Report on

Study of the feasibility of investigating any potential relationship between the supply of discoloured water and gastrointestinal illness

Part 1: A critical literature review
Part 2: A proposed study design

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Critical review of the literature on the association between water discolouration and acute gastrointestinal illness in settings relevant to the United Kingdom

Executive Summary

Background It is not clear whether events resulting in a sudden obvious discolouration of the water supply could result in an increase in the risk of gastrointestinal illness among those exposed. The reviewers were commissioned by the Drinking Water Inspectorate to evaluate the available evidence relevant to the potential association between drinking water discolouration at point of use and risk of acute gastrointestinal illness. The purpose of this review is to assess the existing evidence, to make recommendations regarding the need for a further study in England and Wales, and to inform the design of such a study.

Questions Addressed by the Review This review sought to find evidence relevant to a UK setting of an association between discoloured tap water and the risk of acute gastrointestinal illness and to determine what proportion of acute gastrointestinal illness in England and Wales might be attributable to exposure to discoloured water.

Review Methods The reviewers identified peer reviewed research from structured searches of databases, and non-peer reviewed research from relevant websites. Studies were included in the review if they addressed the question of the association between discolouration of a public water supply in a setting relevant to the United Kingdom and risk of acute gastrointestinal illness. Eligible papers were assessed independently by two reviewers in terms of appropriateness of the study design and analysis, results, interpretation, and the overall quality of the paper.
Results of the Review Time-series studies investigating the effect of temporal variations in turbidity at the treatment plant and subsequent incidence in acute gastrointestinal illness were the only category of studies reviewed deemed sufficiently rigorous to be informative; in general these studies provided evidence of an increase in acute gastrointestinal illness at varying lags following days of high turbidity (range 3 to 29 days). One study investigating the effect of turbidity at point of use was identified, but was judged to be insufficiently rigorous to be informative.

Discussion No studies were identified in this review that specifically addressed the issue of water discolouration incidents and their effect on the risk of acute gastrointestinal illness. With the exception of contamination with highly toxic metals, we have found no evidence that water discolouration at point of use is associated with acute gastrointestinal illness. Only one poor-quality study identified in this review measured water quality at point of use and this study found no association. Most studies identified in this review investigated the effect of variations in turbidity within normal limits at point of treatment or water treatment works final water. These studies address a fundamentally different hypothesis and their results cannot be used to assess the likely effect of water discolouration at point of use. In order to establish whether there is an association between discolouration events and risk of acute gastrointestinal illness in the United Kingdom setting, an epidemiological study specifically designed to address this question would be needed.

Conclusions There is some evidence that increases in turbidity of final water are associated with subsequent increases in the incidence of acute gastrointestinal illness at varying lags. Lags of between 4 and 13 days were commonly reported. A peak in acute gastrointestinal illness at certain lags following days of high final water turbidity was consistently found across studies. This association is unlikely to be the result of measurement error, bias or random error and warrants further investigation. The potential for residual confounding (due to inadequate
adjustment for time-varying confounding factors such as seasonal effects, temperature and precipitation) in these studies remains unclear. Further methodological work in this area could clarify this issue. No evidence was found as to whether turbidity levels at point of use- in the absence of increased turbidity at the treatment plant- increase the risk of acute gastrointestinal illness.

**Background**

Forty-four incidents of discoloured water affecting approximately 1.4 million people were reported in 2003 to the Drinking Water Inspectorate of England and Wales (DWI), based on complaints from members of the public (1). This was down from 95 such incidents reported in 1999. Most of these incidents involved concomitant increases in turbidity far above the threshold of action of 4 Nephelometric Turbidity Units (NTU). Discolouration of tap water can be caused by the disturbance of sediment, particularly corrosion products, within the distribution systems. Poor control of the distribution system and ingress into the distribution system may coincide with presence of gastrointestinal bacterial species such as *Salmonella, Campylobacter, Shigella* or toxigenic strains of *E. coli*, or the parasites *Giardia* or *Cryptosporidium* that transiently pass through the system.

The reviewers were commissioned by the DWI to write a study proposal investigating the potential association between incidents of drinking water discolouration at point of use and risk of acute gastrointestinal (GI) illness. The purpose of this review is to evaluate the existing evidence in order to establish what is known on this topic, to make recommendations regarding the need for a further study addressing this issue in England and Wales, and to inform the design of the proposed study.
Several databases of published literature were searched using a specific search strategy; websites of key organisations were searched for unpublished studies. Researchers and professionals in the field of water research were contacted for information on any other studies not identified by the reviewers. Potentially relevant studies were assessed by two independent reviewers (CT, AM). Based on this critical appraisal, the evidence for an association between water discolouration and risk of acute GI illness was evaluated. Findings of the review were summarised and recommendations made as to the need for and design of the proposed study.

