Human virus and microbial indicator occurrence in public-supply groundwater systems: meta-analysis of 12 international studies

G. Shay Fout\(^1\), Mark A. Borchardt\(^2\), Burney A. Kieke Jr\(^3\), and Mohammad R. Karim\(^4\)

\(^1\)US Environmental Protection Agency, 26 Martin Luther King Dr, Cincinnati, OH 45268, USA
\(^2\)US Department of Agriculture, 2611 Yellowstone Dr, Marshfield, WI 54449, USA
\(^3\)Marshfield Clinic Research Foundation, 1000 Oak Ave, Marshfield, WI 54449, USA
\(^4\)City of Santa Cruz, Public Works Department, 110 California St, Santa Cruz, CA 95060, USA

Abstract

Groundwater quality is often evaluated using microbial indicators. This study examines data from 12 international groundwater studies (conducted 1992–2013) of 718 public drinking-water systems located in a range of hydrogeological settings. Focus was on testing the value of indicator organisms for identifying virus-contaminated wells. One or more indicators and viruses were present in 37 and 15% of 2,273 samples and 44 and 27% of 746 wells, respectively. *Escherichia coli* (*E. coli*) and somatic coliphage are 7–9 times more likely to be associated with culturable virus-positive samples when the indicator is present versus when it is absent, while F-specific and somatic coliphages are 8–9 times more likely to be associated with culturable virus-positive wells. However, single indicators are only marginally associated with viruses detected by molecular methods, and all microbial indicators have low sensitivity and positive predictive values for virus occurrence, whether by culturable or molecular assays, i.e., indicators are often absent when viruses are present and the indicators have a high false-positive rate. Wells were divided into three susceptibility subsets based on presence of (1) total coliform bacteria or (2) multiple indicators, or (3) location of wells in karst, fractured bedrock, or gravel/cobble settings. Better associations of some indicators with viruses were observed for (1) and (3). Findings indicate the best indicators are *E. coli* or somatic coliphage, although both indicators may underestimate virus occurrence. Repeat sampling for indicators improves evaluation of the potential for viral contamination in a well.

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.

Compliance with ethical standards

Publisher's Disclaimer: Disclaimer This report has been subjected to the US Environmental Protection Agency’s peer and administrative review and has been approved for publication. The mention of trade names or commercial products in this report does not constitute endorsement or recommendation for use.
Keywords
Contamination; Drinking water; Enteric virus; Microbial indicators; Health

Introduction

Groundwater is an important source of drinking water in both developed and developing countries. It constitutes about 95% of the world’s accessible freshwater (Chilton and Seiler 2006; Howard et al. 2006; McKay 2011) and is often used with little or no treatment (Pedley et al. 2006). Contamination of groundwater with human enteric viruses is a global issue (Blaschke et al. 2016; Gotkowitz et al. 2016; Hynds et al. 2014; USEPA 2006a, b), as consumption of contaminated water can result in elevated rates of endemic illness and waterborne disease outbreaks in affected communities (Beer et al. 2015; Borchardt et al. 2011; Cho et al. 2014; Guzman-Herrador et al. 2015; Hilborn et al. 2013; Jack et al. 2013; Wallender et al. 2014; Zhou et al. 2012). Enteric viruses implicated in waterborne outbreaks include enteroviruses, hepatitis A, rotavirus, and norovirus (Craun et al. 2010; Hejkal et al. 1982), but others such as adenoviruses, Aichi virus 1, hepatitis E, and reoviruses, potentially are capable of groundwater-borne transmission. Waterborne viruses cause a wide range of illnesses, including gastroenteritis, paralysis, aseptic meningitis, conjunctivitis, diabetes, fevers, herpangina, rash, myocarditis, and respiratory illness, (Kitajima and Gerba 2015; WHO 2011).

Several nonpathogenic bacteria that are normal flora of the human intestine and other warm-blooded animals can be detected using simple and inexpensive techniques. These include total coliform bacteria, fecal (thermotolerant) coliform bacteria, E. coli, enterococci, and bacterial endospores (Ashbolt et al. 2001; Locas et al. 2007; Tallon et al. 2005). Because pathogens, and especially viral pathogens, occur sporadically and are difficult to detect, these bacteria have been used as indicators of fecal pollution. Total coliform bacteria and E. coli are the most commonly employed water quality indicators, but of the two, E. coli is a more definitive indicator of fecal pollution (Edberg et al. 2000; Tallon et al. 2005).

Coliphages (bacteriophages that infect coliform bacteria) are present in wastewater and have been suggested to be a useful conservative indicator of fecal and viral pollution of groundwater (Deborde et al. 1998a, b; Lucena et al. 2006). They are grouped into two major categories—F-specific and somatic coliphages. F-specific coliphages are similar in size (about 26 nm in diameter) and structure (icosahedral) to human enteric viruses (about 28–38 nm in diameter for enteroviruses and noroviruses), while somatic coliphages have more variability. Unlike enteric viruses, coliphages are detectable by simple, inexpensive and rapid techniques (Lucena et al. 2006). Phages of Bacteroides have also been evaluated as useful indicators (Jofre et al. 1986; Johnson et al. 2011; McMinn et al. 2014).

Although bacterial indicators are used for general water quality monitoring, they are not considered to be adequate indicators of viral contamination because the structure, composition, morphology, size (about 1,000 nm in diameter x 2,000 nm in length for E. coli), and survival characteristics of viruses differ fundamentally from those of bacteria (Ashbolt 2015; Payment and Locas 2011; WHO 2011). Viruses survive longer than
vegetative bacteria in the environment (Nasser et al. 1993) and have different transport properties. The usefulness of anaerobic bacterial endospores is limited by very long survival times, the ubiquity of Clostridium species in soil, and by transport properties (Meschke 2001). Aerobic endospores also have major differences in transport and survival properties (Headd and Bradford 2016; Pang et al. 2005).

Viral and other pathogens and microbial indicators enter aquifers through multiple sources and pathways, including leachates from sanitary landfills, on-site septic waste treatment discharges, broken sewage lines, runoff from urban, agricultural and natural areas, and water reuse by direct injection of inadequately treated wastewater into aquifers (Borchardt et al. 2012; Costan-Longares et al. 2008; Gotkowitz et al. 2016). Fecal contamination from the surface may also get into groundwater through improperly constructed, protected, or maintained wells (Hynds et al. 2014).

Despite the potential public health impact from drinking untreated groundwater, the regulations guiding the quality of groundwater supplies used for drinking water are highly varied among countries. Many use the World Health Organization Guidelines for Drinking Water Quality (WHO 2011), which covers drinking water from all sources and suggests that drinking water contain no thermotolerant coliform bacteria or E. coli per 100 ml. Canada’s guidelines state that drinking water should have no detectable total coliforms or E. coli per 100 ml (Health Canada 2017). Australia’s guidelines add intestinal enterococci and coliphages in addition to thermotolerant bacteria and E. coli as agents that should not be detected in 100 ml and suggest that sanitary surveys be conducted for groundwater systems (National Health and Medical Research Council 2016). Sanitary surveys are used to examine a system’s deficiencies that could cause vulnerability to microbial contamination. The surveys are intended to identify deficiencies caused by poor source water, inadequate well construction or maintenance, and improper system operation. Monitoring of water quality is required by regulation in some countries—for example, Japan requires monitoring for E. coli (Wakayama 2016). Korea does periodic monitoring for enteric viruses and their Groundwater Act requires monitoring of groundwater for E. coli (Lee and Kwon 2016; Lee H et al. 2011; Lee S et al. 2011). The European Union requires regulatory authorities to monitor all public drinking-water systems for coliform bacteria and E. coli at a frequency that depends on water volume pumped and/or population served. The United States regulatory instrument is the Ground Water Rule (GWR; USEPA 2006a, b). The key components of the GWR are: sanitary surveys, triggered source water monitoring, corrective actions, and compliance monitoring. Source monitoring may be required rather than triggered if the source water quality is uncertain. The triggered water monitoring provision requires that untreated groundwater systems must conduct triggered source water monitoring for the presence of at least one of the three fecal indicators: E. coli, enterococci, or coliphage following a Revised Total Coliform Rule (RTCR)-positive sample. The RTCR requires regular monitoring of all public systems for total coliforms using 100-ml samples at a frequency that depends on the population served. Any regular monitoring sample that contains any amount of total coliforms must be followed by additional tests for both total coliforms and E. coli. The compliance monitoring provision of the GWR requires that systems that provide 4-log treatment of viruses must conduct compliance monitoring to demonstrate continual treatment effectiveness. The GWR uses a risk-targeted approach to
address the likelihood of viral contamination of wells—i.e., well susceptibility to contamination (note: susceptibility herein is defined by the entry of fecal contamination into an aquifer as measured by the demonstrated presence of virus in any sample from an associated well). It suggests that the agencies that implement the rule consider groundwater from aquifers in limestone, igneous and metamorphic rock, and gravel as potentially susceptible. However, among aquifers in gravel settings, gravel/cobble aquifers are more susceptible than those consisting of gravel/sand (Berger 2008). Aquifers in gravel settings have rock grain sizes of 4 mm or larger, whereas those in gravel/cobble settings have grain sizes up to about 256 mm (Wentworth 1922).

While a good indicator would be expected to be positively correlated to virus presence (i.e., always present when viruses are present and absent when viruses are absent), these types of correlations are not always observed in groundwater (Payment and Locas 2011). This might be due to indicator and virus differing in die-off properties, transport characteristics, waste treatment practices before release to the environment, source concentrations—e.g., indicators are constantly present and at higher concentrations in wastewater than are viruses (Berg et al. 1978)—and virus infection dynamics and shedding rates in the host population. In addition, in an analysis of studies correlating indicators and pathogens in water types other than groundwater, Wu et al. (2011) showed that the strength of correlations was related to study sample size and the number of pathogen-positive detections. Mindful of these limitations, this study examined the relationship between indicators and viruses by combining data from 12 studies of public drinking-water groundwater systems. In addition to conventional tests of correlation, the study evaluated the association between indicator and virus occurrence by logistic regression and calculated standard performance measures for diagnostic tests, namely, sensitivity, specificity, positive predictive value, and negative predictive value. These measures were further evaluated for three subsets of wells considered susceptible to fecal contamination based on hydrogeological setting and US regulations.

Materials and methods

Site selection

Raw data from 12 studies on viruses in groundwater were used to determine the association between microbial indicators and virus occurrence. Groundwater studies were selected primarily on the basis of availability of raw data supplied in publications or from the corresponding authors (Table 1). Secondly, studies were selected to give a broad range of hydrogeological settings in North America, Europe, and Asia, and to avoid over-representing one location or study team in the combined dataset.

