Evaluation of Two Triplex One-Step qRT-PCR Assays for the Quantification of Human Enteric Viruses in Environmental Samples
Farkas, Kata; Peters, Dafydd E.; McDonald, James; de Rougemont, Alexis; Malham, Shelagh; Jones, David
Food and Environmental Virology

DOI: 10.1007/s12560-017-9293-5
Published: 01/09/2017

Publisher's PDF, also known as Version of record

Citation for published version (APA):
Farkas, K., Peters, D. E., McDonald, J., de Rougemont, A., Malham, S., & Jones, D. (2017). Evaluation of Two Triplex One-Step qRT-PCR Assays for the Quantification of Human Enteric Viruses in Environmental Samples. Food and Environmental Virology, 9(3), 342-349. https://doi.org/10.1007/s12560-017-9293-5

General rights
Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

- Users may download and print one copy of any publication from the public portal for the purpose of private study or research.
- You may not further distribute the material or use it for any profit-making activity or commercial gain
- You may freely distribute the URL identifying the publication in the public portal

Take down policy
If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.
Evaluation of Two Triplex One-Step qRT-PCR Assays for the Quantification of Human Enteric Viruses in Environmental Samples

Kata Farkas · Dafydd E. Peters · James E. McDonald · Alexis de Rougemont · Shelagh K. Malham · Davey L. Jones

Received: 13 January 2017 / Accepted: 28 March 2017 © The Author(s) 2017. This article is an open access publication

Abstract Human enteric viruses are responsible for waterborne and shellfish-associated disease outbreaks worldwide. Quantitative reverse transcription PCR (qRT-PCR) is often used to assess the health risks associated with shellfish and environmental water, but viral titres in sediments are less commonly investigated. In this study, we developed and validated two multiplex qRT-PCR assays for aquatic sediment and shellfish samples targeting viruses that are a common cause of gastroenteritis (norovirus GI, GII and hepatitis A virus), two emerging viruses (sapovirus and hepatitis E virus), along with mengovirus (MgV), which is often used as a sample process control for the assessment of RNA extraction efficiency. Singleplex and multiplex assays demonstrated comparable PCR efficiencies and gave reliable results over a wide concentration range. The multiplex assays showed remarkable sensitivity with a limit of detection of 1 RNA copy/μL nucleic acid extract for all target viruses and limits of quantification of 3–18 RNA copies/μL for the targeted human pathogenic viruses and 20–40 RNA copies/μL for MgV. The results demonstrated the veracity of multiplex qRT-PCR for the estimation of viral titres in sediment and shellfish, allowing the rapid assessment of viral infection risks associated with environments exposed to wastewater contamination.

Keywords Multiplex real-time reverse transcription PCR · Nucleic acid quantification · Enteric viruses · Sediment · Shellfish

Introduction

Enteric viruses represent a major risk to human health, being responsible for numerous local and global disease outbreaks. For example, noroviruses (NoV) are responsible for approx. 3 million acute gastroenteritis cases in the UK and 20 million cases in the USA each year (Centers for Disease Control and Prevention 2015; Tam et al. 2012) placing an enormous burden on health care systems and the wider economy. Hepatitis A and E viruses (HAV and HEV), causing acute liver disease, have also been shown to be associated with large disease outbreaks. HAV used to be the most frequent cause of hepatitis worldwide; however, with improved sanitation and vaccination it has become less frequent in developed countries (Tahaei et al. 2012). Nonetheless, HAV, along with NoV, still represents the major cause of foodborne viral outbreaks (Bosch et al. 2016), including those associated with shellfish (Bellou et al. 2013). HEV was responsible for the largest viral waterborne outbreak in New Delhi in December 1955–January 1956, where more than 29,000 cases were
registered with HEV infection (Purcell 1996; Rao and Melnick 1986). Recently, HEV and sapoviruses (SaV), which cause gastroenteritis similar to NoV infection, have been responsible for sporadic cases in developed countries and are now considered as emerging pathogens (Dalton et al. 2013; Yates 2014).

