CrAssphage for fecal source tracking in Chile: Covariation with norovirus, HF183, and bacterial indicators

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A B S T R A C T

Anthropogenic fecal pollution in urban waterbodies can promote the spread of waterborne disease. The objective of this study was to test CrAssphage, a novel viral human fecal marker not previously applied for fecal source tracking in Latin America, as a fecal pollution marker in an urban river in Chile. Human fecal markers CrAssphage CPQ_064 and Bacteroides HF183, the human pathogen norovirus GII, and culturable fecal indicator bacteria (FIB) were quantified at six locations spanning reaches of the Mapocho River from upstream to downstream of Santiago, as well as in repeated sub-daily frequency samples at two urban locations. Norovirus showed positive correlation trends with CrAssphage (τ = 0.57, p = 0.06) and HF183 (τ = 0.64, p = 0.03) in river water, but not with E. coli or enterococci. CrAssphage and HF183 concentrations were strongly linearly related (slope = 0.97, p < 0.001). Chlorinated wastewater effluent was an important source of norovirus GII genes to the Mapocho. Precipitation showed non-significant positive relationships with human and general fecal indicators. Concentrations of CrAssphage and HF183 in untreated sewage were 8.35 and 8.07 log10 copy/100 ml, respectively. Preliminary specificity testing did not detect CrAssphage or HF183 in bird or dog feces, which are predominant non-human fecal sources in the urban Mapocho watershed. This study is the first to test CrAssphage for microbial source tracking in Latin America, provides insight into fecal pollution dynamics in a highly engineered natural system, and indicates river reaches where exposure to human fecal pollution may pose a public health risk.

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1. Introduction

Rivers are the primary source of renewable freshwater for humans and ecosystems globally, yet they face many anthropogenic threats, including extraction and pollution (Vörössmarty et al., 2010). In particular, urbanization around rivers tends to degrade water quality, increasing potential for the spread of waterborne disease (Meybeck, 2003). Waterborne disease risks are most commonly assessed by measuring concentrations of fecal indicator bacteria (FIB), such as E. coli or enterococci, which are recommended for monitoring fecal pollution globally (World Health Organization, 2011). While monitoring FIB has many benefits, FIB are markers of general fecal pollution and do not indicate specific animal sources of feces, which differ in their human health risks (Soller et al., 2010b). As bacteria, FIB may also have limited utility for indicating the presence of viruses (Harwood et al., 2013), which are thought to be predominant etiologies of recreational waterborne illness (Soller et al., 2017, 2010a).

To improve detection of health-relevant microbial pollution, methods to assess human fecal pollution have been developed, primarily using genetic microbial source tracking (MST) markers. Among the highest performing MST markers are genetic targets from human-associated organisms, which are quantified by quantitative polymerase chain reaction (qPCR) (Warish Ahmed et al., 2016; Boehm et al., 2013; Mayer et al., 2018). Recently, MST methods were developed (Stachler et al., 2017) for a highly abundant human-gut-associated bacteriophage called CrAssphage (Dutilh et al., 2014). As a virus, CrAssphage could serve as a superior indicator of human viral pollution compared to bacterial indicators,
though little data on the co-occurrence of crAssphage and human viral pathogens currently exists (Farkas et al., 2018). CrAssphage MST markers have been tested in the USA (Ahmed et al., 2018a; Stachler et al., 2018), Nepal (Malla et al., 2019), Thailand (Kongprajug et al., 2019), Australia (Ahmed et al., 2018b), the UK (Farkas et al., 2019), Spain (García-Aljaro et al., 2017), and Japan (Malla et al., 2019), but not in any Latin American countries. Although crAssphage and high-performing human bacterial markers, such as Bacteroides HF183, Bacteroidales BacHum, and Firmicutes Lachno2, have been shown to be quite specific to human feces in most geographic contexts tested thus far, cross-reaction with non-human fecal sources has been reported (Ahmed et al., 2018a; Warish Ahmed et al., 2016; Boehm et al., 2013; Kildare et al., 2007; Malla et al., 2019; Mayer et al., 2018; Stachler et al., 2017). Thus, cross-reaction with locally relevant fecal sources should be examined prior to application in new regions (Stachler et al., 2017). CrAssphage MST marker abundance in sewage should also be examined to assess its local sensitivity for detecting sewage pollution because although crAssphage sequences in wastewater metagenomes have been found to be globally ubiquitous, there is also tremendous geographic sequence diversity (Edwards et al., 2019), and evidence of strong geographic dependence of crAssphage abundance in sewage (Stachler and Bibby, 2014).