Summary of included studies

A summary of each study is provided with a focus on settings and virus occurrence. The American Water Works Association Research Foundation (AWWARF, now Water Research Foundation) and USEPA conducted a study of viruses in groundwater in the US public systems during 1992–1994 (Dahling 2002; Fout et al. 2003; Lieberman et al. 2002; USEPA 2006a, b). This study focused on 30 wells, with 13 located in karst, fractured bedrock, or
course gravel settings; 11 in alluvial settings with frequent microbial indicator-positive samples, and 6 in unknown or in alluvial settings without a record of frequent indicator detections. All but one of the seven culturable virus-positive wells were in karst or fractured bedrock locations, while only 11 of the 25 sites positive for virus by molecular tests were in karst or fractured bedrock settings. A culturable virus-positive sample means that infectious viruses are present in water from the aquifer. Culturable virus methods detect only a narrow range of those viruses that cause waterborne disease, so a negative result should not be interpreted to mean that all infectious viruses are absent. Molecular methods can detect most of the viruses that cause waterborne illness, but most studies only assay for a limited number of them. The primary limitation of molecular methods is their inability to determine whether detected viruses are infectious. They also may be affected by water chemistries that cause a false-negative reaction.

The US Geological Survey (USGS) and Public Drinking Water Program of the Missouri Department of Natural Resources conducted a two-phase study of 182 public water supplies in the Ozark plateaus aquifer system in Missouri during 1997–1998 (Davis and Witt 2000; Davis and Witt 1998; Femmer 2000). Public supply wells were selected to represent settings in primary karst, secondary karst, the confined Ozark aquifer, or in alluvium. One sample was positive for culturable virus and that was from a source located in an area of confined primary karst; however, only three of the 13 samples positive by molecular assays were in karst settings.

Francy et al. (2004a, b) conducted a study of 38 groundwater-supply wells during 1999–2001. This study targeted public systems that served a population size between 25 and 3,300 in Silurian-Devonian sand, gravel, and clay aquifers in southeastern Michigan, USA. The aquifers in this area consist typically of dual layers of glacial outwash and till with unconfined upper layers and semi-confined lower layers. Culturable virus was detected at two of these sites and virus by molecular tests at seven.

The USGS conducted a study of microbial indicators and virus from 60 non-community public water supplies in Pennsylvania during 2000–2001 (Lindsey et al. 2002). Twenty-five sites were located in karstic limestone or dolomite, with another 25 sites situated in fractured bedrock. Five sites each were in siliciclastic settings and in unconsolidated sediments. Culturable viruses were detected in two wells in a karstic setting, at one well in a fractured bedrock setting, and at two wells in a siliciclastic setting.

The American Water Service Company performed a study of 20 public drinking water wells from 11 US states during 2001–2002. Wells were selected from the first round of a larger study (Abbaszadegan et al. 2003) based upon the presence of culturable virus (five wells), viral nucleic acid (six wells), indicator bacteria (five wells), and an absence of both indicators and virus (four wells). Three of the wells were in fractured bedrock settings, with most others being in alluvial or glacial sand and gravel. Culturable virus was present at seven sites and virus by molecular assays at 15 sites; two of the three wells in fractured bedrock were virus-positive, one for culturable virus and the other for virus measured by PCR.
A small study of springs and wells was conducted in Tennessee (USA) during 2004 (Johnson et al. 2011). Three wells were selected from a larger group of wells to represent "low" susceptibility to fecal contamination. This was based upon the absence of E. coli during an initial period of monitoring of the wells for microbial indicators (Johnson 2005). One of these wells was in a Pre-Conasagua group carbonate aquifer overlain by 35 m of residuum. Another was in a Copper Ridge dolomite (Knox Group) aquifer, overlain by 30–46 m of overburden. The third was in Chilhowee Group Sandstone and Conglomerate aquifer, overlain by 0–2 m of residuum. The fourth well, located in a Knox Group Carbonate aquifer, overlain by 9 m of residuum, was chosen to represent a susceptible well based upon the prior sampling. Culturable virus was detected at two of the three wells in the low susceptibility group and in the well in the high susceptibility group.

A year-long study of 12 municipal wells in Quebec, Canada, was conducted during 2003–2004 to examine the influence of different aquifers, soil types, and well depths on virus and indicator occurrence (Locas et al. 2007). Groups of public systems were selected consisting of (1) wells tapped into glacial deposits with and without confinement and no history of microbial detections; (2) wells in glacial deposits or fractured bedrock with sporadic detection of total coliforms; and (3) wells in unconfined sand and gravel aquifers or fractured sandstone with frequent detections of total and fecal indicators. A follow-up study was conducted during 2006 and 2012 on 24 municipal wells in three provinces in Canada (Locas et al. 2008; Payment and Locas 2005; P. Payment, Institut Armand-Frappier, 2015, personal communication), including two group three sites from the first study. No data were collected on the hydrogeology of the additional wells, but none appeared to be in karst or fractured bedrock locations (Ford 1997). Overall, culturable virus was detected at four sites, two of which were in karst or fractured bedrock sites. Norovirus was detected at three unconfined aquifers.

A small study of eight groundwater wells in karstic settings was conducted during 2005 in central Italy by the University of Rome Tor Vergata (Gabrieli et al. 2009). Samples were tested for virus using only molecular assays. All were negative for fecal indicators, while three wells (38%) were positive for norovirus.

A Japanese study of 46 wells was conducted by the University of Tokyo during 2005–2006 in the eastern lowland area of Tokyo (Katayama 2008). About half of the wells received water from the unconfined Yurakucho alluvial sand aquifer, while the other half obtained water from deeper confined aquifers. Samples were tested for adenovirus using real time PCR with four wells being positive, two of which were in confined and two in unconfined aquifers.

A study of 36 public wells that supplied untreated drinking water to 14 small communities (population of 1,363–8,300) was conducted by the Marshfield Clinic Research Foundation in Wisconsin (USA) during 2006–2007 (Borchardt et al. 2012). Wells were primarily located in non-karstic, sandstone settings (Lambertini et al. 2011). The setting of six communities was sand and gravel or mixtures of sand, gravel, and sandstone. Two communities had limestone and dolomite or sandstone with limestone and dolomite aquifers. Two communities were located close to regions in the state that are karstic, but do not appear to
be influenced by karst. Well depths ranged from 19 to 173 m and pumping rates from about 500,000 L/day for the smallest community to 14,500,000 L/day for the largest community. All samples were tested for adenovirus, enterovirus, hepatitis A virus, norovirus genogroups I and II, and rotavirus by real time PCR. Samples that were positive for adenovirus and enterovirus by PCR were also tested for culturable viruses using integrated cell culture-quantitative PCR. Overall, 31 of the wells were positive for virus and about a quarter of the PCR-positive samples tested contained culturable virus.

South Korea’s National Institute of Environmental Research (NIER) initiated several groundwater monitoring studies in metropolitan areas and provinces during 2007–2011 (Jung et al. 2011; Lee et al. 2013; Lee H et al. 2011; Lee S et al. 2011) with data being provided for 220 sites. No information was given on hydrogeology of the sites, but four were in areas with porous volcanic rock or with a high likelihood of being in karst regions (University of Auckland 2008). Other sites would be primarily alluvial as the hydrogeology of most of the Korean peninsula is poorly permeable crystalline granitic and metamorphic rocks (Won et al. 2005), and thus associated with low groundwater yields. Wells were frequently positive for viruses by PCR, with 30% being positive for norovirus, 13% for adenovirus, and 8% for enterovirus. No virus was present in the volcanic or karst sites. More than half of the norovirus positive samples were from genogroup I genotypes.

A study of public systems in Iowa (USA) was conducted in the spring of 2013 (Hruby et al. 2015). A total of 63 systems (with 66 wells) were chosen covering the major hydrogeological areas of the state. These included aquifers in alluvial settings (18% of wells), sand and gravel (11%), Cambrian-Ordovician sandstone (18%), Dakota Cretaceous sandstone (11%), Mississippi sandstone and carbonate (8%), and Silurian-Devonian carbonate (35%). Although a few areas in the northeast portion of the state have strong karst features (Horick 1984), all of the studied wells appear to be outside the karst areas. Samples were analyzed for microbial indicators, pathogens, and numerous chemical contaminants. Despite the large number of wells in carbonate and sandstone aquifers, there was virtually no microbial contamination of the wells. One site had norovirus genogroup II, but all were negative for adenoviruses, enteroviruses, or hepatitis E virus. Samples were collected following a severe drought during 2012, which may have been a contributing factor to the low microbial contamination observed during the study. This study was the first to include measurements of pepper mild mottle virus (PMMV). This virus comes from foods eaten with peppers and occurs at high levels in sewage (Kuroda et al. 2015). Eleven samples in the Iowa study were positive for PMMV, suggesting that this virus may be a better conservative indicator of microbial contamination from human sources than bacterial or bacteriophage indicators; however, Kuroda et al. (2015) suggest the detection rate for PMMV in groundwater is lower than that of human enteric viruses.

Data handling

This study analyzed available raw data from the 12 groundwater virus occurrence studies described in the preceding, covering 2,273 samples from 718 drinking water systems and 746 wells (Table 1). Table 2 describes the quantity and type of data provided by each study. Culturable virus occurrence and virus occurrence measured by PCR methods (hereafter...
designated PCR-virus) was provided by six studies. One study provided only culturable virus occurrence data, while five studies provided only PCR-virus data. Some studies provided qualitative data for some analytes. For purposes of analysis, all qualitative data were converted to quantitative data as detailed in Tables 3 and 4. In addition, all concentrations were normalized to 100 ml for making comparisons across the studies.

**Statistical analysis**

Spearman rank order correlations (SigmaPlot) between concentrations of viruses and indicators were evaluated at two levels, sample and well. By well, the sum of virus concentrations was compared to the sum of indicator concentrations. As many samples were negative for virus or indicator (i.e., below the detection limit of the assays) which might result in spurious correlations, the analyses were repeated using a dataset restricted to culturable virus and PCR-virus positive samples.

The utility of water quality indicators as indicators of virus-contaminated groundwater was evaluated by testing the association between indicators and virus and by calculating the four conventional performance measures of a diagnostic test: sensitivity (i.e. true positive rate), specificity (i.e., true negative rate), positive predictive value, and negative predictive value (Borchardt et al. 2003) for the 12-study combined dataset. For these association and performance analyses, each of the six indicators—total coliforms, *E. coli*, enterococci, F-specific phage, somatic phage, spores (anaerobic spores + aerobic spores)—was treated as a dichotomous variable (detect or non-detect) and compared to two measures of virus contamination: culturable virus and PCR-virus. Both outcome measures were also treated as dichotomous variables (i.e., detect/non-detect). The strength of the indicator-virus association additionally was quantified by calculating the risk ratio (positive predictive value/[1-negative predictive value]). The null value for the risk ratio is 1.0 with values greater than 1.0 representing the relative elevation in the virus detection rate if the indicator is detected versus when the indicator is not detected. The preceding analyses were conducted separately for data at the level of the sample and well and all results were derived from logistic regression models.