Questions Addressed by the Review

The central question addressed by this review was the following: is there evidence relevant to a United Kingdom (UK) setting of an association between discoloured tap water and the risk of acute GI illness and, if so, what proportion of acute GI illness in England and Wales could be attributed to exposure to discoloured water? This question can be further broken down as follows:

- What published peer-reviewed or non-peer-reviewed literature exists about the potential association between water discolouration and acute GI illness?
- What is the evidence for or against an association?
- What is the quality of this evidence?

The results of the review were used to make recommendations about the need for a further study in a UK setting, as well as to guide the design of any such study should one be deemed necessary by the DWI.
Review Methods

Study selection

The review included studies that investigated the effect of exposure to some aspect of
discolouration of drinking water from a treated public water supply (i.e. not a private well) on the
risk of acute GI illness in the population(s) served by the affected supply. Studies were
considered eligible if they met the following criteria:

1. They were conducted in a setting relevant to the UK in terms of water supply
   infrastructure and incidence and aetiology of acute GI illness (namely Europe, North
   America, and Australia).
2. The exposure was defined as some aspect of discolouration of the water supply at any
   point in the distribution system (i.e. pre-treatment, leaving the treatment works (water
   treatment works final water, or WTW final water), in transit (e.g. reservoirs), or at point
   of use (tap water)). This included colour (defined either qualitatively (presence or
   absence) or quantitatively using colour units) and turbidity, one possible aspect of
   discolouration.
3. The outcome was defined according to symptoms associated with acute GI illness.

Reports of outbreaks of acute GI illness were excluded. In general, outbreak investigations are
triggered by increases in the incidence of acute GI illness over a short time-period. Reports of
such investigations are likely to comprise highly unusual events that are, by definition, associated
with illness, but are unlikely to be representative of breaches in water quality generally. A review
of outbreak reports would be biased, as investigations of events that are not associated with illness are not generally conducted.

Data sources and search strategy

A number of key papers were identified in an initial exploratory phase of the review. These papers were read and subject headings associated with each paper obtained by entering their citations into the Pubmed database. A preliminary search strategy based on these subject headings was conducted in Pubmed. In discussion with the study team, the strategy was modified slightly (Annex A). Papers identified were initially categorised by setting: those not relevant to the UK setting (as defined by the eligibility criteria above) were excluded at this stage. Abstracts of remaining papers were read, a shortlist of potentially relevant papers generated, and the full texts of these articles retrieved to determine whether or not they were eligible. The same procedure was repeated in other key databases and on websites holding non-peer reviewed publications (grey literature) (Annex A). A list of eligible studies was generated using this process and the reference lists of these papers checked for further potentially relevant studies. This process was repeated until no new papers were identified. At this time, the reviewers also contacted, via email, a list of experts in the field of drinking water quality, as well as authors of eligible studies identified, to request information on any other published or unpublished studies relevant to the review (Annex C). At a later date, a separate search for papers related to a known incident of aluminium contamination was performed and one additional study identified.

Study quality assessment

Salient points of eligible studies were summarised by one reviewer (AM). Each study was then assessed by two independent reviewers (CT, AM) according to a list of criteria defined a priori
(Annex B). The results of these independent assessments were compared and summarised in text form.

Data synthesis

We classified all studies according to whether discoloration/turbidity was measured at point of treatment or point of use (in the latter case, provided that discoloration/turbidity readings at point of treatment were within the range considered to be acceptable in that setting). We classified all studies according to whether discoloration/turbidity was measured at point of treatment or point of use (in the latter case, provided that discoloration/turbidity readings at point of treatment were within the range considered to be acceptable in that setting). Studies were then divided into two categories, according to whether the outcome was hypothesised to have a microbial or chemical aetiology, and subdivided according to the specific research question addressed and the study design. A number of studies in the microbial aetiology category all addressed a similar question using similar methodology, but were subdivided into two groups (Groups 1 and 2) according to the type of health outcome used. A further three studies in this category all addressed different questions and had different designs, and so were reviewed individually (Groups 3-5). Each group of studies was assessed according to four broad categories: appropriateness of the study design and analysis, results and interpretation- including possible alternative explanations for the results seen- and the overall quality of the paper.
Details of the included and excluded studies

Figure 1. Flow-diagram of process of identifying key papers.

<table>
<thead>
<tr>
<th>Total number of citations identified after initial electronic search: Pubmed 3226 ORS, 7799 NORS, Embase 748 ORS, 1980 NORS, ASFA/IAMA 6265, Sigle 13 (n=20,031)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potentially relevant citations after screening of electronic search: Pubmed 20 ORS, 2 NORS, Embase 8 ORS, 33 NORS, ASFA/IAMA 7, Sigle 10, Grey literature 9 (n=89)</td>
</tr>
<tr>
<td>Citations excluded after reading full-text revealed they did not meet inclusion criteria (n=72)</td>
</tr>
<tr>
<td>Key papers identified (n=16)</td>
</tr>
<tr>
<td>Citations excluded: MacKenzie 1994, Stirling 2001 (outbreak investigation reports) (2)</td>
</tr>
<tr>
<td>Papers included in critical review (14)</td>
</tr>
</tbody>
</table>

