTP)

Tehran is the capital city of Iran with 22 distinct geographic regions and it is the largest city in this country. The population of Tehran according to Statistical Centre of Iran (SCI) is over 13 million [51]. This city is located in latitude of 35.6892° N and longitude of 51.3890° E (Fig. 1) [52–54]. Sampling was performed in inlets (untreated/raw municipal wastewater) and outlets (treated municipal wastewater) of Southern Tehran WWTP. The Southern Tehran WWTP is placed in the southeast of Tehran, Iran (in latitude of 35.5724° N and longitude of 51,5539° E) (Fig. 1) [51, 55]. The Southern Tehran WWTP is planned for three phases for treatment of wastewater with a capacity of $4.2 \times 10^6$ people. The first phase of this WWTP (i.e., from module one to four) the extended aeration activated sludge (EAAS) system. The second phase (i.e., from module five to six) is planned using the oxidation ditch method.

![Fig. 1 The locations and sampling sites of each WWTP: A: East Anzali WWTP, B: Southern Tehran WWTP and C: Qom WWTP No. 3](image-url)
Another phase (i.e., from module seven to eight) will be established in the future [55]. In addition, it should be noted that sampling were performed in the first phase (i.e., one to four modules) and the second phase (i.e., from module five to six). The major specifications and design parameters of this WWTP are shown in Table 1.

East Anzali wastewater treatment plant (east Anzali WWTP)

Anzali city, also known as Bandar-e Anzali, is located in the north of Iran, in Gilan Province. The population of Anzali based on Statistical Centre of Iran (SCI) in 2016 was over 118,564. This city is located in latitude of 37.4639° N and longitude of 49.4799° E (Fig. 1). Sampling was performed in inlets (untreated/raw municipal wastewater) and outlets (treated municipal wastewater) of East Anzali WWTP. This WWTP is located in the east of Anzali city, (in latitude of 35.8055° N and longitude of 41.4831° E) (Fig. 1). The major specifications and design parameters of East Anzali WWTP are shown in Table 1, an activated sludge process in used. Activated sludge of this WWTP is the biological wastewater treatment process, that comprises two grit and grease removal, two circular primary sedimentation tanks, a wastewater pumping station, two biological units, two secondary sedimentation tanks and two coagulation unit tanks, four gravity sand filter units, and a UV disinfection unit.

Qom wastewater treatment plant no. 3 (Qom WWTP no. 3)

Qom city (is) situated 140 km to the south of Tehran (34.6416° N, 50.8746° E). Qom is the seventh largest city of Iran (Fig. 1). The population of Qom according to Statistical Centre of Iran (SCI) in 2016 was 1,201,158. Sampling was performed in inlet and outlet of Qom WWTP No. 3, like both previous WWTPs. Target WWTP is placed in the east of Anzali city, (in latitude of 35.8055° N and longitude of 41.4831° E) (Fig. 1). The major specifications and design parameters of East Anzali WWTP are shown in Table 1, an activated sludge process in used. Activated sludge of this WWTP is the biological wastewater treatment process, that comprises two grit and grease removal, two circular primary sedimentation tanks, a wastewater pumping station, two biological units, two secondary sedimentation tanks and two coagulation unit tanks, four gravity sand filter units, and a UV disinfection unit.

Untreated and treated wastewater sampling in three WWTPs

In this study, sampling points were selected in inlets (untreated/raw municipal wastewater) and outlets (treated municipal wastewater) of three WWTPs in the three cities of Iran: Tehran, Qom and Anzali. Due to the COVID-19 outbreak in these cities the first cases were found in Qom and regarding the RNA viruses entrance into, sampling in inlets (raw municipal wastewater) and outlets (treated municipal wastewater) of three WWTPs can be a representative of the covered population according to pervious works [50, 56]. For these reasons, taking composite samples of inlet and outlet of each WWTPs could be considered to be sufficient to estimate the probability of the presence or absence of the RNA viruses in the municipal wastewater. Sampling was performed in spring between April 4, 2020 and April 17, 2020 (a two-week period) and between April 20, 2020 and May 2, 2020 (a two-week period). Twelve composite untreated wastewater samples with a frequency of one composite sample each week were collected from each WWTPs during a two-week period (from April 4, 2020 to April 17, 2020). Hence, we collected a total of 12 untreated/raw wastewater samples during a two-week period owing to three composite sample at the 3 sites ((3 sites +3 inlets) × a two-week period = 12) (Table 2).