While measuring MST markers is useful for assessing fecal contamination, directly measuring human pathogens informs the understanding of specific health risks posed by water contact or consumption. Epidemiological and microbial risk assessment modeling indicate that human norovirus is a predominant etiology of waterborne illness. Norovirus is a leading cause of gastroenteritis outbreaks globally, and primary outbreak cases are often linked to food or water exposure (Patel et al., 2009). Available data indicate that norovirus has been a leading cause of acute gastroenteritis in Chile since at least 2000. A study conducted in Santiago between 2000 and 2003 found that 45% of gastroenteritis outbreaks investigated were caused by human calcivirus, a family of viruses to which norovirus belongs, with the most common etiologic agent being norovirus GI (Vidal et al., 2005). A 2008 study among hospitalized children with diarrhea in Concepción, Chile found that norovirus prevalence was higher (25.5%) than that of other tested viruses, rotavirus and adenovirus (Montenegro et al., 2014). In 2010, an outbreak led to 31,036 reported cases of norovirus GI infection in the Antofagasta region. Primary cases in this outbreak were attributed to consumption of raw vegetables irrigated with treated wastewater containing low levels of residual free chlorine (Díaz et al., 2012). From 2013 to 2017, norovirus GI was the most prevalent enteric virus in outbreak, diarrhea monitoring, and shellfish samples collected by the Chilean epidemiological monitoring network (Instituto de Salud Pública de Chile, 2018). Quantitative microbial risk assessment (QMRA) models of US waters also consistently indicate that norovirus drives health risk in recreational waters polluted with sewage (Boehm et al., 2015; Wynn-Jones et al., 2011) and stormwater runoff (Soller et al., 2017). Thus, accurately assessing norovirus contamination is key to understanding risks to people who contact urban environmental waters.

The objective of this study was to test crAssphage for assessing fecal pollution in an urban river in Chile. This is the first study to apply human Bacteroides HF183 and crAssphage MST markers in Chile, and it adds new data on the co-occurrence of crAssphage and norovirus. Overall, this research extends knowledge of the utility of crAssphage in a new geographic context, provides insight into fecal pollution dynamics in a highly engineered natural system, and indicates river reaches where exposure to human fecal pollution may pose a public health risk.

2. Methods

2.1. Research questions

The specific research questions (RQ) investigated were: (RQ1) Do crAssphage and HF183 indicate human fecal pollution in the Mapocho River? Low concentrations of these markers in untreated wastewater or detection in non-human feces could suggest that these markers are poor indicators of human fecal contamination. (RQ2) Is crAssphage correlated with norovirus in Mapocho River water samples? To contextualize this correlation, correlations of norovirus with HF183 and culturable E. coli and enterococci were calculated as well. Norovirus is expected to be most correlated with crAssphage, since both are PCR-based measurements of viruses. (RQ3) Are crAssphage and HF183 correlated in Mapocho River water samples? Lack of correlation could indicate differential sources or environmental fate and transport, possibly caused by differences in viral and bacterial physiology. Finally, (RQ4) how do concentrations of fecal indicators vary with as a function of water turbidity, precipitation, and time of day? Correlation with these factors can provide insight into human and environmental processes governing fecal pollution in the Mapocho River.

2.2. Site description and sampling campaigns

Samples were collected from the Mapocho River in the greater metropolitan area of Santiago, Chile (33.440°W, 70.649°S), which has an estimated population of 7.1 million inhabitants (Instituto Nacional de Estadísticas de Chile, 2017). The Mapocho flows from its headwaters in the Andes Mountains east of Santiago, through the city, and then joins the Maipo River west of Santiago before discharging into the Pacific Ocean. Upstream of Santiago, the Mapocho’s catchments comprise 854 km² of mountainous, undeveloped land, classified as 23% grassland, 23% shrubland, 51% barren, and <1% each cropland and impervious (Figure S1) (Alvarez-Garretón et al., 2018). The Mapocho system within and to the west of Santiago, however, is highly engineered: its urban reaches are channelized, large fractions of its flow are extracted for irrigation, and it receives large volumes of wastewater treatment plant effluent downstream of the city. It also receives input from the Maipo River via the San Carlos canal in central Santiago (indicated in Fig. 1), although this canal was closed for maintenance from May 18 through June 2, 2019. Thus, the Mapocho did not receive input from the San Carlos canal during the spatial sampling campaign described below. Santiago has separate municipal stormwater and sewage collection systems, and stormwater is discharged to the Mapocho River as well.

Three sampling campaigns were undertaken to assess spatial and temporal water quality variation. The spatial campaign comprised 6 sampling sites along the length of the Mapocho River (Fig. 1), referred to from upstream to downstream as sites S1–S6 (detailed description in Table 1). Spatial sampling was conducted on 4 days during Fall 2019: May 22, 23, 25, and 26. S1 is located upstream of the city, in an unchannelized reach surrounded by low-density rural housing. S2–S4 are located in urban areas where the Mapocho is channelized. Much of the river’s flow is extracted for irrigation prior to reaching S2, and the San Carlos Canal replenishes the Mapocho’s flow by transferring water from the Maipo River to the Mapocho downstream of S2 but upstream of S3. S5 and S6 are located in agricultural areas on the western edge of Santiago, where the Mapocho is unchannelized. S5 receives chlorinated WWTP effluent from the Farfana WWTP, which performs activated sludge treatment and clarification before chlorination and discharge (Suez, 2020). S6 receives WWTP effluent from the Mapocho-Trelal WWTP, which also performs activated sludge treatment and
secondary clarification prior to chlorination and discharge (Suez, 2020). The Farfana and Mapocho-Trebal plants each have annual median treatment capacity of 8.8 m³/s of raw sewage, a population equivalent of 3.7 million people (Suez, 2020).

Temporal sampling was split between two campaigns: a sub-daily campaign and an hourly campaign. The sub-daily campaign comprised two sampling sites, T1 and T2 from upstream to downstream, both located just downstream of S3. These sites were sampled at 7:30 and 13:30 h daily for 15 days in 2019 between May 22 and June 5. The hourly campaign was conducted at site T2 on a single day; hourly samples were collected from 7:30 to 16:30 on June 6, 2019.