Due to the high risks associated with the consumption of shellfish contaminated with enteric viruses, a standard method is now available for the quantification of NoV GI, GII and HAV (International Organization for Standardization 2013; Lees and CEN WG6 TAG4 2010). This method recommends the use of a one-step quantitative reverse transcription PCR (qRT-PCR) for the enumeration of viral RNA in the extract of shellfish digestive gland. However, while this approach does not give information on the infectivity status of the virus particles, it is the most sensitive and accurate method available for those viruses whose propagation in vitro is challenging, e.g. human NoV and SaV. Recent efforts have also been made to use molecular methods for enteric virus detection in sediment (Miura et al. 2011; Staggemeier et al. 2015a, b). Nonetheless, these methods often lack thorough validation, and the recoveries may vary due to the contrasting chemical composition of the samples. Therefore, the use of extraction and sample process controls (such as mengovirus) is recommended (Hennechart-Collette et al. 2015; International Organization for Standardization 2013; Martin-Latil et al. 2012); however, they are not routinely used.

The lack of proper validation and use of controls may be associated with the high costs and extended labour of the individual quantification of RNA viruses using qRT-PCR. Methods are available for the quantification of enteric viruses from shellfish, water and faecal samples using duplex and multiplex assays (Farkas et al. 2015; Fuenteres et al. 2014; Lee et al. 2016; Martin-Latil et al. 2012; van Maraseveen et al. 2010; Yan et al. 2013); however, no method has been validated for sediment. In this study, we validated two triplex qRT-PCR assays for the simultaneous quantification of the RNA of the most prevalent NoV genogroups (GI and GII), HAV, two emerging pathogens, SaV GI and HEV, and a commonly used extraction control, mengovirus (MgV), in sediment and shellfish nucleic acid extracts. One assay targeted the NoV GI, SaV and HEV and the other assay targeted the NoV GII, HAV and MgV.

Materials and Methods

Target Viral RNA

NoV GI RNA, MgV strain VMC0 and HAV strain pHM17543c were kindly provided by Dr. Lisa Cross (Centre for Environment Fisheries and Aquaculture Science; CEFAS UK). NoV GI RNA was derived from a pool of nucleic acid extracts of shellfish samples processed according to the ISO/TS 152016-1 (2013) standard. The sample was tested for all viruses targeted in this study using the singleplex qRT-PCR assays detailed below and contained no other target viruses. SaV GL2 was obtained from clinical stool samples and genotyped by the National Reference Centre for Enteric Viruses, Dijon, France. The viral sample was generated by the preparation of a 10% solution, using phosphate-buffered saline (pH 7.4), which was subsequently filtered through a 0.2-μm filter. Norovirus sample (diluted and filtered faecal matter from a patient with confirmed NoV GII infection) and HEV RNA (ORF3 segment) were provided by Prof. Ian Goodfellow (University of Cambridge, UK). When necessary, viral RNA was extracted from a 0.5 mL viral sample using the NucliSENS® MiniMag® Nucleic Acid Purification System (bioMérieux SA, France) and eluted in 100 μL molecular grade water. Viruses and viral RNA were stored at −80 °C. Nucleic acid extracts derived from faecal matter were tested for all viruses targeted in this study using the singleplex qRT-PCR assays detailed below. Results confirmed that the samples contained no target virus other than NoV GII or SaV GI.

Spiking Environmental Samples

Sediment and mussel samples were collected in the Conwy estuary (53°17’37.5”N 3°50’22.0”W), North Wales, at low tide. A sediment sample was processed using the elution–concentration method described elsewhere (Farkas et al. 2017; Lewis and Metcalf 1988). In brief, five aliquots of 10 g sediment sample were mixed with 30 mL 3% beef extract in 2 M NaNO3 (pH 5.5) for 30 min and the solid matter was removed by centrifugation at 2500 × g for 10 min. The pH of the eluent was adjusted to 7.5, then incubated in 15% polyethylene glycol 6000 and 2% NaCl overnight at 4 °C and centrifuged at 10,000 × g for 30 min at 4 °C. The digestive tissue (DT) of 30 mussels was extracted and homogenised. Aliquots of 2 g of the digestive tissue mix were treated with proteinase K according to the ISO/TS 152016-1 (2013) standard. Viral nucleic acids of the sediment and shellfish DT concentrates were extracted using the NucliSENS® MiniMag® Nucleic Acid Purification System (bioMérieux SA, France). Preliminary findings confirmed that the samples were negative for all target viruses prior to spiking. A pool of viral nucleic acids (NoV GI and GII, SaV, HAV, HEV, MgV) were added to the nucleic acid extracts at the ratio of 1:50 to reach a final concentration of approx. 10⁶ RNA copies/μL. That sample was further diluted in sediment or shellfish extract to achieve the final concentrations of 10³, 10⁴, 10⁵, 10⁶, 60, 40, 20, 10, 5 and 1 RNA copies/μL.
qRT-PCR Assay