Two logistic regression model formulations that addressed the non-independent nature of the data were employed. The primary analyses entailed fitting mixed models with random intercepts for study and well (sample-level analyses) and study only (well-level analyses). These models also incorporated robust variance estimation (Chavance and Escolano 2016; Morel et al. 2003). The second model formulation involved fitting population-averaged models with robust variance estimation (Morel et al. 2003). With respect to the point estimates of the risk ratio, sensitivity, specificity, positive predictive value and negative predictive value; the mixed model formulation can be considered adjusted (i.e., for study in all analyses and additionally for well in the sample-level analyses), whereas the latter formulation can be viewed as unadjusted. To facilitate model convergence, the independence covariance structure was used in all models. All logistic regression modeling was performed using PROC GLIMMIX of the SAS software (SAS Institute, Inc., Cary, NC).

The aforementioned analytic framework was applied separately to all wells and to three separate non-mutually-exclusive subsets of wells satisfying each susceptibility criterion.
The first susceptibility subset included all wells that either have or have potential to have multiple violations of the RTCR (hereafter, TCR). The second susceptibility subset included all wells located in karst, fractured bedrock, or gravel/cobble hydrogeological settings. The last subset was based on the triggered source water monitoring provision of the GWR. This included wells with total coliforms and any of the additional follow-up indicators specified by the GWR.

When conducting well-level analyses, an attempt was made to include an adjustment in the regression models for the number of times the well was sampled (NTWS), since this could affect the corresponding probability of detection for the well; however, this resulted in model convergence problems. Within each study NTWS was generally homogeneous, either a small number of similar values or dominated by a single value. It is possible that the model adjustment for study lessened any effect of NTWS, but this may be a limitation of the well-level analyses.

**Results**

Table 6 shows the percent of samples and wells that were positive for indicators and virus from the 12 groundwater studies. Total coliforms were detected in 31% of samples and 36% of wells. Overall, 37% of samples and 45% of wells were positive for any indicator, while 15% of samples and 27% of wells were positive for any enteric viruses. The average titer of positive samples containing culturable- or PCR-virus from the 12 studies was 0.4 infectious units and 16 genomic copies per liter, respectively (data not shown).

Spearman Rank Order tests showed that among the indicators there were positive and moderately strong correlations (ρ ≥0.5, P < 0.001, n > 1,300) for combinations of total coliforms, *E. coli*, and enterococci, and between enterococci and somatic phage (data not shown). Between the indicators and culturable virus concentrations on a per-sample basis the trends were positive but weak (ρ ≤0.3, P < 0.001, n > 1,200; Table 7). Correlations on a per-well basis were also weak with the highest between somatic coliphage and culturable virus (ρ = 0.46, P < 0.001, n = 355). Correlations among indicators and PCR-virus (Table 7) were always weaker than to culturable virus.

The analysis was repeated by restricting the data to virus-positive samples (or wells) to minimize the effect of non-detects on the correlations. With this restricted data set, the highest correlations to culturable virus were *E. coli* (ρ = 0.62, P < 0.001, n = 144) and somatic coliphage (ρ = 0.54, P < 0.001, n = 141) on a per-sample basis and somatic coliphage (ρ = 0.48, P < 0.001, n = 76) on a per-well basis; however, restricting the data set did not improve the correlations among the indicators and virus detected by molecular assays (data not shown).

Subsequently, the association between indicator and virus occurrence was evaluated (detect or non-detect) using logistic regression models and the data from the 12 studies. Culturable viruses were associated at statistically significant levels with the indicators except for total coliforms measured at the sample level and spores measured at the well level (Table 8). In contrast, PCR-viruses and indicators at the sample level were never statistically associated.
and at the level of well only three of the six indicators (total coliforms, *E. coli*, and somatic phage), were significantly associated with PCR-viruses. The two most commonly used indicators in the US—total coliforms and *E. coli*—were associated at the well level with both culturable viruses and PCR-viruses, albeit the *E. coli* and culturable virus association was marginally not significant (*P* = 0.087; Table 8).

The risk ratios reported in Table 8 give the relative increase in the probability of detecting a virus-positive sample (or well) when an indicator is detected compared to when an indicator is not detected—for example, a positive *E. coli* sample is associated with a seven-times greater chance of detecting culturable viruses in a corresponding sample compared to a negative *E. coli* result. For PCR-viruses, the chance of detection in a sample is similar whether a corresponding *E. coli* sample is positive or negative (risk ratio = 0.9, *P* = 0.79). However, at the level of well, an *E. coli* positive well is associated with a 60% greater chance the well, at some time, will be positive for PCR-viruses (risk ratio = 1.6, *P* = 0.008).

Table 8 reports the associations and risk ratios adjusted for study (i.e., accounting for underlying differences among the 12 studies) and the sample-level analyses additionally include an adjustment for wells. Analyses were also conducted without these adjustments (data not shown). Generally, the adjustments resulted in lower estimated risk ratios and similar conclusions regarding statistical significance of the indicator-virus association, suggesting the probability of detecting a virus when an indicator is present differed across the individual studies and the wells within each study. The effect of the adjustments on the risk ratios could reflect differences in hydrogeological settings, virus contamination sources, laboratory methods, or several unknown factors at the level of study or well that are related to virus and indicator occurrence. Nonetheless, for this evaluation of indicators and viruses in groundwater, it is evident it was important to account for the effects of both study and well.

Indicator test performance measures were examined next. Sensitivities of the indicators for signifying whether a virus was detected in a sample or well were low (2–30%), with estimates being relatively higher when virus positivity was determined by culture (11–30%, Table 9) versus PCR (2–12%, Table 10); note that all test performance measures reported here are adjusted for study and well (sample-level analyses) or study only (well-level analyses). By culture, between 11 and 30% of virus-positive samples and between 37 and 73% of virus-positive wells could be correctly identified as virus-positive by an indicator. In other words, many samples and wells negative for an indicator were, in fact, positive for virus. The corresponding numbers for PCR were ≤12% for samples and ≤39% for wells.

Positive predictive values were also low for the indicators in predicting the detection of both culturable virus and PCR-virus. Many samples and wells that were positive for an indicator were, in fact, negative for virus (i.e., the indicators had a high rate of false positives). Positive predictive values at the well level were higher (but still mostly <50%) than at the sample level. Approximately 30 to 40% of the wells that were positive for an indicator were contaminated with virus at some point during the sampling period.
In contrast, specificities of the indicators were high, with values often near 90% or greater for culturable virus and PCR-virus. For example, *E. coli* had 95 and 97% specificity at the sample level for culturable virus and PCR-virus, respectively. In other words, a negative *E. coli* result correctly identified 95–97% of the samples that were negative for viruses. Specificity values at the well level were consistently lower than at the sample level for all six indicators, but the magnitude of the differences were usually not large.

Negative predictive values for the indicators were generally greater than 90%, with the exception of PCR-viruses at the well level, where estimates ranged from 50 to 76%. An indicator with a high negative predictive value means that when a sample or well is indicator-negative, it is likely to be also virus-negative (i.e. the indicator has a low rate of false negatives).

The sample and well data were reanalyzed according to the TCR, Hydrogeology, and GWR susceptibility subsets described in Table 5. Table 11 shows the ratio of the percent of virus-positive samples or wells in each subset to the percent of all virus positive samples or wells. Culturable viruses were twice as likely to be present in samples obtained from wells in each subset as in samples from all wells. Viruses measured this way were only 1.3–1.5 times more likely in wells from the TCR and hydrogeology subsets, but 4 times more likely in the GWR subset. PCR virus was poorly associated with the different subsets (ratios of 0.9–1.6).

Table 12 shows the indicator test performance measures for those combinations of indicators and susceptibility categories that had statistically significant indicator-virus associations. The chance of detecting virus was elevated when total coliform or enterococci bacteria or F-specific bacteriophage were detected in a hydrogeological susceptibility setting and when enterococci or somatic bacteriophage were detected for wells considered susceptible by having a TCR violation. No single indicator improved the chance of detecting virus for those wells considered susceptible by the GWR. Compared to the analyses that included all samples or wells, sensitivities and positive predictive values generally were higher for indicators measured in wells in all three susceptibility categories for both culturable virus and PCR-virus.

**Discussion**

This study addresses the relationship between microbial indicators and human enteric virus in groundwater. The focus of the study is on public systems in the US, Canada, Europe, and Asia for which raw data from 746 wells and 2273 samples were available. One or more indicator was found in 44% of the wells, while culturable virus was detected in 8% and PCR virus in 30% of the wells.

Many studies have examined correlations between concentrations of indicators and viruses and often times the correlations are weak or non-existent (see references in Table 1). When 12 studies were combined, the correlations between indicator and virus concentrations were statistically significant, but the low rho values showed the correlations were weak. Contributing factors to the weak relationships are the use of different methodologies, the amalgamation of wells from various hydrogeological settings, and the large number of...
samples or wells in the dataset that were negative for indicators or viruses. Excluding most negative values from the Spearman rank order test did increase the rho values pertaining to culturable virus, but as negative values cannot be excluded a priori such a relationship has no practical value for well testing.

Hynds et al. (2014) pooled data from 39 studies of private groundwater wells that included both microbial indicator and pathogen data and compared the percentage of indicator positive wells to the percentage of pathogen positive wells. The correlations tended to be weak and some were opposite of what would be expected, for example, viruses in groundwater were negatively correlated with enterococci. These authors recommend direct testing for virus. From 540 sets of indicator-pathogen correlations in studies of surface water, Wu et al. (2011) modeled the probability of finding a significant correlation and concluded that many studies do not find a correlation because of small sample size. Payment and Locas (2011) compared virus concentrations to concentrations of six indicators in 242 samples from 25 groundwater sites and found no correlations.

Logistic regression was used as an alternative approach to examine the association between dichotomous representations of indicator and virus detections. This approach allowed the calculation of risk ratios (i.e. the ratio of virus detection probabilities in the presence versus the absence of an indicator) while adjusting the association for underlying differences among the 12 studies and among wells within a study. Risk ratios and tests for association evaluate the overall correspondence between indicator and virus occurrence. A more complete interpretation of an indicator’s value with respect to viruses is provided by calculating standard test performance measures (e.g., sensitivity, specificity, etc.) that convey the likelihood of false positives and false negatives (Borchardt et al. 2003).

The risk ratios show that, in general, culturable viruses are more likely to be detected when there is a positive indicator result, especially for the indicators E. coli, enterococci and somatic phage at the sample level and F-specific and somatic phage at the well level. For example, the data indicate that it is nine times more likely to find culturable virus in a sample or well that also has somatic coliphage. In contrast, no significant risk ratios were found among indicators and PCR-virus at the sample level and the strongest risk ratios at the well level were less than two. Similarly, in examining the factors associated with statistically significant correlations between indicators and pathogens in surface waters, Wu et al. (2011) showed that studies that use molecular methods for pathogen detection were less likely to find significant correlations.