### 2.3. Sample collection and processing

500 mL samples were collected in 1 L polycarbonate bottles, which were cleaned with isopropyl alcohol before sampling. Sample bottles were triple rinsed with sample water before transferring a final sample to a sterile, disposable Whirl-Pak bag. Samples were collected mid-river at the surface. Samples processed for FIB were stored in the dark at 4 °C for <6 h until lab processing. Samples from the spatial campaign were stored at 4 °C for up to 9 h prior to processing for molecular analysis. River temperature was measured in situ using a handheld probe (YSI-30, YSI, Xylem, Rye Brook, NY). Turbidity was measured using a bench top turbidity meter (HF Scientific DRT-15CE, Fort Myers, FL). E. coli and enterococci were quantified by defined substrate fluorescence assay (Colilert-18 and Enterolert, respectively, IDEXX, Westbrook, ME), per the manufacturer’s instructions, in sample water diluted 10-fold with deionized water. The lower and upper limits of detection for the IDEXX assays were 10 and 24,192 most probable number (MPN) per 100 mL of water sample.

For molecular analysis, virus and bacteria from 100 mL of each sample were concentrated by filtering water through a 47 mm, 0.45 μm-pore size mixed cellulose ester membrane filter (MilliporeSigma, Burlington, MA) using sterile disposable filtration funnels (Nalge Nunc International, Rochester, NY). To aid virus adsorption, prior to filtration samples were augmented with MgCl₂ to a final concentration of 100 mM (Lukasik et al., 2000). Following filtration, 0.5 mL of RNA stabilization solution (RNAlater, Invitrogen, Waltham, MA) were added to the filter and incubated for 5 min before being aspirated through the filter. Process blanks tested the effectiveness of the sampling bottle and filter forceps cleaning procedure; a cleaned sampling bottle was used to collect deionized water, which was then processed along with river samples. Filters were stored in cryotubes overnight at 4 °C, and then transferred...
to –20 °C and stored for up to 1 month. Filters were then shipped to Stanford, CA, USA and stored at –80 °C for 4 months until nucleic acid extraction. This storage time is short compared to that of other studies which stored samples as long as 12–18 months (Li et al., 2019; Shanks et al., 2010). DNA and RNA were extracted simultaneously and directly from filters using the AllPrep PowerViral DNA/RNA kit (Qiagen, Hilden, Germany), which includes bead beating and β-mercaptoethanol denaturation of RNases. According to the manufacturer, this kit is designed for both bacterial and viral nucleic acid extraction. One extraction blank was included with every 15 samples to test for contamination in reagents or during the extraction process. Extracts were divided into 30 μl aliquots, stored at –80 °C, and then thawed only once for qPCR within 2 months. Microbial recovery was not quantified in this study. However, previous work indicated that extraction recovery efficiency measured by qPCR following direct extraction of filtered MgCl2-treated surface water was approximately 7% for non-enveloped virus (MS2 coliphage) and 17% for the bacterium E. faecium (Vlai et al., 2011). Recent work also extracting nucleic acids directly from MgCl2-treated samples found whole process RNA recovery, including concentration, of murine hepatitis virus (an enveloped virus) of approximately 66% (Ahmed et al., 2020).

An untreated sewage influent sample was collected from the Farfana WWTP. Sewage was allowed to settle, and 25 ml were filtered following the filtration protocol above. Wastewater nucleic acid extracts were quantified fluorometrically (Qubit 2.0, Life Technologies, Thermo Fisher Scientific, Waltham, MA) and diluted to 1 ng/μl of dsDNA. Fresh bird (n = 5) and dog (n = 5) feces were collected aseptically from concrete surfaces near the Mapocho River. Fecal samples were transported to the lab and stored at –20 °C within 1 h of collection. Nucleic acids were extracted from each fecal sample in an extraction run containing no water samples, diluted to 1 ng dsDNA/μl, and then composited in equal mass ratios to create one sample per feces type at 1 ng dsDNA/μl. Compositing or pooling samples for MST assay sensitivity and specificity is well-established in the literature (Boehm et al., 2016; Shanks et al., 2009, 2008).

2.4. qPCR assays

Primers, probes, and standard materials for all assays are given in Table S1. HF183/BacR287 (Green et al., 2014) and crAssphage CPQ_064 (Stachler et al., 2017) reactions were 25 μl, consisting of 1X Environmental Master Mix 2.0 (ABI, Thermo Fisher Scientific, Waltham, MA), 1 μM of each primer, 0.08 μM probe, 0.2 mg/ml BSA, 5 μl template, and the remainder water. Thermal cycling was performed in an ABI Step One Plus and comprised a 10-min hold at 95 °C, followed by 40 cycles of 95 °C for 15 seconds and 60 °C for 1 min. Standards were synthetic ssDNA ultramers (IDT, Coralville, IA). Norovirus GII ORF1-ORF2 (Loisy et al., 2005) reactions were 25 μl, consisting of 1X AgPath-ID 1-step RT-qPCR master mix (ABI, Thermo Fisher Scientific, Waltham, MA), 0.2 μM of each primer and probe, 5 μl template, and the remainder water. Thermal cycling conditions comprised 30-min at 50 °C for reverse transcription and a 10-min hold at 95 °C, followed by 45 cycles of 95 °C for 15 seconds and 60 °C for 1 min. Standard material was synthetic ssRNA spanning the ORF1-ORF2 junction (product no. VR-32355D, ATCC, Manassas, VA).