The same exact mode of operation used to the treated wastewater sampling, yielding 16 total treated wastewater samples (Note: the inlet Southern Tehran WWTP is divided into two outlets (Tables 1 and 2)). As there were three WWTPs, we collected 28 total treated and untreated wastewater samples altogether.

Wastewater samples were moved on ice to the laboratory and kept at 4 °C until further analysis. It should be emphasized that authors have used standard personal protective equipment (PPE) for wastewater sampling and analyzing samples in laboratory, like N95 mask, steel capped boots, face shields, hard hats, long pants safety glasses/goggles and gloves. A summary of the locations and sampling sites in each of the WWTPs is shown in Fig. 1.

Preparation of sewage samples using the phase-separation method (preparation of half a liter of sewage sample)

The concentration of sewage samples for the isolation of viral RNA were performed based on WHO guidelines [27, 57, 58]. The preparation of the sewage samples and its inoculation into cell culture should be done in two consecutive days [27, 57]. The first day includes three steps, while the second day includes two steps.

The first day, which includes three steps:

Step 1: Sample preparation

At the first, sewage sample was allowed to sediment for five minutes for removing large solid materials. Then, about 500 mL of the sample was divided into two 250 mL centrifuge bottles within a biosafety cabinet (class II). Those bottles were centrifugated at 1500 g for 20 min at +4 °C. After that, the supernatant was poured in a 1 l beaker. In addition, for further using, the closed
### Table 1 The major specifications and design parameters of Southern Tehran WWTP, East Anzali WWTP and Qom WWTP No. 3

#### Southern Tehran WWTP

| Parameters                              | Numbers of samples | Unit |
|-----------------------------------------|--------------------|------|
| Number of inhabitant served             | 4.2 million people  |      |
| Maximum flow                            | 450,000 m³/day     |      |
| Mean flow                               | 18,750 m³/h        |      |
| A capacity of each module               | 525 thousand people|      |
| Wastewater treatment process (1–4 modules) | Extended aeration activated sludge (EAAS) system |      |
| Wastewater treatment process (5–6 modules) | Oxidation ditch method |      |
| Wastewater treatment process (7–8 modules) | Will be established |      |
| Average inlet Biochemical Oxygen Demand (Inlet-BOD5) | 31 | 215 mg/L |
| Average outlet Biochemical Oxygen Demand (Outlet-BOD5) | 31 | 24 mg/L |
| Average inlet Chemical Oxygen Demand (Inlet-COD) | 31 | 385 mg/L |
| Average outlet Chemical Oxygen Demand (Outlet-COD) | 31 | 58 mg/L |
| Average inlet Suspended Solids (Inlet-SS) | 31 | 179 mg/L |
| Average outlet Suspended Solids (Outlet-SS) | 31 | 30 mg/L |
| *Disinfection unit for 1–4 modules       | Chlorine disinfection |      |
| *Disinfection unit for 5–6 modules       | Ultraviolet (UV) disinfection |      |
| Wastewater treatment process (7–8 modules) | Will be established |      |
| Application of the wastewater effluent   | Irrigate the Varamin plain and aquifer recharge |      |