All six indicators that were evaluated tended to have low sensitivity and low positive predictive values, but high specificity and high negative predictive values. From a practical standpoint, this means that if a well has an unknown virus contamination problem, it is unlikely to be identified by a positive indicator (low sensitivity). Additionally, a positive indicator result does not necessarily mean the well is virus-contaminated; it could likely be a false positive (low positive predictive value). If the well does not have a virus contamination problem, there is a reasonable chance the indicator results will be negative, confirming there is no problem (high specificity). And a well that is indicator negative is unlikely to be virus contaminated, especially for culturable viruses (high negative predictive value). In summary,
the downside of the indicators is that many virus positive wells can be missed and there
could be many false positives. On the other hand, wells that are indicator-negative are
unlikely to have virus contamination problems. Moreover, sensitivities and positive
predictive values at the well level were higher for all six indicators compared to the sample
level, for the most part without sacrificing a large drop in specificity or negative predictive
value. This suggests that multiple samples from a well improves indicator performance in
assessing a well’s susceptibility to virus contamination. Indicator performance might also be
improved by considering not just the detection of the indicator, but also its concentration.
Payment and Locas (2011) showed the probability of human virus detection in the Saint-
Lawrence River in Quebec, Canada, increased with increasing concentrations of the
indicators thermotolerant coliforms or Clostridia perfringens. An indicator’s sensitivity
performance measure may depend on the indicator concentration with higher sensitivities at
higher concentrations.

The indicator-virus associations were evaluated for three well susceptibility categories to
examine if indicators were more informative based on prior knowledge of well-specific data
(Table 5). The TCR category was chosen because total coliforms are the most common
indicator used to evaluate well susceptibility to pathogens. The hydrogeology category was
selected due to the known susceptibility of karst, fractured bedrock, and gravel/cobble
settings, and the GWR category was selected because the criteria for well susceptibility
includes two indicators that typically had not been tested in groundwater, enterococci and
coliphage. Among the wells in the three susceptibility categories, culturable virus was
detected twice as frequently in samples and 1.3–3.9 times more in wells compared to all
samples or wells (Table 11). In contrast, PCR-viruses were detected only 0.7–1.6 times more
in samples or wells in the susceptibility categories. This difference may be a function of
virus fate in the subsurface. It may be easier to detect viruses by PCR in less susceptible
settings as the viruses can still be detected long after infectivity is lost and at greater
distances from sources of contamination (Ogorzaly et al. 2010).

Sensitivities, positive predictive values, and risk ratios tended to be higher for total
coliforms, enterococci, F-specific phage, and somatic phage when they were measured in
samples or wells in the TCR or hydrogeology susceptibility categories. In other words, for
wells with two or more TCR violations or wells located in susceptible hydrogeological
settings, like karst, these indicators performed better at identifying samples and wells that
were true virus-positives and with a lower false positive rate. However, for wells that would
have met the criteria under the GWR for additional monitoring, none of the single indicators
performed any better than when they were measured in all wells regardless of susceptibility
status.

In interpreting the data presented in the preceding, it needs to be stressed that there is a
degree of uncertainty in the placement of wells into the susceptibility categories, particularly
the hydrogeology category. Interpretation is weakened by the lack of uniformity in the types
of stressors measured by individual studies as well as by the different methodologies used.
Future studies that examine indicator and virus relationships should measure viruses by both
culture and molecular assays, and include at least total coliform, E. coli, enterococci, aerobic
spores, F-specific and somatic coliphages. Standard methods should be used as much as possible.

The finding of a virus by either culturable or molecular assays is not always easily translatable to the public health risk presented by a particular well. Virus concentration in groundwater can vary rapidly over time (Bradbury et al. 2013). A single data point can fall anywhere within the possible range of virus concentrations during transient contamination, from having no virus, and thus underestimating risk, to by chance collecting the sample during peak concentration and overestimating risk. In addition, as virus recovery from water by various concentration procedures rarely achieves 100% (Cashdollar and Wymer 2013; Gibbons et al. 2010; Ikner et al. 2012, 2011; Karim et al. 2009; Sobsey and Glass 1984; Wu et al. 2013) virus concentration, and therefore risk, can be underestimated. Different virus groups and even different members of the same species can have different recovery efficiencies, while recovery efficiency of a single virus type will vary over time with changes in turbidity and other water quality factors.

To pose a health risk, virus concentrations in the well water must be high enough that individuals drinking the water ingest more virus than the minimum infectious dose necessary to initiate infection. Apart from egregious virus contamination of groundwater that results in an outbreak (e.g. Borchardt et al. 2011), most surveys of groundwater in developed countries show culturable virus concentrations that typically are very low (e.g., combined data from this study), suggesting the health risk is low. On the other hand, culturable virus assays can underestimate virus concentrations and occurrence. Virus in water can be aggregated with each aggregate registering as a single infectious unit in culturable virus assays (Gassilloud and Gantzer 2005); however, aggregates may have a sufficient number of virus particles to cause an infection, if consumed. Culturable assays also underestimate risk in that they do not detect many important waterborne enteric viruses, and cell culture does not necessarily replicate the same favorable conditions for virus replication as when a virus infects a host.

Because PCR detects virus genomes and not necessarily infectious virions, PCR measurements of viruses in ground-water are often thought to overestimate the health risk. On the other hand, virus present can maintain infectivity for months at temperature of 12 °C or less (Charles et al. 2009; Nasser and Oman 1999). PCR also may underestimate risk because enteric viruses, and especially RNA viruses, evolve constantly, leading to genetic variability that prevents their detection by PCR (i.e., false negative results). Nonetheless, the presence of viral genomes in groundwater shows that an aquifer is susceptible to fecal contamination. Importantly, Borchardt et al. (2012) compared the fraction of acute gastrointestinal illness (AGI) associated with drinking untreated groundwater in 14 communities with PCR-virus from community wells. The concentration of enteroviruses in the tap water of households in the communities was associated with AGI in adults, and tap water concentrations of norovirus genogroup I were associated with AGI in all ages. Using quantitative microbial risk assessment, the authors estimate that between 6 and 22% of the overall AGI in the communities and up to 63% of AGI in children less than 5 years of age was associated with drinking untreated tap water. The AGI incidence increased by 63% when the mean norovirus concentration in the communities’ tap water increased from 0 to about 3 genomic copies per liter. Thus, without results to the contrary, the possibility of
adverse health risks based upon PCR positive findings should be considered when there are septic tanks, sewer lines, or other sources of contamination near aquifers that reside in similar hydrogeologic settings to those in Wisconsin.

Conclusions

A meta-analysis of 12 groundwater studies was conducted. Overall, one or more indicators were present in 37% of samples and 44% of wells, culturable virus in 4% of samples and 8% of wells, and PCR-virus in 14% of samples and 30% of wells. All six indicators examined in this study were associated with culturable virus at one or both levels of analysis (sample or well). None of the indicators were associated with PCR-measured viruses at the sample level, and only total coliforms, E. coli, and somatic phage were associated with PCR-viruses at the well level. Judging by the risk ratios, the best indicator for culturable virus and PCR-virus in groundwater was somatic phage. However, all the indicators tended to have low sensitivities and positive predictive values, but high specificity and negative predictive values, which means groundwater that tests negative for the indicators is unlikely to be virus contaminated, but positive indicator tests do not necessarily mean there is virus contamination. That the sensitivities and positive predictive values of the indicators were higher at the well-level of analysis matches intuition: evaluating well susceptibility to virus contamination is improved by sampling for indicators multiple times from the well. In combining the 12 studies, it was learned that the statistical models had to account for underlying differences among the studies and wells, and furthermore, the indicator-virus associations changed when the analysis was restricted to wells classified as susceptible by the total coliform rule or hydrogeology. This suggests the strength of associations between indicators and viruses are specific to hydrogeological setting. Thus, for optimal management of groundwater sanitary quality, the indicator associations and test performance measures, as reported in this study, should be characterized for the specific region under management. This would give groundwater managers the greatest confidence in interpreting indicator test results for their region. This study focused upon the relationships of single microbial indicators to virus occurrence in groundwater samples and wells. Future studies should examine whether multiple microbial indicators would provide stronger correlation to virus presence.

Acknowledgements

Mohammad R. Karim was supported through an appointment to the Research Participation Program at the US Environmental Protection Agency (EPA), National Exposure Research Laboratory, administered by the Oak Ridge Institute for Science and Education through an Interagency Agreement between the US Department of Energy and the EPA. The authors thank Drs. Hiroyuki Katayama, Keisuka Kuroda, Soon-Young Paik, and Pierre Payment for supplying raw study data, and Donna Francy, Bruce Lindsey, and Richard Lieberman for supplying site identification data. The authors also acknowledge the many authors who have investigated and reported on factors affecting virus and indicator occurrence in groundwater and on the virus occurrence studies mentioned in this paper. Without their efforts this paper would not be possible.

References

Abbaszadegan M, Lechevallier M, Gerba C (2003) Occurrence of viruses in US groundwaters. J Am Water Works Assoc 95(9):107–120
Ashbolt NJ (2015) Microbial contamination of drinking water and human health from community water systems. Curr Environ Health Rep 2(1):95–106. doi:10.1007/s40572-014-0037-5 [PubMed: 25821716]

Ashbolt NJ, Grabow WOK, Snozzi M (2001) Indicators of microbial water quality In: Fewtrell L, Bartram J (eds) Water quality: guidelines, standards and health. World Health Organization, IWA, London, pp 289–316

Beer KD, Gargano JW, Roberts VA, Hill VR, Garrison LE, Kutty PK, Hilborn ED, Wade TJ, Fullerton KE, Yoder JS (2015) Surveillance for waterborne disease outbreaks associated with drinking water: United States, 2011–2012. MMWR Morb Mortal Wkly Rep 64(31):842–848 [PubMed: 26270059]

Berg G, Dahling DR, Brown GA, Berman D (1978) Validity of fecal coliforms, total coliforms, and fecal streptococci as indicators of viruses in chlorinated primary sewage effluents. Appl Environ Microbiol 36(6):880–884 [PubMed: 104657]

Berger P (2008) Viruses in ground water In: Hlavinek J, Mahrikova I (eds) Dangerous pollutants (xenobiotics) in urban water cycle. Springer, Dordrecht, The Netherlands, pp 131–149

Blaschke AP, Derx J, Zessner M, Kirnbauer R, Kavka G, Strelec H, Farneitner AH, Pang L (2016) Setback distances between small biological wastewater treatment systems and drinking water wells against virus contamination in alluvial aquifers. Sci Total Environ 573:278–289. doi:10.1016/j.scitotenv.2016.08.075 [PubMed: 27570196]