Six-point triplicate standard curves and no-template controls (NTC) were quantified for crAssphage, HF183, and norovirus GII on 8, 7 and 3 instrument runs, respectively, including on each plate of samples. Mixed effects regression, with instrument run as a random intercept, was used to combine data across multiple runs while retaining plate-specific standard equation estimates (Sivaganesan et al., 2010). The limit of detection (LOD) was defined as the lowest concentration that can be reliably detected, i.e., at which at least 95% of standard reactions are expected to amplify. The lower limit of quantification (LLOQ) was defined as the lowest standard that can be quantified with a coefficient of variation less than 0.35 (Klymus et al., 2020). See SM for details. All samples were run in duplicate reactions. To test for PCR inhibition, for each assay at least 20% of extracts were run at 1:1 and 1:10 dilutions. Inhibition was defined as ΔCt <2.3 between 1:1 and 1:10 dilutions.

2.5. Precipitation, flow, and historical FIB data

Cumulative six-hour precipitation measurements were obtained from the Meteorological Directorate of Chile online database, station ID 330019 (Chilean Directorate of Meteorology, n.d.), which is located near sampling station S3. To compute cumulative precipitation in the 24-hr period prior to each water sample, each 6-hr cumulative measurement was assumed to comprise 6 1-hr intervals of equal precipitation. The preceding 24 of these 1-hr intervals was then summed. The river's historical fecal coliform data were obtained courtesy of Aguas Andinas S.A. (internal monitoring program for river health). Enumeration of historical fecal coliform samples was performed by an external certified lab following Standard Method 9222 for membrane filtration (Rice et al., 2012).

2.6. Data analysis

This study's data presented several statistical properties common in environmental monitoring that were accounted for to accurately assess patterns: non-detects, serial correlation, and spatial clustering. If the data subset used to investigate a research question contained >15% non-detects, statistical methods that explicitly account for censoring were used. If statistically significant serial correlation in residuals was found (applying the astsa::acf2() function (Shumway and Stoffer, 2017) in R to model residuals), appropriate models were used. All analysis was performed on log10-transformed microbial concentrations. Non-detect values were assigned half the limit of detection in log space. Spatial clustering by sampling location was assessed with the Kruskall-Wallis rank sum test, and further analyses accounted for spatial clustering in several ways. Details of these methods are described below.

For RQ2, to examine the correlation between the four indicators with norovirus, concentrations of all five microbes in all available field samples were used. In these data, norovirus was not detected in >15% of samples, and both serial correlation (autocorrelation coefficient significant at p < 0.05) and spatial clustering (by sampling site, p < 0.05) were evident. Suitable rank-based regression or correlation routines that can account for clustered data are not available in R as of February 2020. Thus, concentrations were averaged for each of the eight sampling sites, yielding eight values for each microbe, and Kendall’s tau-b was computed. Kendall’s tau-b was used for computing rank correlations because it accounts for rank ties, which are common in censored data, while Spearman’s rho does not (Agresti, 2010). Tau is expected to be about 0.15 units smaller than rho for the same degree of correlation (Helsel and Hirsch, 2002).

For RQ3, to test the correlation between crAssphage and HF183, all available crAssphage and HF183 concentrations from field samples were used. This included the temporal and spatial sampling campaigns, but not the hourly campaign since only FIB were measured. In these data, <15% of values were non-detects and spatial clustering (by sampling site, p < 0.05) was evident. Thus, a generalized estimating equation (GEE) with clustering by sampling site, an exchangeable correlation error structure, and robust
standard errors was used. Serial correlation was not evident in model residuals.

For RQ4, to examine variation in indicator concentrations as a function of turbidity and precipitation, all concentrations of the four indicator organisms measured in the temporal campaign were used. In these data, <15% of values were non-detects, so ordinary least squares models were fit to indicator concentrations, with precipitation (cumulative over preceding 24 hours) and \( \log_{10} \) turbidity as independent variables. The temporal sampling campaign comprised two proximate sampling sites, and model residuals were highly correlated between these two sites. Thus, to avoid pseudo-replication, concentrations were averaged across the two sites at each time point. Model residuals did not show serial correlation, but crAssphage and HF183 model residuals showed a temporal trend, so time was included as a continuous covariate for these two models.

Multi-collinearity was assessed by computing variance inflation factors (VIF) on models with multiple covariates. Bonferroni correction was applied to family-wise error rates, where a family of tests was defined as a group of analogous tests used to investigate an individual research question. Bonferroni was selected as a conservative correction because tests within a family were generally not independent (e.g., correlation of A with B and A with C). The pre-adjusted alpha value was 0.05. Data analysis was performed in R version 3.5.1 using Tidyverse packages (Wickham et al., 2019). Analyses can be reproduced from code available at https://github.com/williejennings/chile_mapocho_2019.