#### East Anzali WWTP

| Parameters                              | Numbers of samples | Unit |
|-----------------------------------------|--------------------|------|
| Number of inhabitant served             | 40 thousand people  |      |
| Mean daily inlet flow                   | 7800 m³/day        |      |
| Mean monthly flow                       | 241,800 m³/month   |      |
| Wastewater treatment process (Inlet-BOD5) | Extended aeration activated sludge (EAAS) system |      |
| Average inlet Biochemical Oxygen Demand (Inlet-COD) | 30 | 180 mg/L |
| Average outlet Biochemical Oxygen Demand (Outlet-BOD5) | 30 | 10 mg/L |
| Average inlet Chemical Oxygen Demand (Inlet-COD) | 30 | 331 mg/L |
| Average outlet Chemical Oxygen Demand (Outlet-COD) | 30 | 19 mg/L |
| Average outlet Suspended Solids (Outlet-SS) | 30 | 170 mg/L |
| **Disinfection unit                     | Chlorine disinfection |      |
| Application of the wastewater effluent   | Discharged into Anzali lagoon |      |

#### Qom WWTP No. 3

| Parameters                              | Numbers of samples | Unit |
|-----------------------------------------|--------------------|------|
| Number of inhabitant served             | 249 thousand people|      |
| Mean daily inlet flow                   | 51,756 m³/day      |      |
| Mean monthly flow                       | 1,542,354 m³/month |      |
| Wastewater treatment process (Inlet-BOD5) | Conventional Activated Sludge (CAS) system with diffuser aeration |      |
| Average inlet Biochemical Oxygen Demand (Inlet-COD) | 31 | 401 mg/L |
| Average outlet Biochemical Oxygen Demand (Outlet-BOD5) | 31 | 36 mg/L |
| Average outlet Chemical Oxygen Demand (Outlet-COD) | 31 | 254 mg/L |
| Average inlet Suspended Solids (Inlet-SS) | 31 | 10 mg/L |
| Application of the wastewater effluent   | Agriculture (mainly irrigation) |      |

*It should be noted that sampling were performed in the first phase (i.e., one to four modules) and the second phase (i.e., from module five to six).

**It should be noted that sampling were performed in this unit.
centrifuge bottles containing the solid pellet were kept in the refrigerator at 4 °C [27, 57].

**Step 2: Adjusting the pH of supernatant**

At first, the pH solution within a 1 l beaker were adjusted by 1 N HCl and 1 N NaOH (pH = 7.0–7.5). Then, a 500 mL of adjusted supernatant by HCl and NaOH was transferred to a 1 l beaker for further study. The rest of the supernatant and the raw sewage samples were stored as a backup sample at 4 °C until the cells inoculated with the concentrate under the microscope showed no toxicity. Finally, after ensuring that the samples were not toxic, they were stored at 20 °C until the full test result is known [27, 57].

**Step 3: Glass separatory funnel with different separation phases**

At first, a 39.5 mL of 22% dextran, 287 mL of 29% polyethylene glycol (PEG 6000), and 35 mL 5 M NaCl were added to 500 mL of the supernatant obtained from the previous step, respectively. Then, the mixture obtained at this stage was stirred with sufficient speed using a magnetic stirrer (vortex formation) for one hour at +4 °C. After that, a sterile one liter conical glass separatory funnel were prepared for each samples. The funnel stood on the foundation. A very thin layer of the stopcock grease was prepared on the sliding glass surface of the valves but did not clog the holes (note: Teflon valves did not need grease). Finally, the stirred mixture was slowly poured into a 1 L separatory funnel and then kept at +4 °C overnight for separating mixture to three phases (upper phase, interphase and lower phase). Polyethylene glycol created the more hydrophilic, denser and lower phase, whereas dextran created the more hydrophobic, less dense and upper phase (Fig. 2) [27, 57, 58].

The second day, which includes two steps:

**Step 1: Interphase separation**

In the second day, the bottom of the glass separatory funnel was carefully checked. The lower and intermediate phase (interphase) were seen with solids. The entire lower phase and the interphase were poured firstly dropwise into a sterile 50 mL centrifuge tube. Then, the combined volume (lower phase and the interphase) was measured and recorded. In this step, the volume of the solid precipitate was recorded in the lower layer and/or in the interphase, whether it was more than half a mL. It should also be pointed out that when the interphase was too thick, it may be collected separately, because it will be hard to gather. In addition, it should be noted that once the interphase layer was collected, the glass separatory funnel and upper phase were disinfected using appropriate disinfectants (such as sodium hypochlorite (NaOCl)) for one hour [27, 57].