Borchardt MA, Bertz PD, Spencer SK, Battigelli DA (2003) Incidence of enteric viruses in groundwater from household wells in Wisconsin. Appl Environ Microbiol 69(2):1172–1180 [PubMed: 12571044]

Borchardt MA, Bradbury KR, Alexander EC, Jr, Kolberg RJ, Alexander SC, Archer JR, Braatz LA, Forest BM, Green JA, Spencer SK (2011) Norovirus outbreak caused by a new septic system in a dolomite aquifer. Ground Water 49(1):85–97. doi:10.1111/j.1745-6584.2010.00686.x [PubMed: 20199588]

Borchardt MA, Spencer SK, Kieke BA, Lambertini E, Loge FJ (2012) Viruses in nondisinfected drinking water from municipal wells and community incidence of acute gastrointestinal illness. Environ Health Perspect 120(9):1272–1279. doi:10.1289/ehp.1104499 [PubMed: 22659405]

Bradbury KR, Borchardt MA, Gotkowitz M, Spencer SK, Zhu J, Hunt RJ (2013) Source and transport of human enteric viruses in deep municipal water supply wells. Environ Sci Technol 47(9):4096–4103. doi:10.1021/es400509b [PubMed: 23570447]

Cashdollar JL, Wymer L (2013) Methods for primary concentration of viruses from water samples: a review and meta-analysis of recent studies. J Appl Microbiol 115(1):1–11. doi:10.1111/jam.12143 [PubMed: 23360578]

Charles KJ, Shore J, Sellwood J, Laverick M, Hart A, Pedley S (2009) Assessment of the stability of human viruses and coliphage in groundwater by PCR and infectivity methods. J Appl Microbiol 106(6):1827–1837. doi:10.1111/j.1365-2672.2009.04150.x [PubMed: 19298517]

Chavance M, Escolano S (2016) Misspecification of the covariance structure in generalized linear mixed models. Stat Methods Med Res 25(2):630–643. doi:10.1177/0962280214526859 [PubMed: 23070599]

Chilton J, Seiler K-P (2006) Groundwater occurrence and hydrogeological environments In: Schmoll O, Howard G, Chilton J, Chorus I (eds) Protecting groundwater for health managing the quality of Drinking-water systems. World Health Organization, Geneva, pp 21–47

Cho HG, Lee SG, Kim WH, Lee JS, Park PH, Cheon DS, Jheong WH, Jho EH, Lee JB, Paik SY (2014) Acute gastroenteritis outbreaks associated with ground-waterborne norovirus in South Korea during 2008–2012. Epidemiol Infect 142(12):2604–2609. doi:10.1017/S0950268814000247 [PubMed: 24534556]

Costan-Longares A, Montemayor M, Payan A, Mendez J, Jofre J, Mujeriego R, Lucena F (2008) Microbial indicators and pathogens: removal, relationships and predictive capabilities in water reclamation facilities. Water Res 42(17):4439–4448. doi:10.1016/j.watres.2008.07.037 [PubMed: 18762313]

Craun GF, Brunkaed JM, Yoder JS, Roberts VA, Carpenter J, Wade T, Calderon RL, Roberts JM, Beach MJ, Roy SL (2010) Causes of outbreaks associated with drinking water in the United States

Hydrogeol J. Author manuscript; available in PMC 2018 September 19.
from 1971 to 2006. Clin Microbiol Rev 23(3):507–528. doi:10.1128/CMR.00077-09 [PubMed: 20610821]

Dahling DR (2002) An improved filter elution and cell culture assay procedure for evaluating public groundwater systems for culturable enteroviruses. Water Environ Res 74(6):564–568 [PubMed: 12540097]

Davis JV, Witt EC, III (1998) Microbiological quality of public-water supplies in the Ozark plateaus aquifer system. Missouri. US Geol Surv Fact Sheet 028–98

Davis JV, Witt EC, III (2000) Microbiological and chemical quality of ground water used as a source of public supply in southern Missouri: phase I, May 1997–March 1998. US Geol Surv Water Resour Invest Rep 00–4038

Deborde DC, Woessner WW, Lauerman B, Ball P (1998a) Coliphage prevalence in high school septic effluent and associated ground water. Water Res 32(12):3781–3785. doi:10.1016/S0043-1354(98)00171-7

Deborde DC, Woessner WW, Lauerman B, Ball PN (1998b) Virus occurrence and transport in a school septic system and unconfined aquifer. Ground Water 36(5):825–834. doi:10.1111/j.1745-6584.1998.tb02201.x

Edberg SC, Rice EW, Karlin RJ, Allen MJ (2000) Escherichia coli: the best biological drinking water indicator for public health protection. Symp Ser Soc Appl Microbiol 29:106S–116S

Femmer SR (2000) Microbiological and chemical quality of ground water used as a source of public supply in southern Missouri-phase II, April–July, 1998. US Geol Surv Water Resour Invest Rep 00–4260

Ford DC (1997) Principal features of evaporite karst in Canada. Carbonate Evaporite 12(1):15–23

Fout GS, Martinson BC, Moyer MW, Dahling DR (2003) A multiplex reverse transcription-PCR method for detection of human enteric viruses in groundwater. Appl Environ Microbiol 69(6):3158–3164. doi:10.1128/AEM.69.6.3158-3164.2003 [PubMed: 12788711]

Francy DS, Bushon RN, Stopar J, Luzano EJ, Fout GS (2004a) Environmental factors and chemical and microbiological water-quality constituents related to the presence of enteric viruses in ground water from small public water supplies in southeastern Michigan. US Geological Survey, Reston, VA, pp 1–54

Francy DS, Bushon RN, Stopar J, Luzano EJ, Fout GS (2004b) Environmental factors and chemical and microbiological water-quality constituents related to the presence of enteric viruses in ground water from small public water supplies in southeastern Michigan US Geol Surv Sci Invest Rep 2004–5219, pp 1–54

Gabrieli R, Maccari F, Ruta A, Pana A, Divizia M (2009) Norovirus detection in groundwater. Food Environ Virol 1(2):92–96. doi:10.1007/s12560-009-9014-9

Gassilloud B, Gantzer C (2005) Adhesion-aggregation and inactivation of poliovirus 1 in groundwater stored in a hydrophobic container. Appl Environ Microbiol 71(2):912–920. doi:10.1128/AEM.71.2.912-920.2005 [PubMed: 15691948]

Gibbons CD, Rodriguez RA, Tallon L, Sobsey MD (2010) Evaluation of positively charged alumina nanofibre cartridge filters for the primary concentration of noroviruses, adenoviruses and male-specific coliphages from seawater. J Appl Microbiol 109(2):635–641. doi:10.1111/j.1365-2672.2010.04691.x [PubMed: 20202019]

Gotkowitz MB, Bradbury KR, Borchardt MA, Zhu J, Spencer SK (2016) Effects of climate and sewer condition on virus transport to groundwater. Environ Sci Technol 50(16):8497–8504. doi:10.1021/acs.est.6b01422 [PubMed: 27434550]

Guzman-Herrador B, Carlander A, Ethelberg S, Freiesleben de Blasio B, Kuusi M, Lund V, Lofsdahl M, MacDonald E, Nichols G, Schonning C, Sudre B, Tronberg L, Vold L, Semenza JC, Nygard K (2015) Waterborne outbreaks in the Nordic countries, 1998 to 2012. Euro Surveill 20(24). doi:10.1515/2015111501

Headd B, Bradford SA (2016) Use of aerobic spores as a surrogate for cryptosporidium oocysts in drinking water supplies. Water Res 90:185–202. doi:10.1016/j.watres.2015.12.024 [PubMed: 26734779]

Health Canada (2017) Guidelines for Canadian drinking water quality: summary table. www.hc-sc.gc.ca/ewh-smt/ pubs/water-eau/sum_guide-res_recom/index-eng.php Accessed January 2017
Hejkal TW, Keswick B, LaBelle RL, Gerba CP, Sanchez Y, Dreesman G, Hafkin B, Melnick JL (1982) Viruses in a community water supply associated with an outbreak of gastroenteritis and infectious hepatitis. J AWWA 84(6):318–321

Hilborn ED, Wade TJ, Hicks L, Garrison L, Carpenter J, Adam E, Mull B, Yoder J, Roberts V, Gargano JW (2013) Surveillance for waterborne disease outbreaks associated with drinking water and other nonrecreational water: United States, 2009–2010. Morb Mortal Wkly Rep 62(35):714–720

Horick PJ (1984) Silurian-Devonian aquifer of Iowa Iowa Geological Survey, University of Iowa, Iowa City, IA

Howard G, Bartram J, Pedley S, Schmoll O, Chorus I, Berger P (2006) Groundwater and public health In: Schmoll O, Howard G, Chilton J, Chorus I (eds) Protecting groundwater for health: managing the quality of drinking-water sources. IWA, London, pp 3–19

Hruby CE, Libra RD, Fields CL, Kolpin DW, Hubbard LE, Borchardt MR, Spencer SK, Wichman MD, Hall N, Schueller MD, Furlong ET, Weyer PJ (2015) 2013 Survey of Iowa groundwater and evaluation of public well vulnerability classifications for contaminants of emerging concern. Iowa Geological and Water Survey Technical Information Series 57, Iowa Department of Natural Resources, Des Moines, IA

Hynds PD, Thomas MK, Pintar KD (2014) Contamination of groundwater systems in the US and Canada by enteric pathogens, 1990–2013: a review and pooled-analysis. PLoS One 9(5):e93301. doi:10.1371/journal.pone.0093301 [PubMed: 24806545]

Ikner LA, Soto-Beltran M, Bright KR (2011) New method using a positively charged microporous filter and ultrafiltration for concentration of viruses from tap water. Appl Environ Microbiol 77(10):3500–3506. doi:10.1128/AEM.02705-10 [PubMed: 21441329]

Ikner LA, Gerba CP, Bright KR (2012) Concentration and recovery of viruses from water: a comprehensive review. Food Environ Virol 4(2):41–67. doi:10.1007/s12560-012-0980-2 [PubMed: 23412811]

Jack S, Bell D, Hewitt J (2013) Norovirus contamination of a drinking water supply at a hotel resort. NZ Med J 126(1387):98–107

Jofre J, Bosch A, Lucena F, Girones R, Tartera C (1986) Evaluation of Bacteroides fragilis bacteriophages as indicators of the virological quality of water. Water Sci Technol 18(10):167–173

Johnson TB (2005) Detection of enteric viruses in east Tennessee public ground water systems. MSc Thesis, University of Tennessee, Knoxville, TN

Johnson TB, McKay LD, Layton AC, Jones SW, Johnson GC, Cashdollar JL, Dahlinger DR, Villegas LF, Fout GS, Williams DE, Sayler G (2011) Viruses and bacteria in karst and fractured rock aquifers in East Tennessee, USA. Ground Water 49(1):98–110. doi:10.1111/j.1745-6584.2010.00698.x [PubMed: 20331750]