All qRT-PCR assays were carried out in a QuantStudio™ Flex 6 Real-Time PCR System (Applied Biosystems, USA). Standards for NoV GI, GII and HAV (prepared according to ISO/TS 15216-1:2013 standard) were kindly provided by Dr. Lisa Cross (CEFAS, UK). SaV and MgV standards were derived from cloning qRT-PCR amplicons into pGem-T Easy vector (Promega, USA). A pSV plasmid incorporating the HEV ORF3 gene was kindly provided by Prof. Ian Goodfellow (University of Cambridge, UK). Plasmids were transformed to Alpha Select Bronze Competent Cells (Bioline, UK) and isolated using the ISOLATE II Plasmid Mini Kit (Bioline, UK). Recombinant plasmids were quantified using NanoDrop ND-1000 (NanoDrop, USA) and 10-fold serially diluted. Standard dilutions ranging from 10^5 to 10^0 DNA copies/μL in triplicate were used to generate standard curves for qRT-PCR quantification. Amplification efficiency, slope and R^2 were determined based on the standard curve of each reaction and calculated by the QuantStudio™ Real-time PCR software (Applied Biosystems, USA).

Primers and probes were adapted from previous studies; however, the reporters and/or quenchers were replaced for multiplex applications.

All singleplex and multiplex qRT-PCR assays were based on a single-step TaqMan-based assay described in the ISO/TS 15216-1:2013 standard (International Organization for Standardization 2013) using the RNA UltraSense One-step qRT-PCR kit (Invitrogen, USA). The 20 μL qRT-PCR reaction mix contained 1xRNA UltraSense Reaction Mix with 1 μL RNA UltraSense Enzyme Mix, 10 pmol of the forward and the reverse primers, 5 pmol of the probe/probes, 0.1 μL ROX reference dye, 1 μg bovine serum albumin (BSA) and 3 μL of the sample/standard. Negative controls (3 μL molecular grade water) were included in each run. Due to the differences in the melting temperatures of the primers and probes, two assays were validated: one with the annealing temperature (Ta) of 56 °C for the detection of NoV GI, SaV and HEV and another with Ta of 60 °C for NoV GII, HAV and MgV. The qRT-PCR assay consisted of a 60-min reverse transcription step at 55 °C followed by a 5-min step of denaturation at 95 °C, and 45 cycles of amplification consisting of 95 °C for 15 s, 56 °C for 60 s and 65 °C for 60 s. The baseline (cycle threshold; Ct) was manually adjusted after each run.