**Step 2: Adding pellet to the interphase solution**

In this step, 5 mL of the interphase solution under a biosafety cabinet (class II) was added to a sterile centrifuge tube which contained pellets and dissolved well. Then, this dissolved solution (containing the pellet) was returned to the interphase solution, prepared in step one in the second day. Hence, the final prepared solution of interphase was prepared and used for RNA extraction [27, 57]. In addition, it should be noted that in case which the sewage samples are very clean, it helps to add a small amount of solid from pallet to facilitate the identification of the interfaces in the glass separatory funnel.

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Table 2  Detection of SARS-CoV-2 RNA in untreated (raw) and treated wastewater samples in Southern Tehran WWTP, East Anzali WWTP and Qom WWTP No. 3

| Sampling (Inlet) | Sampling date | Types of samples | Presence (+) or absence (−) of SARS-CoV-2 RNA |
|-----------------|---------------|------------------|-----------------------------------------------|
| Untreated wastewater | April 4, 2020 | Composite sample | + (Southern Tehran WWTP: Qom WWTP No. 3 East Anzali WWTP) |
| Untreated wastewater | April 17, 2020 | Composite sample | + |
| Untreated wastewater | April 20, 2020 | Composite sample | + |
| Untreated wastewater | May 2, 2020 | Composite sample | + |

| Sampling (Outlet) | Sampling date | Types of samples | Presence (+) or absence (−) of SARS-CoV-2 RNA |
|-------------------|---------------|------------------|-----------------------------------------------|
| Treated wastewater | April 4, 2020 | Composite sample | – (1-4 modules: Chlorine) |
| Treated wastewater | April 17, 2020 | Composite sample | – |
| Treated wastewater | April 20, 2020 | Composite sample | + |
| Treated wastewater | May 2, 2020 | Composite sample | + |

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One-step real-time PCR analysis

Total RNA was extracted from 200 μL of the concentrated samples using FastPure Viral RNA Mini Kit (Vazyme-China) according to the manufacturers’ procedures. Briefly, the sample was mixed with lysis buffer and then loaded onto the filter membrane column. Then, the filter column was washed twice with washing solution. Finally, the RNA was eluted in 50 μL of RNase free water and stored at −80 °C until required.

For qualitative detection of SARS-CoV-2, One-Step Taqman Real-Time PCR was used to amplify the ORF1ab and N gene according to Sansure SARS-CoV-2 Nucleic Acid Test Kits (Sansure Biotech- China). To confirm extraction of samples, the RNase P gene was selected as Endogenous control and positive and negative samples were included in each PCR reaction (Fig. 2).

The final PCR volume reactions were performed in 50 μL reaction mixtures containing 4 μL of enzyme mix, 26 μL PCR Master mix contain primers and probe and 20 μL RNA template. Amplification was carried out under the following temperature program: cDNA synthesis at 50 °C for 30 min, preheat at 95 °C for 1 min, followed by 45 two-step cycles of 95 °C for 15 s and 60 °C for 30s on Step One Plus™ Real-Time PCR System (Applied Biosystems, California, USA) instrument. The cycle threshold (Ct value) was calculated automatically.

Results and discussions

In this study, Taq-man Multiplex Real-Time PCR was used to detect SARS-CoV-2 RNA. The Kit targets ORF1ab and N genes (Sansure Biotech- China) are used for detection of SARS-CoV-2 RNA. The N gene encoded nucleocapsid that surrounded the genomic RNA and the orf1ab gene encodes RNA-dependent RNA polymerase required for viral RNA replication. This kit provides positive and negative controls to ensure correct kit performance. In this kit, human RNase P was used as an endogenous control gene. The use of endogenous control indicates the validation of RNA extraction and presence of inhibitors in the RNA sample. Plus human RNase P gene was taken as internal and extraction control (Fig. 3). Furthermore, amplification plot for one positive sample with positive and negative controls are shown in Fig. 3. Accordingly, a positive sample is a sample that both orf1ab and N genes are positive. In addition, it should be noted that newly reported Taq-man Multiplex Real-Time PCR has been utilized for finding of human coronaviruses such as SARS-CoV-2 RNA and human respiratory syncytial virus (HRSV) in samples of wastewater and hospitalized children [59–61].