Jung JH, Yoo CH, Koo ES, Kim HM, Na Y, Jheong WH, Jeong YS (2011) Occurrence of norovirus and other enteric viruses in untreated groundwaters of Korea. J Water Health 9(3):544–555. doi:10.2166/wh.2011.142 [PubMed: 21976201]

Karim MR, LeChevallier MW, Abbaspazegan M, Alum A, Sobrinho J, Rosen J (2004) Field testing of USEPA methods 1601 and 1602 for coliphage testing in groundwater. AWWA Research Foundation, Denver

Karim MR, Rhodes ER, Brinkman N, Wymer L, Fout GS (2009) New electropositive filter for concentrating enteroviruses and noroviruses from large volumes of water. Appl Environ Microbiol 75(8):2393–2399. doi:10.1128/AEM.00922-08 [PubMed: 19218410]

Katayama H (2008) Detection of microbial contamination in groundwater In: Takizawa S (ed) Groundwater management in Asian cities: technology and policy for sustainability. Springer, Tokyo, pp 151–169

Kitajima M, Gerba CP (2015) Aichi virus 1: environmental occurrence and behavior. Pathogens 4(2):256–268. doi:10.3390/pathogens4020256 [PubMed: 25996404]

Kuroda K, Nakada N, Hanamoto S, Inaba M, Katayama H, Do AT, Nga TT, Oguma K, Hayashi T, Takizawa S (2015) Pepper mild mottle virus as an indicator and a tracer of fecal pollution in water environments: comparative evaluation with wastewater-tracer pharmaceuticals in Hanoi, Vietnam. Sci Total Environ 506–507:287–298. doi:10.1016/j.scitotenv.2014.11.021

Hydrogeol J. Author manuscript; available in PMC 2018 September 19.
Lambertini E, Spencer SK, Kieke BA, Jr, Loge FJ, Borchardt MA (2011) Virus contamination from operation and maintenance events in small drinking water distribution systems. J Water Health 9(4):799–812. doi:10.2166/wh.2011.018 [PubMed: 22048438]

Lambertini E, Borchardt MA, Kieke BA, Jr, Spencer SK, Loge FJ (2012) Risk of viral acute gastrointestinal illness from nondisinfected drinking water distribution systems. Environ Sci Technol 46(17):9299–9307. doi:10.1021/es301592t [PubMed: 22839570]

Lee GC, Jeong WH, Kim MJ, Choi DH, Baik KH (2013) A 5-year survey (2007–2011) of enteric viruses in Korean aquatic environments and the use of coliforms as viral indicators. Microbiol Immunol 57(1):46–53. doi:10.1111/j.1348-0421.2012.00515.x [PubMed: 23046241]

Lee H, Kim M, Lee JE, Lim M, Kim M, Kim JM, Jeong WH, Kim J, Ko G (2011) Investigation of norovirus occurrence in groundwater in metropolitan Seoul, Korea. Sci Total Environ 409(11): 2078–2084. doi:10.1016/j.scitotenv.2011.01.059 [PubMed: 21440930]

Lee J-Y, Kwon K (2016) Current status of groundwater monitoring networks in Korea. Water 8(4):168. doi:10.3390/w8040168

Lee SG, Jeong WH, Suh CI, Kim SH, Lee JB, Jeong YS, Ko G, Jang KL, Lee GC, Paik SY (2011) Nationwide groundwater surveillance of noroviruses in South Korea, 2008. Appl Environ Microbiol 77(4):1466–1474. doi:10.1128/AEM.01996-10 [PubMed: 21183642]

Lieberman RJ, Shadix LC, Newport BS, Frebis CP, Moyer MWN, Safferman RS, Stetler RE, Lye D, Fout GS, Dahlberg DR (2002) Microbial monitoring of vulnerable public groundwater supplies. American Water Works Association, Denver, CO

Lindsey BD, Rasberry JS, Zimmerman TM (2002) Microbiological quality of water from noncommunity supply wells in carbonate and crystalline aquifers of Pennsylvania. US Geol Surv Water Resour Invest Rep 01-426

Locas A, Barthe C, Barbeau B, Carriere A, Payment P (2007) Virus occurrence in municipal groundwater sources in Quebec, Canada. Can J Microbiol 53(6):688–694. doi:10.1139/w07-034 [PubMed: 17668028]

Locas A, Barthe C, Margolin AB, Payment P (2008) Groundwater microbiological quality in Canadian drinking water municipal wells. Can J Microbiol 54(6):472–478. doi:10.1139/w08-028 [PubMed: 18535633]

Lucena F, Ribas F, Duran AE, Skraber S, Gantzer C, Campos C, Moron A, Calderon E, Jofre J (2006) Occurrence of bacterial indicators and bacteriophages infecting enteric bacteria in groundwater in different geographical areas. J Appl Microbiol 101(1):96–102. doi:10.1111/j.1365-2672.2006.02907.x [PubMed: 16834595]

McKay LD (2011) Foreword: pathogens and fecal indicators in groundwater. Ground Water 49(1):1–3. doi:10.1111/j.1745-6584.2010.00763.x [PubMed: 21039447]

McMinn BR, Korajkic A, Ashbolt NJ (2014) Evaluation of Bacteroides fragilis GB-124 bacteriophages as novel human-associated faecal indicators in the United States. Lett Appl Microbiol. doi:10.1111/lam.12252

Meschke JS (2001) Comparative adsorption, persistence, and mobility of Norwalk virus, poliovirus type 1, and F+RNA coliphages in soil and groundwater. PhD. Thesis, University of North Carolina, Chapel Hill, NC

Morel JG, Bokossa MC, Neerchal NK (2003) Small sample correction for the variance of GEE estimators. Biom J 45(4):395–409. doi:10.1002/bimj.200390021

Nasser AM, Oman SD (1999) Quantitative assessment of the inactivation of pathogenic and indicator viruses in natural water sources. Water Res 33(7):1748–1752. doi:10.1016/S0043-1354(98)00380-7

Nasser AM, Tchorch Y, Fattal B (1993) Comparative survival of E. coli, F+ bacteriophages, HAV and poliovirus 1 in wastewater and groundwater. Water Sci Technol 27(3–4):401–407

National Health and Medical Research Council (2016) Australian drinking water guidelines 6, version 3.2 NHMRC, Canberra, Australia

Ogorzaly L, Bertrand I, Paris M, Maul A, Gantzer C (2010) Occurrence, survival, and persistence of human adenoviruses and F-specific RNA phages in raw groundwater. Appl Environ Microbiol 76(24):8019–8025. doi:10.1128/AEM.00917-10 [PubMed: 20952644]

Hydrogeol J. Author manuscript; available in PMC 2018 September 19.
Pang L, Close M, Goltz M, Noonan M, Sinton L (2005) Filtration and transport of Bacillus subtilis spores and the F-RNA phage MS2 in a coarse alluvial gravel aquifer: implications in the estimation of setback distances. J Contam Hydrol 77(3):165–194 [PubMed: 15763354]

Payment P, Locas A (2005) Évaluation et contrôle de la qualité virologique des eaux souterraines [Evaluation and monitoring of the virological quality of groundwater]. INRS-Institut Armand-Frappier, Quebec

Payment P, Locas A (2011) Pathogens in water: value and limits of correlation with microbial indicators. Ground Water 49(1):4–11. doi:10.1111/j.1745-6584.2010.00710.x [PubMed: 20477877]

Pedley S, Yates M, Schijven JF, West J, Howard G, Barrett M (2006) Pathogens: health relevance, transport and attenuation In: Schmoll O, Howard G, Chilton J, Chorus I (eds) Protecting groundwater for health managing the quality of drinking-water systems. IWA, London, pp 49–80

Sobsey MD, Glass JS (1984) Influence of water quality on enteric virus concentration by microporous filter methods. Appl Environ Microbiol 47(5):956–960 [PubMed: 6331310]

Tallon P, Magajna B, Lofranco C, Leung KT (2005) Microbial indicators of faecal contamination in water: a current perspective. Water Air Soil Pollut 166(1–4):139–166. doi:10.1007/s11270-005-7905-4

University of Auckland (2008) Karst. http://web.env.auckland.ac.nz/our_research/karst/ Accessed August 2016

USEPA (2006a) 40 CFR parts 9, 141, and 142 National Primary Drinking Water Regulations: ground water rule; final rule. Fed Reg 71(216):65574–65660

USEPA (2006b) Occurrence and monitoring document for the final ground water rule: 1–1 to A-2. USEPA, Washington, DC

Wakayama H (2016) Revision of drinking water quality standards in Japan. Office of Drinking Water Quality Management http://www.nilim.go.jp/lab/bcg/siryou/tnn/tnn0264pdf/ks0264011.pdf Accessed January 2017

Wallender EK, Ailes EC, Yoder JS, Roberts VA, Brunkard JM (2014) Contributing factors to disease outbreaks associated with untreated groundwater. Ground Water 52(6):886–897. doi:10.1111/gwat.12121 [PubMed: 24116713]

Weary DJ, Doctor DH (2014) Karst in the United States: a digital map compilation and database. US Geological Survey https://pubs.usgs.gov/of/2014/1156/ Accessed 2016

Wentworth CK (1922) A scale of grade and class terms for clastic sediments. J Geol 30(5):377–392

WHO (2011) Guidelines for drinking-water quality, 4th edn World Health Organization, Geneva

Won J-H, Kim J-W, Koh G-W, Lee J-Y (2005) Evaluation of hydrogeological characteristics in Jeju Island, Korea. Geosci J 9(1):33–46. doi:10.1007/bf02910552

Wu J, Long SC, Das D, Dorner SM (2011) Are microbial indicators and pathogens correlated? A statistical analysis of 40 years of research. J Water Health 9(2):265–278. doi:10.2166/wh.2011.117 [PubMed: 21942192]

Wu J, Simmons OD, 3rd, Sobsey MD (2013) Uncertainty analysis of the recovery of hollow-fiber ultrafiltration for multiple microbe classes from water: a Bayesian approach. J Microbiol Methods 93(3):161–167. doi:10.1016/j.mimet.2013.03.005 [PubMed: 23524155]