3. Results

3.1. qPCR and MST performance

Assay performance characteristics are given in Table S2. No processing blanks, extraction blanks, or NTCs amplified for any targets. Assays efficiencies were 95%, 92%, and 85% for HF183, crAssphage, and norovirus GI respectively. No inhibition was observed for crAssphage or HF183. For norovirus, three samples amplified in 1:10 dilutions, and two of those samples showed inhibition. Given that norovirus concentration estimates were generally low, concentration estimates from undiluted samples were used for analysis, recognizing that these estimates may be biased low due to RT-qPCR reaction inhibition.

Neither crAssphage nor HF183 amplified in composite dog or bird fecal samples. CrAssphage and HF183 concentrations in raw sewage were estimated at 8.35 (\( \sigma \) of qPCR replicates = 0.002) and 8.07 (0.02) \( \log_{10} \) cp/100 ml, respectively.

3.2. Turbidity, temperature, and precipitation

During the temporal campaign, turbidity (Table S3) was lowest at upstream sites S1 (mean = 0.91 \( \log_{10} \) NTU, standard deviation = 0.11 \( \log_{10} \) NTU) and S2 (1.44, 0.23 \( \log_{10} \) NTU). Turbidity was highest at mid-city sites S3 (1.92, 0.81 \( \log_{10} \) NTU) and S4 (2.35, 0.74 \( \log_{10} \) NTU). Water temperature (Table S4) showed an increasing trend from upstream sites to downstream sites, with lowest temperature at S1 (mean = 6.5°C, standard deviation = 2.3°C) and highest temperature at S5 (16.4°C, 1.3°C). No precipitation occurred during or in the week prior to the spatial sampling campaign. During the temporal sampling campaign, turbidity was very similar at sites S1 (1.9, 0.5 \( \log_{10} \) NTU) and S2 (1.5, 0.4 \( \log_{10} \) NTU), and temperature was not measured. Mean precipitation in the 24 hours preceding samples was 1.2 mm and standard deviation was 2.6 mm (Table S4 shows data disaggregated by time of day). Linear regression of turbidity on time of day and precipitation indicated that turbidity was significantly associated with time of day (p < 0.001) but not 24-hr cumulative precipitation (p = 0.06). Controlling for precipitation, expected turbidity was 0.61 \( \log_{10} \) NTU higher at 13:30 than 7:30.

3.3. Variation of molecular indicators, FIB, and norovirus

Seventy-eight field samples were tested for molecular and culturable targets. An additional 14 samples were tested only for culturable targets. Across all samples, the median (25th, 75th percentile) E. coli and enterococci concentrations were 2.70 (2.43, 3.04) and 2.29 (1.85, 2.80) \( \log_{10} \) MPN/100 ml, respectively. Median crAssphage, HF183, and norovirus concentrations were 3.85 (3.38, 4.46), 4.21 (3.80, 4.79), and <1.78 (<1.78, <1.78) \( \log_{10} \) cp/100 ml, respectively. Norovirus was detected in 14 of 78 samples. During the spatial sampling campaign, median concentrations of crAssphage and HF183 were highest (6.69 and 6.68 \( \log_{10} \) cp/100 ml, respectively) at downstream sites S5 and S6 and lowest at mid-city sites S3 and S4 (<3.52 and 1.91 \( \log_{10} \) cp/100 ml, respectively) (Fig. 2). Norovirus was detected in three of eight upstream (S1 and S2) samples, one of eight mid-city samples, and all eight downstream samples, where medians were approximately 2.6 \( \log_{10} \) cp/100 ml. E. coli and enterococci did not significantly vary by site (p > 0.05). Median E. coli concentrations ranged from <1.00 (S5) to 3.34 (S2) \( \log_{10} \) MPN/100 ml. Median enterococci concentrations ranged from <1.00 (S1, S5) to 2.61 (S6) \( \log_{10} \) MPN/100 ml.

Median \( \log_{10} \) unit/100 ml concentrations during the temporal campaign across sites T1 and T2 were 3.82 for crAssphage, 4.13 for HF183, 2.69 for E. coli, and 2.49 for enterococci (Fig. 4). Norovirus was not detected at sites T1 or T2 during the temporal campaign. During the temporal campaign, median measurements at sites T1 and T2 differed by 7% or less for each indicator.

CrAssphage and HF183 concentrations in river water were strongly correlated. A 1 \( \log_{10} \) unit increase in crAssphage concentration (per 100 ml) was associated with a 0.97 (95% CI 0.89, 1.05; p < 0.001) \( \log_{10} \) unit increase in HF183 concentration. This estimate describes covariation of HF183 and crAssphage both within an individual sampling site and between sampling sites, while accounting for spatial clustering and temporal autocorrelation.

Norovirus was consistently detected only at wastewater effluent-impacted sites, where crAssphage and HF183 were highest. However, no indicator was significantly correlated with norovirus after adjusting for multiple comparisons. Scatter plots of norovirus against human fecal and general fecal indicator concentrations are shown in Fig. 3. Correlation coefficients between norovirus and indicator concentrations across spatial and temporal sampling campaigns were as follows: norovirus with crAssphage, 0.57 (Kendall’s tau-b, p = 0.06); with HF183, 0.64 (p = 0.03); with E. coli, –0.11 (p = 0.70); and with enterococci, –0.19 (p = 0.52). The alpha value used was 0.0125 (0.05 adjusted for four comparisons). Site-averaged values at the six spatial and two temporal sites were used for these estimates, as described in Methods.