Detection of SARS-CoV-2 RNA in wastewater samples

East Anzali WWTP

In the case of East Anzali WWTP, among two wastewater samples collected in raw wastewater (inlets) during a two-week period, two samples (100%) were positive for SARS-CoV-2 RNA using Taq-man Real-Time PCR method (Table 2). Besides, the results of four collected outlet (treated) samples from chlorine disinfection of East Anzali WWTP on April 20, 2020 and on May 2, 2020 showed that none of these samples were positive for SARS-CoV-2 RNA. Recent work has described that 71% gathered untreated wastewater samples (5 from 7 samples) from seven treatment plants in Istanbul (Turkey) was positive for SARS-CoV-2
RNA using polyethylene glycol 6000 (PEG 6000) adsorption SARS-CoV-2 concentration method, which the amount of SARS-CoV-2 in seven untreated wastewater samples ranged from $1.80 \times 10^4$ (for Pasakoy WWTP) to $8.26 \times 10^3$ (for Ambarli WWTP) [62]. Moreover, a study of Gonzalez et al. (2020), a three RT-ddPCR assays (N1, N2, N3) were applied to identify SARS-CoV-2 RNA in weekly samples from 9 WWTPs in Virginia (a southeastern U.S. state) [63]. They found that during the twenty-one week study, SARS-CoV-2 concentrations ranged from $10^1$ to $10^4$ copies $100\text{ mL}^{-1}$ in samples where viral RNA was measured [63].

Qom WWTP no. 3

The results of raw wastewater samples gathered from the Qom WWTP No. 3 were positive for SARS-CoV-2 RNA. However, four collected outlet (treated) samples from chlorine disinfection of Qom WWTP No. 3 during a two-week period showed that these samples were negative (Table 2). Likewise, Kocamemi et al. (2020) collected two composite samples from sewage manholes placed close to the pandemic hospitals in in Istanbul (Turkey), which both samples were examined positive (100%). In addition, the amount of SARS-CoV-2 in sewage manholes ranged from $4.49 \times 10^3$ to $9.33 \times 10^4$ [62]. In addition, Cascella et al., (2020) declared that SARS-CoV-2 virus can be efficaciously inactivated using chlorine, ultraviolet rays, heat, and ethanol, which is consistent with our results [26]. Furthermore, Wurtzer et al. (2020) collected twenty three raw and eight treated wastewater samples from 3 main WWTP of the Parisian area and they found all raw wastewater samples scored positive for SARS-CoV2 [64]. In addition, they reported that six out of eight samples from treated wastewater scored positive by RT-qPCR [64]. Moreover, treated wastewater effluents displayed a 100 times reduction in the viral load compared to the corresponding raw wastewater samples [64]. Additionally, previous work studied the presence of SARS-CoV-2 RNA in untreated and treated wastewater samples in southern Louisiana (USA). SARS-CoV-2 RNA was identified in two out of fifteen wastewater samples using two reverse transcription-quantitative polymerase chain reaction (RT-qPCR) assays (CDC N1 and N2) [65]. Trottier et al. (2020) identified the amount of SARS-CoV-2 RNA at the inflow point of the major WWTP of Montpellier, France [66]. They found that amounts of SARS-CoV-2 RNA at the WWTP from mid-June on started increasing [66]. Generally, results of past works showed that the SARS-CoV-2 RNA existed in raw and treated wastewaters [67, 68].