### Table 1 Primers and probes used for the singleplex/multiplex qRT-PCR assays

| Virus           | Assay | Primers/probes          | Sequence (5'–3') | Amplicon length | References                  |
|-----------------|-------|--------------------------|------------------|-----------------|-----------------------------|
| Norovirus GI    | Assay 1| QNIF4-F                  | CGCTGGATGCGCNNTTCCAT  | 86 bases         | Da Silva et al. (2007)      |
|                 |       | NV1LC-R                  | CCTTAGACGCCATCATCATTTAC  |                 | Svraka et al. (2007)        |
|                 |       | TM9-P                    | FAM-TGGACAGGAGATCGCNFQMGB  |                 | Hoehne and Schreier (2006)  |
| Sapovirus GI    | Assay 1| CU-SV-F                  | TTGGCCCTCAGCCACCTAC  | 101 bases        | Chan et al. (2006)          |
|                 |       | CU-SV-R                  | CCCCCATYTCAAACTACTWTTTG  |                 | Chan et al. (2006)          |
|                 |       | CU-SV-P                  | CAAATTAGTGTGATGAGGGG  |                 | Chan et al. (2006)          |
| Sapovirus GI    | Assay 1| CU-SV-P                  | VIC-TGGTTCATAGTGTTAC-NFQMGB*  |                 | Chan et al. (2006)          |
| Hepatitis E virus| Assay 1| JVHEV-F                  | GGTTGGTTCTGGGTGAC  | 71 bases         | Jothikumar et al. (2006)    |
|                 |       | JVHEV-R                  | AGGGGTTCAGTGATGAA  |                 | Jothikumar et al. (2006)    |
|                 |       | JVHEV-P                  | ABY-TGATTTCAGCCTTCG-GQSY*  |                 | Jothikumar et al. (2006)    |
| Norovirus GII   | Assay 2| QNI2-F                   | ATGTTCAAGTGGTATGAGGCTTCWGA  | 89 bases         | Loisy et al. (2005)         |
|                 |       | COG2-R                   | TCGACGCACTCCTCACA  |                 | Kageyama et al. (2003)      |
|                 |       | QNI5S-P                  | FAM-AGACATGGGGGAGGCATCG-QSY  |                 | Loisy et al. (2005)         |
| Hepatitis A virus| Assay 2| HAV68-F                  | TCACCGCCTGGCTTAC  | 173 bases        | Costafreda et al. (2006)    |
|                 |       | HAV240-R                 | GGAGGCCCTGGGAAAGAA  |                 | Costafreda et al. (2006)    |
|                 |       | HAV150-P                 | VIC-CCTGGACTGAGGAATTAA-QSY*  |                 | Costafreda et al. (2006)    |
| Mengovirus      | Assay 2| Me110-F                  | GCGGGTCCTGGCCGAAGT  | 100 bases        | Pinto et al. (2009)         |
|                 |       | Me209-R                  | GGATGAAATAGTAGGACGCCACAC  |                 | Pinto et al. (2009)         |
|                 |       | Me147-P                  | ABY-ATCATATTACTGCCCAGAG-NFQMGB*  |                 | Pinto et al. (2009)         |

*Original fluorescent dyes used for probes were replaced for multiplex assay development.

**F** forward primer, **R** reverse primer, **P** probe, **FAM** 6-Carboxyfluorescein, **NFQMGB** non-fluorescent quencher/minor groove binder
Reproducibility and Sensitivity of the Multiplex qRT-PCR Assay

To investigate reproducibility, the serial dilutions of spiked sediment and shellfish extracts were assayed using qRT-PCR on two plates in duplicate on each plate. The nominal concentrations of the spiked samples were $10^5$, $10^4$, $10^3$, $10^2$ and $10^1$ RNA copies/µL. To avoid RNA degradation, dilution series were freshly prepared before each run.

The limit of detection (LOD) and the limit of quantification (LOQ) were estimated according to the EN 24790 guideline (European Network of GMO laboratories 2011). Replicates of ten of the shellfish and sediment samples spiked with viral RNA were used at the nominal concentrations of 60, 40, 20, 10, 5 and 1 RNA copies/µL in each singleplex and multiplex assay. In order to determine RNA concentration in the samples, dilution series of the plasmid standards covering the concentration range of the samples were used. The lowest concentration where all replicates were positive was the estimated LOD. LOQ was estimated as the lowest concentration where the coefficient of variation (CV) amongst replicates was below 0.25.

Results

Multiplex Assay Performance

For each virus type, dilution series of plasmid standards incorporating the target genes were used for validation and quantification in singleplex and multiplex qRT-PCR assays. All standards showed excellent negative linearity in the range tested in both singleplex and multiplex assays (Fig. 1). The $R^2$ values ranged from 0.942 to 1 and the assay efficiency (E) ranged from 91.5 to 113.2%. The difference in these values were negligible between multiplex and singleplex assays. The corresponding standard curves overlapped in most cases, except in the SaV assays where lower Ct values were observed for all dilutions in the multiplex assay than in the singleplex.