Zhou X, Li H, Sun L, Mo Y, Chen S, Wu X, Liang J, Zheng H, Ke C, Varma JK, Klena JD, Chen Q, Zou L, Yang X (2012) Epidemiological and molecular analysis of a waterborne outbreak of norovirus GI.4. Epidemiol Infect 140(12):2282–2289. doi:10.1017/S0950268812000374 [PubMed: 22400795]
Table 1
Microbial indicator and virus occurrence sources used in this study

| Study name                                      | Number of drinking water systems | Number of wells | Number of samples | Study date range (month/year) | Reference |
|-------------------------------------------------|----------------------------------|-----------------|-------------------|-------------------------------|-----------|
| USEPA/AWWARF                                    | 29                               | 30              | 333               | 9/92–12/94                    | Dahling 2002; Fout et al. 2003; Lieberman et al. 2002 |
| USGS (Missouri)                                 | 180                              | 182             | 322               | 5/97–7/98                     | Davis and Witt 2000; Femmer 2000 |
| USGS (Ohio)/USEPA                               | 38                               | 38              | 169               | 6/99–7/01                     | Francy et al. 2004 |
| USGS (Pennsylvania)                             | 60                               | 60              | 60                | 9/00–2/01                     | Lindsey et al. 2002 |
| American Water Service Company                  | 20                               | 20              | 235               | 3/01–6/02                     | Karim et al. 2004 |
| University of Tennessee (USA)                   | 4                                | 4               | 6                 | 3/04–8/04                     | Johnson et al. 2011 |
| Institut Armand-Frappier (Canada)               | 36                               | 36              | 243               | 3/04–12/12                    | Locas et al. 2007; Locas et al. 2008; Payment and Locas 2005; P. Payment, Institut Armand-Frappier, 2015, personal communication |
| University of Rome Tor Vergata (Italy)          | 8                                | 8               | 14                | 6/05–12/05                    | Gabrieli et al. 2009 |
| University of Tokyo (Tokyo, Japan)              | 46                               | 46              | 46                | 11/05–1/06                    | Katayama 2008; H. Katayama, University of Tokyo, 2015, personal communication |
| Marshfield Clinic Research Foundation (USA)     | 14                               | 36              | 391               | 4/06–11/07                    | Borchardt et al. 2012 |
| NIER (South Korea)                              | 220                              | 220             | 383               | 7/07–12/08                    | Jung et al. 2011; Lee et al. 2013; S. Lee et al. 2011; Lee S et al. 2011; S-Y. Paik, The Catholic University, Seoul, 2015, personal communication |
| Iowa Department of Natural Resources (USA)      | 63                               | 66              | 71                | 3/13–6/13                     | Hruby et al. 2015 |
| Total                                           | 718                              | 746             | 2,273             | -                             | -         |

*All wells were from public drinking water systems, e.g., those used by multiple individuals rather than a single household. In the US, a public system is one with at least 15 service connections or which serves 25 persons for 60 days or more.*

*Hydrogeol J. Author manuscript; available in PMC 2018 September 19.*
### Table 2

Assays and data types for combined studies

| Study name                                      | TC  | EC  | Ent | Spores | F+   | SomPh | CulVir | PCRVir | Type |
|------------------------------------------------|-----|-----|-----|--------|------|-------|--------|--------|------|
| USEPA/AWWARF                                   | QN  | QN  | QN  | QN\(^b\) | QN   | QN    | QN     | QL     | Yes  |
| USGS (Missouri)                                | QN\(^c\) | QN  | QN  | ND    | QN   | QN    | QL\(^d\) | QL     | Yes\(^d\) |
| USGS (Ohio)/USEPA                              | QN  | QN  | QN  | ND    | QN   | QN    | QL     | QL     | Yes  |
| USGS (Pennsylvania)                            | QN  | QN  | QN  | ND    | QN   | QN    | QN     | ND     | ND   |
| American Water Service Company                 | QL  | QL  | QL  | QN\(^b\) | QN   | QN    | QL     | Yes    |      |
| University of Tennessee (USA)                  | QN  | QN  | ND  | ND    | ND   | QN    | QL     | Yes    |      |
| Institut Armand-Frappier (Canada)              | QN  | QN  | QN  | QN\(^e\) | QN   | QN    | QL     | Yes    |      |
| University of Rome Tor Vergata (Italy)         | QN  | QN  | QN  | ND    | ND   | ND    | QL     | Yes    |      |
| University of Tokyo (Tokyo, Japan)             | QN  | QN  | ND  | ND    | ND   | ND    | QL     | Yes    |      |
| Marshfield Clinic Research Foundation (USA)    | QL\(^f\) | QL  | ND  | ND    | ND   | ND    | QN     | Yes    |      |
| NIER (South Korea)                             | QL  | QL  | ND  | ND    | QN   | QN    | QL     | Yes    |      |
| Iowa Department of Natural Resources (USA)     | QN  | QN  | QN  | ND    | ND   | ND    | QN     | QN     | Yes  |

\(^a\)Abbreviations: TC total coliforms; EC *E. coli*; Ent enterococci; F+ F-specific phage; SomPh somatic phage; CulVir culturable virus; PCRVir PCR-virus; Type individual virus types identified by PCR; QN quantitative assays; QL qualitative assays (presence/absence); ND assay not performed or data not available

\(^b\)Anaerobic spores measured

\(^c\)Fecal rather than total coliforms measured

\(^d\)This study was performed in two phases with no molecular assays being done in the 2nd phase

\(^e\)Aerobic spores measured

\(^f\)Community TCR data were used to estimate the possible presence of total coliform and *E. coli* in community wells of this study. All wells in a community were assumed to be negative if a TCR sample collected within 0–2 days of the day of virus sampling was negative. Likewise, wells were considered positive if a TCR sample was positive during the same timeframe (i.e. 0–2 days). Wells were considered negative for *E. coli* if a total coliform test was negative and *E. coli* data not recorded.
### Table 3

Microbial indicator and culturable virus data conversion

| Indicator/virus   | No. volume options | Volume analyzed | Result | Value/100 ml Used |
|-------------------|--------------------|----------------|--------|-------------------|
| Bacterial indicators | 1                  | 100 ml         | -      | 0 CFU             |
|                   | 1                  | 100 ml         | +      | 5 CFU             |
|                   | 1                  | 100 ml         | TMTC   | 200 CFU           |
|                   | 2                  | 100 ml         | -      | 0 CFU             |
|                   |                    | 1000 ml        | -      |                   |
|                   | 2                  | 100 ml         | -      | 0.5 CFU           |
|                   |                    | 1000 ml        | +      |                   |
|                   | 2                  | 100 ml         | +      | 5 CFU             |
|                   |                    | 1000 ml        | +      |                   |
| Bacteriophage     | 1                  | 100 ml         | TMTC   | 400 PFU           |
|                   | 2                  | 100 ml         | -      | 0 PFU             |
|                   |                    | 1000 ml        | -      |                   |
|                   | 2                  | 100 ml         | -      | 0.5 PFU           |
|                   |                    | 1000 ml        | +      |                   |
|                   | 2                  | 100 ml         | + or ND| 5 PFU             |
|                   |                    | 1000 ml        | +      |                   |
| Culturable virus  | 1                  | 500 L          | -      | 0 MPN             |
| 1. (expressed as +/- per 500 L) | 1    | 500 L          | +      | 0.0002 MPN        |

*a* Qualitative data were converted to quantitative data as indicated so that Spearman correlations could be performed on the broader data set. The converted data were not used for any other statistical analysis.

*b* No. volume options = 1: a 100-ml sample volume was used for microbial indicator tests; No. volume options = 2: both 100-ml and 1,000-ml sample volumes were used.

*c* Quantitative data using 100-ml samples with too many colonies (bacteria) or plaques (bacteriophage) to count

*d* ND – not done
### Table 4

Molecular method qualitative data conversion

| Assay type               | Conversion                                                                                                                                 |
|--------------------------|-------------------------------------------------------------------------------------------------------------------------------------------|
| Integrated cell culture-PCR | Assuming at least one positive cell culture flask (i.e., 1 MPN per volume assayed), MPN/L is calculated by dividing 1 MPN by the equivalent volume of original water sample placed on replicate flasks. Genomic copies (GC)/100 ml is calculated by dividing the MPN/L value by 10 and multiplying by 20. Note: the value 20 is a conservative estimate of the physical to infectious particle ratio. |
| Conventional PCR         | Study-specific data were used to calculate the volume (in liters) of groundwater sample added to each PCR assay. This volume was calculated using the volume of groundwater from which virus was initially concentrated, the volumes and amount of any additional concentration steps, and the volume of final concentrate or extracted nucleic acid added to each PCR assay. GC/100 ml is calculated by dividing 1 by the volume per PCR assay (note that included in this formula is an assumption that the detection limit for PCR is 10 GC. The actual detection limit can vary depending on the presence of PCR inhibitors in a sample). |
### Table 5
Susceptibility categories for evaluating microbial indicators

| Susceptibility subset                        | Description                                                                                                                                                                                                 |
|---------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Total Coliform Rule (TCR)                   | This subset includes all wells from identifiable US groundwater systems (380 of 412 systems) with more than two health-based TCR violations (from EPA’s Safe Drinking Water Information System Fed Data Warehouse database) and all wells from unidentifiable US and international systems that likely would have had violations based on TCR criteria. Two violations were allowed due to the possibility of violations being due to distribution issues rather than from groundwater (Lambertini et al. 2012) |
| Hydrogeology                                | This subset includes all wells located in karst, fractured bedrock, or gravel/cobble settings. Many of the individual studies provided information on the hydrogeology of well settings. For those that did not report this information, karst maps were used to determine the setting (University of Auckland 2008; Weary and Doctor 2014) |
| US Ground Water Rule indicators (GWR)       | This subset includes all wells with total coliforms and either E. coli, enterococci, or coliphage in any sample from a well (e.g., all US and international wells that might be triggered into additional monitoring based on GWR criteria) |
Table 6
Indicator and virus occurrence in groundwater from 12 groundwater virus studies

| Target            | Percent positive (n) | Percent of studies analyzing for the target |
|-------------------|----------------------|---------------------------------------------|
|                   | Samples             | Wells                                       |
| Total coliform    | 30.7 (2,013)         | 36.2 (741)                                  |
| E. coli           | 10.5 (2,009)         | 12.0 (741)                                  |
| Enterococci       | 12.1 (1,389)         | 14.9 (424)                                  |
| Aerobic spores    | 43.1 (188)           | 52.0 (25)                                   |
| Anaerobic spores  | 6.0 (567)            | 26.0 (50)                                   |
| F-specific phage  | 7.5 (1,799)          | 15.2 (644)                                  |
| Somatic phage     | 9.6 (1,801)          | 11.0 (644)                                  |
| Any indicator     | 36.9 (2,015)         | 44.5 (741)                                  |
| Culturable virus  | 3.9 (1,292)          | 7.9 (365)                                   |
| PCR-virus         | 14.1 (2,106)         | 29.6 (611)                                  |
| Any virus         | 14.7 (2,273)         | 26.7 (745)                                  |
| Adenovirus        | 7.7 (781)            | 18.3 (191)                                  |
| Enterovirus       | 5.9 (1,426)          | 13.6 (413)                                  |
| Norovirus         | 7.0 (1,597)          | 21.3 (442)                                  |
| Hepatitis A virus | 1.1 (1,072)          | 9.9 (121)                                   |
| Rotavirus         | 1.4 (927)            | 7.8 (115)                                   |
| Reovirus          | 12.9 (303)           | 33.3 (60)                                   |
Table 7