Southern Tehran WWTP

Among two wastewater samples collected in inlets of Southern Tehran WWTP during a two-week period, two samples (100%) was positive for SARS-CoV-2 RNA by the Taqman Real-Time PCR (Table 2). Furthermore, we collected two outlet samples from chlorine disinfection (module outlet 1 to 4) and two samples from ultraviolet (UV) disinfection (module outlet 5 to 6) of Southern Tehran WWTP during a two-week period and the results showed that only two samples collected from chlorine disinfection unit were positive (Table 2).

For comparison, previous investigation reported that 22% collected samples (two from nine samples) from WWTP in Southeast Queensland (Australia) were positive for SARS-CoV-2 RNA by N_Sarbeco method [50]. In addition,
Wurtzer et al. (2020) collected untreated wastewater samples in WWTP of Parisian area in Paris (France) that the results of their study showed the concentration of SARS-CoV-2 RNA in untreated wastewater samples was about $5.10^4$ genome units/L [69] and the results of the same work concluded that the increase of genome units in untreated wastewater identically pursued the increase of human SARS-CoV-2 cases showed in the study area, which is consistent with the results of our study [69]. Hence, the outbreak of COVID-19 created by SARS-CoV-2 virus has been dispersed swiftly around the world [70, 71]. Previous works reported that the COVID-19 patients can excrete SARS-CoV-2 virus into wastewater via gastrointestinal tract (feces) and those can survive from hours to days in untreated wastewater [30, 50, 69]. In addition, Bogler et al. (2020) reported that SARS-CoVs were present in wastewater for several days, leading to potential health risks via waterborne and aerosolized wastewater pathways [72]. In addition, whereas SARS-CoV-2 virus can be identified in feces samples, Wurtzer et al. (2020) has recently suggested that SARS-CoV-2 RNA in wastewater can be used as a supplementary tool to inspect novel coronavirus circulation in human populations [69]. Hence, the identification of SARS-CoV-2 RNA in wastewater makes it possible to monitor the spread of infections among the population through wastewater-based epidemiology [50, 56].

In addition to the aforementioned, according to results of outlet samples of Southern Tehran WWTP described that the UV disinfection was more effective than chlorine disinfection and for effective disinfection, WWTP operator should increase the concentration of free residual chlorine to amounts greater than or equal to 0.5 mg/L at retention time (RT) thirty minutes at pH lower than eight, which is in agreement with technical brief reported by WHO on March 03, 2020 [73]. Furthermore, based on the recent survey, the UV disinfection has numerous benefits such as formation of no disinfection byproducts, short retention time, effective on a wide range of resilient viruses, and no residues compared with chlorine disinfection [74].

Regarding that in the three WWTPs, activated sludge (AS) processes (biological wastewater treatment process) are used, hence these systems have the ability to remove SARS-CoV-2 RNA, especially WWTP that have suitable disinfection systems (as can be seen in Table 2 and Fig. 4) and the removal efficiency of SARS-CoV-2 RNA in Southern Tehran WWTP, East Anzali WWTP and Qom WWTP No. 3 were 50%, 100% and 100%, respectively.

On the other side, we should not forget that proper sludge management of those WWTPs are essential due to the sludge of these treatment plants can contain active viruses [62, 75–77]. For example, past work described that nine Human coronavirus 229E and one Human coronavirus HKU1 were identified from class B bio-solids in an anonymous US wastewater treatment facility [78]. In addition, during the COVID-19 outbreak, using of untreated wastewater and even treated wastewater in agricultural irrigation poses a serious threat to all members of society, especially in the cities of developing countries.

As a result, it can threaten the health of workers at WWTPs during sludge production, transportation and disposal, and the residents living near the treatment plants and even entire communities. Finally, according to the findings of this research work, it should be emphasized that workers at WWTPs should wear standard personal protective equipment (PPE), like N95 mask, steel capped boots, face shields, hard hats, long pants safety glasses/goggles and gloves.