Validation with Spiked Sediment and Shellfish Extract Samples

The usefulness of the singleplex and multiplex qRT-PCR assays for environmental samples was assessed using a serial dilution of sediment and shellfish nucleic acid extracts spiked with known concentration of the RNA of target viruses. Samples were run in duplicate in two assays to assess intra- and inter-assay variability. All samples tested were positive except the lowest concentration of MgV sample in shellfish extract, which was negative in both singleplex assays, but positive in the multiplex runs (Table 2). The CV values, based on the RNA copy concentration and standard deviation of duplicate samples in two assays ($n = 4$), were all lower than 0.5 in the high-concentration samples, except those calculated for the NoV GII in the sediment extracts and for HAV with the concentration of approx. 1000 copies/µL. The high variation noted for those samples may have been a result of pipetting error during sample preparation. Higher CV values were noted for SaV and HEV with <400 copies/µL and for other viruses with concentration <100 copies/µL.

The LOD and LOQ for the multiplex assays were determined by running ten replicates of low concentrations of spiked sediment and shellfish extracts. The LOD for all virus types was 1 RNA copy/µL. The highest LOQs were observed in sediment and shellfish extracts spiked with MgV (20 and 40 copies/µL, respectively). The LOQ of the other target viruses ranged from 5 to 10 copies/µL in spiked sediment and from 3 to 18 copies/µL in shellfish samples (Table 3).

Discussion

The accurate detection and quantification of a wide range of waterborne enteric viruses is essential to investigate health risks associated with wastewater contamination of environmental matrices. In this study, we evaluated the usefulness of two triplex one-step qRT-PCR assays targeting five strains of human enteric viruses (NoV GI/GII, HAV, HEV, SaV GI) and a murine cardiovirus (MgV), which is often used as an extraction control for shellfish sample processing.

All the primers and probes used have been described elsewhere (see Table 1 for references) and were shown to be specific to the target sequences. For some of the probes, the replacement of the published reporter and quencher dyes was necessary for multiplexing. For the SaV and HAV probes, the 6-carboxyfluorescein (FAM) reporter dye (fluorescent emission at 517 nm) was replaced with VIC (emission at 551 nm), and for the HEV and MgV probes it was replaced with ABY (emission at 580 nm). The TAMRA and the black hole quencher of the NoV GII and HEV probes were replaced with a non-fluorescent quencher, QSY. The replacements did not affect the melting temperature of the probes.

The Ta for the SaV and HEV assays was 56 °C (Chan et al. 2006; Jothikumar et al. 2006) and the primers and probes for those assays failed to align when the Ta was increased to 60 °C. Therefore, the Ta for the NoV GII assay was lowered to 56 °C from the reported value of 60 °C (International Organization for Standardization 2013) for multiplexing. Preliminary results based on the standard curves run at 56 and 60 °C showed no significant
The difference in PCR efficiency when the lower Ta was used (Fig. 1).

Performances of the singleplex and multiplex assays were first compared using a dilution series of DNA plasmids incorporating the target sequences. The overlapping standard curves suggested no competition for resources when multiplex approach was used. The only exception was the SaV assay where the standard curve of the multiplex assay slightly shifted (Fig. 1). However, the difference between the corresponding mean Ct values was below 1 and did not affect accurate quantification. Overall, the results suggested that all singleplex and multiplex assays are suitable for the quantification of the target sequences in a wide concentration range.