Sub-daily sampling at two proximate sites revealed that only enterococci was associated with turbidity, and no indicators were associated with precipitation. A 1-\( \log_{10} \) NTU increase in turbidity was associated (p < 0.001) with a 0.87 (95% CI 0.42, 1.32) \( \log_{10} \) increase in enterococci concentration (Table 2), estimated by linear regression. No other coefficients for either precipitation or turbidity were statistically significant in models of crAssphage, HF183, E. coli, or enterococci (p-values > 0.0125, the alpha value adjusted for four comparisons). All variance inflation factors were near 1, indicating that multi-collinearity was not a problem.

Hourly samples collected between 7:30 and 16:30 on a single day and measured for E. coli and enterococci revealed no significant linear trend over time. A plot of these data (Figure S4) did not suggest other diurnal patterns.
Fig. 2. Microbial concentrations from the spatial sampling campaign. Crassphage, HF183, and norovirus are in units of log_{10} cp/100 ml. E. coli and enterococci are in units of log_{10} MPN/100 ml. Filled circles represent measured concentrations. Red Xs represent non-detects. Boxplot centerlines are medians, bottoms and tops of boxes are 25th and 75th percentiles, and whiskers extend to the furthest data point within a distance of 1.5 x interquartile range from the 25th and 75th percentiles, respectively. Individual points are displaced horizontally to aid visualization. N = 4 for each sampling site (S1 through S6). (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

Fig. 3. Human fecal marker and general fecal indicator concentrations compared to norovirus concentrations. Crassphage, HF183, and norovirus are in units of log_{10} cp/100 ml. E. coli and enterococci are in units of log_{10} MPN/100 ml. Linear regression lines with 95% confidence bands about the mean are shown for illustrative purposes. These trend lines do not represent correlation estimates reported in Results because they are linear rather than rank relationships (so poorly handle the large number of non-detects) and do not account for statistical clustering by sampling location.
4. Discussion

4.1. CrAssphage is useful for indicating wastewater impact in Chile’s Mapocho River

Concentrations of crAssphage in untreated sewage using assay CPQ_064 were high, indicating that crAssphage has the potential to be a sensitive indicator of wastewater impact in Santiago. Concentrations measured in this study were comparable to or greater than concentrations measured in other countries. For example, crAssphage (CPQ_056) was estimated in untreated sewage in Florida, USA at $8.08 - 8.98 \log_{10} \text{cp/100 ml}$ (Ahmed et al., 2018a); in Australia (CPQ_064) at $7.91 \log_{10} \text{cp/100 ml}$ (Ahmed et al., 2018b); in the UK (CPQ_056) at $4 - 8 \log_{10} \text{cp/100 ml}$ (Farkas et al., 2019); and in Thailand (CPQ_056) at $6.57 \log_{10} \text{cp/100 ml}$ (Kongprajug et al., 2019). Although this study did not measure crAssphage concentrations in WWTP effluent samples, studies have reported 0- to 2-log10 (Thailand (Kongprajug et al., 2019) and UK (Farkas et al., 2019)) and 3-log10 (USA (Wu et al., 2020)) reductions of crAssphage concentrations through WWTPs with activated sludge units. More generally, this study’s measurements of crAssphage support the finding that crAssphage is a sensitive human-associated viral MST marker, with concentrations in sewage at least 2–3 orders of magnitude greater than concentrations typically measured of other human-associated viral tracking markers, including human adenovirus, human polyomavirus, and pepper mild mottle virus (W. Ahmed et al., 2016; Farkas et al., 2019; Hughes et al., 2017; Stachler et al., 2018).

CrAssphage was specific to human fecal waste among animal fecal samples tested. It did not amplify in bird or dog feces, two important non-human fecal sources in the Mapocho watershed. However, this study cannot rule out cross-reaction with other fecal sources, which could be important in agricultural areas. Most studies, with the exception of work done in Nepal (Malla et al., 2019), have reported high specificity of crAssphage to human feces, with limited cross-reaction in cat feces and cattle wastewater in Australia (Ahmed et al., 2018b), poultry litter in Florida, USA (Ahmed et al., 2018a), and swine feces in Thailand (Kongprajug et al., 2019).

CrAssphage was highly correlated with the human-associated bacterial fecal marker HF183 in field samples in this study. This suggests that both markers derive from similar sources and have similar environmental fate on the time scale required for water to pass through the urban portion of the Mapocho River. No data are available that describe the degree to which covariance between CPQ_064 and HF183 gene targets may be driven by crAssphage genes being located inside Bacteroides cells or genomes. Other studies have reported mixed results regarding correlations between crAssphage and HF183 in environmental waters. Samples