Table. 2. Detection of SARS-CoV-2 RNA in untreated (raw) and treated wastewater samples in Southern Tehran WWTP, East Anzali WWTP and Qom WWTP No. 3.

**Impact on the composition of wastewater on SARS-CoV-2 RNA**

As reported in the previous work, it is essential to evaluate the effects of the composition of wastewater and meteorological parameters on SARS-CoV-2 RNA during the outbreak [50]. Hence, for this reason we measured important factors such as BOD$_5$, COD and SS in inlet and outlets of three WWTPs. The composition of wastewater in inlets and outlets of Southern Tehran WWTP, East Anzali WWTP and Qom WWTP No. 3 are described in Fig. 4, and it can be seen in this figure that, the high outlet BOD$_5$, COD and SS values in Southern Tehran WWTP compared to East Anzali WWTP and Qom WWTP No. 3 may explain the reason of SARS-CoV-2 RNA founding in the outlet of Southern Tehran WWTP.

In other words, the removal efficiency percentage of parameters such as BOD$_5$, COD and SS in East Anzali WWTP
and Qom WWTP No. 3 were increased. So the question arises: is there a direct relationship between the efficiencies of virus removal and those parameters according to Fig. 4 and Table 1?

Generally, it is essential to investigate the physical and chemical properties of wastewater (e.g., pH, temperature, dissolved oxygen (DO) concentration, total nitrogen (TN) concentration, salinity), composition in waste activated sludge (WAS), and even population of society for providing precise data about transmission of SARS-CoV-2.

Conclusions

This work was concentrated on the presence of SARS-CoV-2 RNA in untreated and treated wastewater during the outbreak of coronavirus disease 2019 (COVID-19) in Southern Tehran WWTP, East Anzali WWTP and Qom WWTP No. 3. The results of this work showed that all samples taken from inlets of Southern Tehran WWTP, East Anzali WWTP and Qom WWTP No. 3 were positive (four positive samples for each of the WWTPs), while the SARS-CoV-2 RNA in outlet samples were two positive samples for Southern Tehran WWTP, negative for East Anzali WWTP and negative for Qom WWTP No. 3. The results of outlet samples of Southern Tehran WWTP described that the UV disinfection was more effective than chlorine disinfection and for effective disinfection; hence, WWTP operator should increase the concentration of free residual chlorine to amounts ≥0.5 mg/L, based on WHO on March 03, 2020. In addition, the high outlet BOD5, COD and SS values in Southern Tehran WWTP compared to East Anzali WWTP and Qom WWTP No. 3 may explain the reason of SARS-CoV-2 RNA founding in the outlet of Southern Tehran WWTP. As a result, the WWTP facilities could pose a threat on the health of workers during sludge production, transportation and disposal, residents living near to the treatment plants. According to the findings of this work, it should be emphasized that workers at WWTPs should wear standard personal protective equipment (PPE), like N95 mask, steel capped boots, face shields, hard hats, long pants safety glasses/goggles and gloves. In addition, we should not forget that proper sludge management of those WWTPs are essential due to the fact that the sludge from these treatment plants may contain active viruses. Finally, it is essential to investigate the physical and chemical properties of wastewater, composition in WAS, and even population of society for providing precise data about transmission of SARS-CoV-2 via water environmental and communities.

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Authors’ contributions

MA, SN and RN supervised of this study, and participated in its design and coordination and participated in writing - review & editing. JY, TMA, AN, and NZSJ participated in methodology, validation, and visualization of this study and detection of SARS-CoV-2 in untreated and treated wastewater using Real-Time PCR analysis. MH, BV, SKAV, and MB collected samples, performed the statistical analysis and participated in the design of the study. ANB participated in the design of the study, prepared to draft the manuscript (writing - original draft) and revised the manuscript (writing - review & editing). ANB, SY and SN participated in the design of the study and in the sequence alignment. All authors read and approved the final manuscript.

Declarations

Consent for publication Not applicable
Ethics approval and consent to participate Not applicable
Conflict of interest The authors declare that they have no conflict of interest.

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