In order to further assess assay sensitivity and reproducibility, nucleic acid extracts of sediment and shellfish samples were spiked with known concentrations of viral RNA. The results suggested that the singleplex and multiplex assays were reproducible between approx. 10 and $10^5$ RNA copies/μL sample concentration, which refers to 50–5 × $10^5$ RNA copies/g sediment and 2 × $10^3$ – 2 × $10^7$ RNA copies/g shellfish assuming that the elution and nucleic acid extraction methods were used with 100% recoveries. These concentrations cover the range of viral nucleic acid concentration common in wastewater-contaminated sediment (Miura et al. 2011) and shellfish (Lowther et al. 2012). Assay sensitivity may be increased by increasing the sample volume or the volume of the qRT-PCR reaction mix (Le Mennec et al. 2016). Results revealed no inhibition due to residual organic matter in the nucleic acid extracts that has been shown to affect reverse transcriptase and polymerase enzymes (Farkas et al. 2017; Iker et al. 2013; Meschke and Sobsey 1998; Rock et al. 2010). The multiplex assay showed high sensitivity with a LOD of 1 RNA copy/μL. This refers to 10 RNA copies/g sediment and 200 RNA copies/g shellfish DT, assuming
100% recovery during viral elution and nucleic acid extraction. The LOQ values, varying between 3 and 40 copies/μL, allowed the accurate quantification of the samples tested.

The LOQ of multiplex qRT-PCR for the MgV was slightly higher (20 and 40 RNA copies/μL in sediment and shellfish extracts, respectively) than that observed for the enteric viruses. The MgV is usually added to a sample before extraction of the nucleic acids in known concentrations. According to the ISO/TS 152016-1 (2013), the extraction is considered successful when the recovery of the extraction control is above 1%, whereas the evaluation of the method used for the elution and concentration of viruses in sediment has shown recoveries above 70% (Farkas et al. 2017; Lewis and Metcalf 1988). Hence, we recommend the addition of $10^6$ intact MgV particles to samples prior to extraction when using the multiplex qRT-PCR (resulting Ct of 23-24) to assess process performance.

### Conclusions

The two multiplex qRT-PCR assays validated in this study allow the accurate quantification of the target viruses in sediment and shellfish nucleic acid extracts. Multiplexing enables the identification of different target pathogens in one assay, lowering the cost and time associated with qRT-PCR. The use of MgV as an extraction control allows the
quality assessment of the extraction method used prior to qRT-PCR, lowering the possibility of false-negative outcomes. Overall, these assays allow the thorough assessment of the target viruses in sediment and shellfish samples and are useful for quantitative risk assessment of wastewater-contaminated environments.

Acknowledgements This work was funded by the Natural Environment Research Council (NERC) and the Food Standards Agency (FSA) under the Environmental Microbiology and Human Health (EMHH) Programme (NE/M010996/1). The authors would like to thank Prof. Ian Goodfellow (University of Cambridge, UK) and Dr. Lisa Cross (CEFAS, UK) for providing virus, RNA and DNA samples. We thank Prof. Alan McCarthy (University of Liverpool) for his helpful review comments on the manuscript.

Compliance with Ethical Standards

Conflict of interest The authors declare that they have no competing interest.

Open Access This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made.

References

Bellou, M., Kokkinos, P., & Vantarakis, A. (2013). Shellfish-borne viral outbreaks: A systematic review. Food and Environmental Virology, 5(1), 13–23.

Bosch, A., Pintó, R. M., & Guix, S. (2016). Foodborne viruses. Current Opinion in Food Science, 8, 110–119.

Centers for Disease Control and Prevention. (2015). Norovirus-U.S. Trends and Outbreaks. http://www.cdc.gov/norovirus/trends-outbreaks.html 2013.

Chan, M. C., Sung, J. J., Lam, R. K., Chan, P. K., Lai, R. W., & Leung, W. K. (2006). Sapovirus detection by quantitative real-time RT-PCR in clinical stool specimens. Journal of Virological Methods, 134(1), 146–153.

Costafreda, M. I., Bosch, A., & Pintó, R. M. (2006). Development, evaluation, and standardization of a real-time TaqMan reverse transcription-PCR assay for quantification of hepatitis A virus in clinical and shellfish samples. Applied and Environmental Microbiology, 72(6), 3846–3855.

da Silva, A. K., Le Saux, J. C., Parnaudeau, S., Pommepuy, M., Elimelech, M., & Le Guyader, F. S. (2007). Evaluation of removal of noroviruses during wastewater treatment, using real-time reverse transcription-PCR: Different behaviors of genotypes I and II. Applied and Environmental Microbiology, 73(24), 7891–7897.

Dalton, H. R., Hunter, J. G., & Bendall, R. P. (2013). Hepatitis E Current Opinion in Infectious Diseases, 26(5), 471–478.