Spearman rank order correlation (rho value) of microbial indicators and viruses from the 12 groundwater virus studies

| Indicator/virus       | Sample-specific |   |   |   |   |
|-----------------------|-----------------|---|---|---|---|
|                       | Culturable virus | PCR-virus | Culturable virus | PCR-virus |
| Total coliform        | 0.22 (1,289) ***| 0.07 (1,844) ** | 0.37 (366) *** | 0.26 (608) *** |
| E. coli               | 0.30 (1,285) ***| 0.02 (1,840)  | 0.31 (366) *** | 0.20 (608) *** |
| Enterococci           | 0.27 (1,228) ***| 0.04 (1,222)  | 0.33 (349) *** | 0.31 (292) *** |
| Spores                | 0.17 (755) ***  | -0.09 (755) **| 0.18 (73)      | 0.00 (73)    |
| F-specific phage      | 0.22 (1,274) ***| 0.08 (1,632) **| 0.38 (355) *** | 0.16 (510) *** |
| Somatic phage         | 0.31 (1,276) ***| 0.12 (1,634) ***| 0.46 (355) *** | 0.29 (510) *** |
| Culturable virus      | -               | 0.13 (1,125) ***| -              | 0.19 (231) ** |

aCorrelations were performed in SigmaPlot; number of samples or wells shown in parentheses;

* P =0.01 to 0.05;

** P = 0.001 to <0.01;

*** P < 0.001; all unmarked correlations are not significant (P > 0.05)

Hydrogeol J. Author manuscript; available in PMC 2018 September 19.
Table 8
Indicator and virus associations from the 12 groundwater virus studies

| Indicator        | Viruses measured by culture | Viruses measured by PCR |
|------------------|-----------------------------|-------------------------|
|                  | Sample-level analyses       | Well-level analyses     | Sample-level analyses | Well-level analyses |
|                  | Risk ratio  | P  | Risk ratio  | P       | Risk ratio | P  | Risk ratio | P |
|------------------|------------|----|------------|---------|------------|----|------------|----|
| Total coliform   | 2.6        | 0.164 | 4.5        | 0.043   | 1.0        | 0.881 | 1.3        | 0.037 |
| E. coli         | 7.0        | <0.001 | 3.1        | 0.087   | 0.9        | 0.795 | 1.6        | 0.008 |
| Enterococcus    | 5.7        | <0.001 | 4.5        | 0.002   | 1.0        | 0.858 | 1.0        | 0.913 |
| F-specific phage| 4.4        | 0.036 | 7.7        | 0.037   | 1.2        | 0.297 | 1.2        | 0.393 |
| Somatic phage   | 9.0        | <0.001 | 9.1        | <0.001 | 1.7        | 0.176 | 1.9        | <0.001 |
| Spores          | 3.3        | <0.001 | 1.5        | 0.549   | 0.8        | 0.592 | 1.4        | 0.342 |

*Associations were evaluated at the level of sample (i.e., comparison between indicator and virus samples collected at the same time from a well) and at the level of well (i.e., designating a well as positive for an indicator or virus based on one or more positive results from multiple samples collected from that well). All analyses were adjusted for the effects of study and well (sample-level analyses) or the effect of study (well-level analyses).

*Risk ratio = positive predictive value/(1-negative predictive value)*

*p is for the overall indicator-virus association*
### Table 9

Indicator test performance for indicating the presence of human viruses measured by culture\(^a\)

| Indicator          | Sample-level analyses | Well-level analyses |
|--------------------|-----------------------|---------------------|
|                    | Sensitivity (%)\(^b\) | Specificity (%)\(^c\) | Positive predictive value \(^d\) | Sensitivity (%) | Specificity (%) | Positive predictive value (%) | Negative predictive value (%) |
| Total Coliform     | 29 (8–65)             | 79 (51–93)          | 7 (2–25) | 97 (88–99) | 73 (28–95) | 64 (32–87) | 23 (7–57) | 95 (79–99) |
| E. coli            | 26 (14–44)            | 95 (90–98)          | 18 (5–48) | 97 (90–99) | 37 (15–66) | 88 (69–96) | 29 (7–70) | 91 (65–98) |
| Enterococcus       | 30 (13–55)            | 92 (85–96)          | 11 (5–23) | 98 (96–99) | 57 (23–85) | 80 (59–92) | 24 (8–54) | 95 (80–99) |
| F-specific phage   | 11 (3–35)             | 97 (92–99)          | 10 (2–39) | 98 (95–99) | 50 (14–86) | 92 (59–99) | 40 (6–86) | 95 (80–99) |
| Somatic phage      | 14 (3–45)             | 97 (90–99)          | 16 (6–36) | 98 (96–99) | 49 (17–82) | 94 (78–99) | 46 (14–81) | 95 (82–99) |
| Spores             | 19 (2–74)             | 95 (62–100)         | 9 (4–18)  | 97 (96–98) | 48 (6–93)  | 71 (42–89) | 32 (17–53) | 79 (46–94) |

\(^a\)All sample and well results from the combined 12-study dataset were included in the analysis. Performance was evaluated at the level of sample (i.e., comparison between indicator and culturable virus samples collected at the same time from a well) and at the level of well (i.e., designating a well as positive for an indicator or virus based on one or more positive results from multiple samples collected from that well). Analyses were adjusted for the effects of study and well (sample-level analyses) or the effect of study (well-level analyses). Values in parentheses are the 95% confidence intervals.

\(^b\)Sensitivity = the percentage of virus-positive samples or wells the indicator correctly identified as virus-positive.

\(^c\)Specificity = the percentage of virus-negative samples or wells the indicator correctly identified as virus-negative.

\(^d\)Positive predictive value = the percentage of indicator-positive samples or wells that were virus positive.

\(^e\)Negative predictive value = the percentage of indicator-negative samples or wells that were virus negative.
Table 10

Indicator test performance for indicating the presence of human viruses measured by molecular methods$^a$

| Indicator           | Sample-level analyses | Well-level analyses |
|---------------------|-----------------------|---------------------|
|                     | Sensitivity (%)       | Specificity (%)     | Positive predictive value (%) | Sensitivity (%) | Specificity (%) | Positive predictive value (%) | Negative predictive value (%) |
| Total Coliform      | 12 (3–35)             | 88 (68–96)          | 9 (4–20)                       | 91 (82–96)      | 33 (14–58)       | 75 (47–91)                   | 33 (13–62)                     |
| E. coli             | 2 (1–10)              | 97 (91–99)          | 9 (3–21)                       | 91 (82–96)      | 12 (5–29)        | 94 (83–98)                   | 40 (15–71)                     |
| Enterococcus        | 4 (1–12)              | 96 (88–98)          | 7 (2–19)                       | 93 (83–97)      | 14 (3–52)        | 85 (50–97)                   | 26 (5–69)                      |
| F-specific phage    | 5 (2–12)              | 96 (90–98)          | 9 (3–22)                       | 93 (83–97)      | 17 (5–44)        | 87 (68–96)                   | 31 (7–74)                      |
| Somatic phage       | 4 (1–18)              | 98 (89–100)         | 12 (3–36)                      | 93 (82–98)      | 15 (5–39)        | 94 (82–98)                   | 45 (14–81)                     |
| Spores              | 5 (1–31)              | 94 (60–100)         | 6 (1–42)                       | 92 (69–98)      | 39 (8–83)        | 74 (41–92)                   | 69 (15–97)                     |

$^a$See footnotes to Table 9 for a description of procedures and test performance measures
Table 11

Ratio of % positive in category/overall % positive$^a$

| Susceptibility category | Samples Culturable virus | PCR-virus | Wells Culturable virus | PCR-virus |
|-------------------------|--------------------------|-----------|-----------------------|-----------|
| TCR                     | 1.9 ($417^b$)            | 1.1 (672) | 1.3 (131)             | 1.2 (148) |
| Hydrogeology            | 1.9 (407)                | 0.7 (333) | 1.5 (131)             | 0.9 (65)  |
| GWR                     | 2.2 (477)                | 1.1 (599) | 3.9 (59)              | 1.6 (118) |

$^a$See Table 5 for category descriptions

$^b$Values in parentheses are the number of samples or wells in the category
Table 12
Indicator test performance measures and risk ratios for wells placed into susceptibility categories<sup>a</sup>

| Indicator | Analysis basis | Category<sup>b</sup> | Sensitivity | Specificity | Positive predictive value | Negative predictive value | Risk ratio | P   |
|-----------|----------------|-----------------------|-------------|-------------|--------------------------|--------------------------|------------|-----|
| **Culturable virus** | | | | | | | | |
| Total coliforms | Samples | All | 29 | 79 | 7 | 97 | 2.6 | 0.164 |
| | | Hydrogeology | 58 | 66 | 14 | 96 | 3.8 | 0.016 |
| | Wells | All | 73 | 64 | 23 | 95 | 4.5 | 0.043 |
| | | Hydrogeology | 88 | 55 | 42 | 93 | 6.2 | 0.016 |
| Enterococci | Wells | All | 57 | 80 | 24 | 95 | 4.5 | 0.002 |
| | | Hydrogeology | 73 | 77 | 41 | 93 | 5.8 | 0.01 |
| | | TCR | 82 | 39 | 36 | 93 | 5.1 | 0.022 |
| F-specific phage | Samples | All | 11 | 97 | 10 | 98 | 4.4 | 0.036 |
| | | Hydrogeology | 15 | 97 | 27 | 96 | 7.2 | 0.004 |
| | Wells | All | 50 | 92 | 40 | 95 | 7.7 | 0.037 |
| | | Hydrogeology | 58 | 96 | 76 | 91 | 8.4 | 0.005 |
| **PCR-virus** | Enterococci | Wells | All | 14 | 85 | 26 | 73 | 1.0 | 0.913 |
| | | TCR | 88 | 38 | 43 | 91 | 4.9 | 0.014 |
| F-specific phage | Samples | All | 5 | 96 | 9 | 93 | 1.2 | 0.297 |
| | | Hydrogeology | 5 | 97 | 10 | 95 | 2.2 | 0.019 |
| Somatic phage | Wells | All | 15 | 94 | 45 | 76 | 1.9 | <0.001 |
| | | TCR | 42 | 86 | 56 | 80 | 2.8 | 0.043 |

<sup>a</sup>Data are reported only for susceptibility categories (see Table 5) and indicators that have statistically significant indicator-virus associations. All values were adjusted for study (i.e., accounting for underlying differences among the 12 studies) and the sample-level analyses additionally include an adjustment for well.

<sup>b</sup>The All/category data are from Tables 9 and 10 and are included here for easy comparison with the susceptibility categories.