Table 2

| Parameter   | Units       | crAssphage Estimate (std err) | p-value | HF183 Estimate (std err) | p-value | E. coli Estimate (std err) | p-value | Enterococci Estimate (std err) | p-value |
|-------------|-------------|------------------------------|---------|--------------------------|---------|----------------------------|---------|-------------------------------|---------|
| Intercept   | log$_{10}$ concentration | 4.09 (0.61) | <0.001 | 4.20 (0.59) | <0.001 | 2.07 (0.33) | 0.77 (0.45) | 0.10 (0.21) | 0.01 |
| Precipitation | mm        | 0.09 (0.05) | 0.09 | 0.12 (0.05) | 0.03 | 0.05 (0.03) | 0.08 (0.05) | 0.21 (0.04) | 0.01 |
| Turbidity   | log$_{10}$ NTU | 0.00 (0.32) | 0.09 | 0.23 (0.32) | 0.47 | 0.35 (0.18) | 0.93 (0.24) | <0.001 | 0.29 |
| Time        | days       | -0.07 (0.03) | 0.04 | -0.10 (0.03) | <0.01 | - | - | - | |
| n           |            | 27 | 27 | 29 | 29 | 29 | 29 | 29 | 29 |
| Adj. R-squared |           | 0.13 | 0.30 | 0.23 | 0.41 | 0.41 | 0.41 | 0.41 | 0.41 |
collected in wastewater-impacted urban river in the USA showed strong correlation between crAssphage CPQ_064 and HF183, and moderate correlation between crAssphage CPQ_056 and HF183 (Stachler et al., 2018). A study in Spain that used a different crAssphage assay found statistically significant linear correlations with HF183 in a creek that does not receive WWTP effluent, but not in a river that does receive WWTP effluent (Ballesta et al., 2019). Indeed, another study in the USA using CPQ_056 found that crAssphage was not correlated with HF183 in storm drain outfalls or a wetland contaminated with sewage in Tampa, Florida (Ahmed et al., 2018a). One possible explanation for the observed differences between HF183 and crAssphage is differential environmental persistence. Both crAssphage decay studies published to date have found evidence that crAssphage markers persist about 2—3 times longer than HF183 markers in freshwater mesocosms (Ahmed et al., 2019; Ballesta et al., 2019).

4.2. Norovirus patterns were more closely tracked by molecular MST markers than FIB

Correlation coefficients of norovirus with indicators were suggestive of positive relationships with crAssphage and HF183, and null relationships with E. coli and enterococci. It is important to note that these correlation analyses used site-averaged concentrations, and so they had low power. The only other study to date that measured crAssphage and norovirus simultaneously found statistically significant positive rank correlations between crAssphage and norovirus GII in wastewater influent and effluent from the UK (Farkas et al., 2019). No other study has examined co-occurrence of norovirus and crAssphage in an environmental water. The lack of correlation between FIB and human viruses in environmental waters is commonly reported (Korajkic et al., 2018) and may be attributable to differential environmental fate and transport (USEPA, 2015) or removal in wastewater treatment plant processes (Rose et al., 1996; Zhang and Farahbakhsh, 2007), as well as differential sources.

4.3. Although FIB pollution in the Mapocho River has declined dramatically since at least 2004, human fecal pollution and norovirus were elevated at some locations

In 2005, Santiago was treating only 4% of its sewage and discharging the rest untreated to the Mapocho River. With the construction of two large conventional treatment plants (activated sludge with chlorine disinfection), by 2016 Santiago was treating 100% of its wastewater (La Farfana WWTP site visit and UNFCCC website (UNFCCC, 2020, n.d.)). This increase in sewage treatment has resulted in a dramatic decline in fecal coliform concentrations in the Mapocho at locations coincident with sites sampled in this study (Figure S3). Compared to historical data, E. coli concentrations measured in this study generally corroborate the lowered levels of fecal coliform concentrations reported in recent years, considering methodological differences (Figure S3).

However, given the consistent detection of norovirus GII genes in the WWTP effluent-impacted waters at sites S5 and S6, these waters may present a health risk to those who contact it directly or who consume products recently irrigated with it. For example, the 2010 norovirus GII outbreak in the Antofagasta region of Chile originated from consumption of raw vegetables irradiated with treated wastewater containing low levels of residual free chlorine (Diaz T. et al., 2012). The national government and citizen groups, who have expressed growing interest in reclaiming the Mapocho for recreational use, should also consider the risk of waterborne illness potentially posed by viral pathogens in river extents receiving large loads of chlorinated effluent—a risk that may not be indicated by FIB. It is important to note that genome-based detection of norovirus does not indicate the infectivity of these viruses, and thus potential health risks cannot be stated unequivocally. However, a recent meta-analysis found that culturable male-specific coliphage often survive secondary treatment and chlorination, and that removal of qPCR-measured norovirus GII (2.7 log_{10} reduction) and culturable surrogate male-specific coliphage (2.9 log_{10} reduction) was similar (Pouillot et al., 2015). More generally, conventional biological treatment with chlorination is thought to be less effective at removing viruses than bacteria (Rose et al., 1996; Zhang and Farahbakhsh, 2007). These studies, along with the low infectious dose (ID) of human norovirus (ID_{50} estimated between 10 and 10^{3} genome copies (Atmar et al., 2014; Teunis et al., 2008)), support the conjecture that the qPCR-measured norovirus concentrations at sites S5 and S6 may pose a human health risk.

Human bacterial and viral fecal pollution concentrations were also high at upstream sites S1 and S2. Since these sites do not receive treated wastewater effluent, this pollution likely comes from untreated human feces, delivered by direct discharge of household-scale wastewater, leaking sewage infrastructure, or dry weather flows from irrigation or other anthropogenic washing activities. No precipitation occurred during or immediately prior to the spatial sampling campaign, which included sites S1 and S2. At S1 and S2, norovirus was detected in three of eight samples, and median crAssphage and HF183 concentrations exceeded QMRA-based risk thresholds for recreational exposure to sewage-polluted waters of 30 illnesses per 1000 bathers (3.67 log_{10} cp/100 ml for crAssphage (Crank et al., 2019) and 3.62 log_{10} cp/100 ml for HF183 (Boehm et al., 2015)).