European Network of GMO laboratories. (2011). EUR 24790 EN Verification of analytical methods for GMO testing when implementing interlaboratory validated methods. JRC Scientific and Technical Reports.

Farkas, K., Hassard, F., McDonald, J. E., Malham, S. K., & Jones, D. L. (2017). Evaluation of molecular methods for the detection and quantification of pathogen-derived nucleic acids in sediment. Frontiers in Microbiology. doi:10.3389/fmicb.2017.00053.

Farkas, T., Singh, A., Le Guyader, F. S., La Rosa, G., Saif, L., & McNeal, M. (2015). Multiplex real-time RT-PCR for the simultaneous detection and quantification of GI, GII and GIV noroviruses. Journal of Virological Methods, 223, 109–114.

Fuentes, C., Guix, S., Pérez-Rodríguez, F. J., Fuster, N., Carol, M., Pintó, R. M., et al. (2014). Standardized multiplex one-step qRT-PCR for hepatitis A virus, norovirus GI and GII quantification in bivalve mollusks and water. Food Microbiology, 40, 55–63.

Hennechart-Collette, C., Martin-Latil, S., Guillier, L., & Perelle, S. (2015). Determination of which virus to use as a process control when testing for the presence of hepatitis A virus and norovirus in food and water. International Journal of Food Microbiology, 202, 57–65.

Hoehne, M., & Schreier, E. (2006). Detection of Norovirus genogroup I and II by multiplex real-time RT-PCR using a 3’mminor groove binder-DNA probe. BMC Infectious Diseases. doi:10.1186/1471-2334-6-69.

Iker, B. C., Bright, K. R., Pepper, I. L., Gerba, C. P., & Kitajima, M. (2013). Evaluation of commercial kits for the extraction and purification of viral nucleic acids from environmental and fecal samples. Journal of Virological Methods, 191(1), 24–30.

International Organization for Standardization. (2013). ISO/TS 15216-1:2013 Microbiological of food and animal feed—Horizontal method for determination of hepatitis A virus and norovirus in food using real-time RT-PCR—Part 1: Method for quantification.

Jothikumar, N., Crombeans, T. L., Robertson, B. H., Meng, X., & Hill, V. R. (2006). A broadly reactive one-step real-time RT-PCR assay for rapid and sensitive detection of hepatitis E virus. Journal of Virological Methods, 131(1), 65–71.
Kageyama, T., Kojima, S., Shinohara, M., Uchida, K., Fukushima, S., Hoshino, F. B., et al. (2003). Broadly reactive and highly sensitive assay for Norwalk-like viruses based on real-time quantitative reverse transcription-PCR. *Journal of Clinical Microbiology, 41*(4), 1548–1557.

Le Mennec, C., Parnaudeau, S., Rumebe, M., Le Saux, J., Piquet, J., & Le Guyader, S. F. (2016). Simultaneous detection of selected enteric viruses in water samples by multiplex quantitative PCR. *Water, Air, and Soil Pollution, 227*(4), 1–11.

Lee, D., Leung, K. T., Lee, H., & Habash, M. B. (2016). Simultaneous screening in shellfish. *Food and Environmental Virology, 2*(3), 146–155.

Lewis, G. D., & Metcalf, T. G. (1988). Polyethylene glycol precipitation for recovery of pathogenic viruses, including hepatitis A virus and human rotavirus, from oyster, water, and sediment samples. *Applied and Environmental Microbiology, 54*(8), 1983–1988.

Loisy, F., Atmar, R. L., Guillon, P., Le Cann, P., Pommepuy, M., & Le Guyader, F. S. (2005). Real-time RT-PCR for norovirus screening in shellfish. *Journal of Virological Methods, 123*(1), 1–7.

Lowther, J. A., Gustar, N. E., Powell, A. L., Hartnell, R. E., & Lees, D. N. (2012). Two-year systematic study to assess norovirus contamination in oysters from commercial harvesting areas in the United Kingdom. *Applied and Environmental Microbiology, 78*(16), 5812–5817.

Martin-Latil, S., Hennechart-Collette, C., Guiller, L., & Perelle, S. (2012). Duplex RT-qPCR for the detection of hepatitis E virus in water, using a process control. *International Journal of Food Microbiology, 157*(2), 167–173.

Meschke, J., & Sobsey, M. (1998). Comparative adsorption of Norwalk virus, poliovirus 1 and F RNA coliphage MS2 to soils suspended in treated wastewater. *Water Science and Technology, 38*(12), 187–189.

Miura, T., Masago, Y., Sano, D., & Omura, T. (2011). Development of an effective method for recovery of viral genomic RNA from environmental silty sediments for quantitative molecular detection. *Applied and Environmental Microbiology, 77*(12), 3975–3981.

Pinto, R. M., Costafreda, M. I., & Bosch, A. (2009). Risk assessment in shellfish-borne outbreaks of hepatitis A. *Applied and Environmental Microbiology, 75*(23), 7350–7355.

Purcell, R. H. (1996). Hepatitis E virus. In B. N. Fields, D. M. Knipe, & P. M. Howley (Eds.), *Fields virology* (pp. 2831–2843). Philadelphia: Lippincott-Raven Publishers.

Rao, V. C., & Melnick, J. L. (1986). Introduction. In V. C. Rao & J. L. Melnick (Eds.), *Environmental virology* (pp. 1–9). Wokingham: Van Nostrand Reinhold.

Rock, C., Alum, A., & Abbasszadegan, M. (2010). PCR inhibitor levels in concentrates of biosolid samples predicted by a new method based on excitation-emission matrix spectroscopy. *Applied and Environmental Microbiology, 76*(24), 8102–8109.

Staggemeier, R., Bortoluzzi, M., da Silva Heck, T. M., da Luz, R. B., Fabres, R. B., Soliman, M. C., et al. (2015a). Animal and human enteric viruses in water and sediment samples from dairy farms. *Agricultural Water Management, 152*, 135–141.

Staggemeier, R., Bortoluzzi, M., da Silva Heck, T. M., da Silva, T., Spilki, F. R., & de Matos Almeida, S. E. (2015b). Molecular detection of human adenovirus in sediment using a direct detection method compared to the classical polyethylene glycol precipitation. *Journal of Virological Methods, 213*, 65–67.

Svraka, S., Duizer, E., Vennema, H., de Bruin, E., van der Veer, B., Dorresteijn, B., et al. (2007). Etiological role of viruses in outbreaks of acute gastroenteritis in The Netherlands from 1994 through 2005. *Journal of Clinical Microbiology, 45*(5), 1389–1394.

Tahaei, S. M., Mohebbi, S. K., & Zali, M. R. (2012). Enteric hepatitis viruses. *Gastroenterology and Hepatology From Bed to Bench, 5*(1), 7–15.

Tam, C. C., Rodrigues, L. C., Viviani, L., Dodds, J. P., Evans, M. R., Hunter, P. R., et al. (2012). Longitudinal study of infectious intestinal disease in the UK (IID2 study): incidence in the community and presenting to general practice. *Gut, 61*(1), 69–77.

van Maarseveen, N. M., Wessels, E., de Brouwer, C. S., Yomena, A. C., & Claas, E. C. (2010). Diagnosis of viral gastroenteritis by simultaneous detection of Adenovirus group F, Astrovirus, Rotavirus group A, Norovirus genogroups I and II, and Sapovirus in two internally controlled multiplex real-time PCR assays. *Journal of Clinical Virology, 49*(3), 205–210.

Yan, Y., Wang, H., Gao, L., Ji, J., Ge, Z., Zhu, X., et al. (2013). A one-step multiplex real-time RT-PCR assay for rapid and simultaneous detection of human norovirus genogroup I and IV. *Journal of Virological Methods, 189*(2), 277–282.

Yates, M. V. (2014). Chapter Twenty-Nine - Emerging Viruses. In S. L. Percival, M. V. Yates, D. W. Williams, R. M. Chalmers, & N. F. Gray (Eds.), *Microbiology of Waterborne Diseases* (2nd ed., pp. 529–533). London: Academic Press.