Median human fecal marker concentrations declined substantially from S2 to S3. This decline may be attributable to dilution from tributary inflows between S2 and S3 (Fig. 1). If the decline were based on microbial decay alone, crAssphage would be expected to decay less than HF183 and culturable FIB given the cold water temperatures at sites S2 and S3 (Ballesta et al., 2019). This expectation is contradicted by observations that crAssphage declined by two to three orders of magnitude between S2 and S3, while HF183 declined by a similar amount and E. coli and enterococci declined by much less (zero to one orders of magnitude). For dilution to be the primary process governing these concentrations reduction, tributary inflows would need to contain similar concentrations of FIB and considerably lower concentrations of human markers compared to the Mapocho at S2. However, further work would be required to confirm that dilution was more important than microbial decay, and also to investigate the role of sediment resuspension, in controlling concentrations between S2 and S3. Furthermore, while the San Carlos canal (Fig. 1), which delivers water from the Maipo River into the Mapocho River just upstream of S3, would typically influence microbial concentrations at S3, it did not contribute any flow during the spatial sampling campaign because it was closed for maintenance activities.

4.4. River channel maintenance activities may elevate FIB concentrations

The Mapocho River receives stormwater from Santiago’s stormwater system, which is separate from its sewage system, and from direct overland drainage. Human and general fecal pollution in stormwater flows have been extensively documented (Brownell et al., 2007; Parker et al., 2010; Sidhu et al., 2013; Tiefenthaler et al., 2011). Thus, concentrations of both human and general indicators of fecal pollution were expected to increase following precipitation events. While HF183, E. coli, and enterococci concentrations showed upward trends with increasing precipitation (p < 0.10), none of these associations were statistically significant after adjusting for multiple
comparisons. Likely, the effect of precipitation cannot be distinguished in this data set ($n = 27$ for molecular markers, $29$ for FIB) because unmeasured anthropogenic and environmental processes cause substantial variation in fecal pollution concentrations.

Turbidin, however, was positively associated with enterococci concentrations, and showed a positive trend with E. coli concentrations. Interestingly, in multiple linear regression, turbidity was strongly associated with time of day ($7:30$ vs $13:30$), but not precipitation (data not shown). In the absence of precipitation driving turbidity levels via stormwater flows or sediment resuspension from elevated river flows, this diurnal turbidity pattern may be related to river channel bed maintenance activities that occurred throughout the sampling campaign. These activities included the use of heavy machinery alongside and occasionally in the river, and were conducted during business hours, implying the potential for greater sediment disturbance at $13:30$ than $7:30$. The association between turbidity and FIB concentrations is supported by laboratory and modeling studies showing that riverine sediment can harbor elevated levels of FIB, and that FIB concentrations in the water column may increase when sediment is resuspended (Droppo et al., 2011, 2008). Time of day was not included in linear regression models of microbial concentrations because time of day was strongly correlated with turbidity.

4.5. Limitations

This field study yielded insight into applying crAssphage and HF183 in a new geographic context, as well as fecal pollution patterns in the Mapocho River. It also has several limitations. First, more extensive characterization of crAssphage and HF183 concentrations in sewage would improve confidence in this study’s assessment of these markers as sensitive indicators of sewage pollution. Repeat sampling of influent sewage and effluent at both the Farfana and Mapocho-Trelbal plants would aid interpretation of this study’s data downstream of these plants. Previous studies have reported no variation in crAssphage concentrations in sewage over the course of a year (García-Aljaro et al., 2017; Malla et al., 2019), so year-round characterization is likely unnecessary. Second, more extensive specificity evaluation of crAssphage and HF183 would be desirable. This would include assessment of these markers’ concentrations in other fecal sources potentially important in the Mapocho watershed. The approach we used of compositing the DNA from the fecal samples could potentially lead to false negatives if the target is diluted to below the assay detection limit when it is combined with the other DNA extracts; however this approach is established in the MST field (Boehm et al., 2016; Shanks et al., 2009; 2008). Third, PCR inhibition of the norovirus GI assay was observed, even though both the extraction protocol and the master mix were selected for their capacity to remove PCR inhibitors. Inhibition potentially yielded underestimates of norovirus concentrations. Dilution of extracts to reduce inhibitory effects was not feasible, given the low concentrations of norovirus present. Filtering larger sample volumes could potentially increase method sensitivity, but could also increase inhibition of PCR reactions. Similar to this study, other studies of urban environmental waters have found little or no inhibition of qPCR assays, yet substantial inhibition of RT-qPCR assays in the same samples (Steele et al., 2018).

5. Conclusions

- CrAssphage and HF183 are useful indicators of human fecal pollution in Santiago, Chile.
- Fecal indicator bacteria are inadequate indicators of viral pollution from chlorinated wastewater effluent.
- Anthropogenic impacts on Mapocho River microbial water quality are extensive.
- Viral pollution must be considered in efforts to reclaim the Mapocho River for recreational activities.

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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Appendix A. Supplementary data

Supplementary data to this article can be found at https://doi.org/10.1016/j.wroa.2020.100071.

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