ORS = obviously relevant setting (to UK), NORS = not obviously relevant setting
ASFA = Aquatic Science and Fisheries Abstracts 3: Aquatic Pollution and Environmental Quality
IAMA = Industrial and Applied Microbiological Abstracts (Microbiology A)

Results of the Review

A total of 14 studies were identified. The expert responses the reviewers received led to the identification of two peer-reviewed publications (8,9) that were subsequently included in the review (Annex C). Of the 14 studies included in the review, 12 were papers in peer-reviewed scientific journals (3,5,6,8-16) and two were Health Canada reports (4,7). Eleven studies measured discolouration/turbidity at point of treatment (3, 5-14) and one of these studies looked additionally at turbidity at the tap (13); one study assessed pre-treatment turbidity (4). Two additional studies assessed the association between copper contamination (15) and aluminium
contamination (16) causing discolouration of tap water. Of studies in which discolouration/turbidity was measured at point of treatment, all dealt with an outcome of hypothesised microbial aetiology.

We identified three studies in European or North American settings that investigated the association between various aspects of water discolouration (turbidity, copper and aluminium contamination respectively) at point of use and risk of acute GI illness (13,15,16). These studies were of poor methodological quality and did not seek to assess the population impact of discolouration (i.e. the proportion of acute GI illness in the population that could be attributed to this exposure). With the exception of the study that assessed pre-treatment water turbidity (4), the remaining studies investigated the putative association between drinking water turbidity leaving the plant and acute GI illness (3, 5-12); one of these studies assessed WTW final water and tap water turbidity, colour, free residual chlorine and iron (13).

Nine of the 11 studies using turbidity as the exposure employed time-series methodology; six of these studies found a positive association between increases in WTW final water turbidity and subsequent increases in incidence of acute GI illness at similar time lags (3,5,6,8,9,10). One study found a positive effect of increased daily raw (pre-treatment) water turbidity and daily counts of acute GI illness at the same lags (4). These significant results were found in different age groups, including children and the elderly.

Seven of the 14 studies used hospital admissions for acute GI illness as the outcome (3-9). Two studies additionally used physician visits (4,7). One study also included visits to long-term care facilities for symptoms related to acute GI illness (7), seven used visits to accident and
emergency (A&E) departments and two included visits to hospital outpatient departments (6,9). In all these cases, acute GI illness was defined on the basis of those ICD-9 codes most plausibly associated with acute GI illness. One further study used A&E visits for putative acute GI illness, but did not define these according to ICD-9 codes (13). Other outcome definitions used were records of sales of prescription and non-prescription medications for acute GI illness (11), self-reported illness from diaries or telephone surveys (10,13-16), frequency of bowel movements and medication use at nursing homes (14), clinical specimens submitted to microbiology laboratories for confirmation of suspected acute GI illness (12), and positive reports of laboratory-confirmed acute GI illness (14).

**Microbial aetiology (Table 1)**

**Group 1 (3-9)**

This group of studies is relevant to the UK setting - the studies were all conducted in North American cities with similar water distribution systems and where incidence and aetiology of GI disease are likely to be comparable to those in the UK. All of these studies investigated the effect of temporal variations in turbidity levels at point of treatment on incidence of acute GI illness in the population served by the treatment plants. The studies were generally of good methodological quality and their results were similar in terms of the magnitude of the associations seen and the lag between exposure and disease.

**Appropriateness of design and analysis**

These studies all used time-series-adapted multivariable regression analysis, which the reviewers judged to be an appropriate method of analysis for the questions addressed by the studies. Two studies also used individual risk factor analysis (binomial Generalised Linear Modelling (GLM))
to assess temporal differences in exposure to turbidity levels between cases of acute GI illness and controls with respiratory illness (4,7). The results of these individual level analyses were consistent with those of the time-series analyses.

In time-series analysis, the population studied serves as its own control at different points in time. Factors that do not change appreciably with time are thus unlikely to influence any observed association between the exposure and the outcome. Only factors that display temporal variation need be considered as potential confounders. These normally include long-term trends in exposure and outcome, seasonal patterns and shorter-term systematic (non-random) variations. The number of such factors, in addition to the exposure and outcome, included in regression models varied between studies. Two studies included day of the week only (6,8), one study also included season (9), while the studies performed in Philadelphia included in the base model terms for long-term time trends, day of the week and temperature, adding season in the earlier of the studies (3,5). In addition to the factors controlled for in the Philadelphia studies, the Canadian studies added terms for precipitation and public holidays (4,7).

Results

Five of seven studies found a positive association between daily finished water turbidity and daily counts of acute GI illness, measured as either relative risk or percent increase in illness for a given increase in turbidity at similar lags (3,5,6,8,9); an association in the same direction and of similar magnitude was found in the study of pre-treatment, raw water turbidity (4). Four studies found a significant association between daily WTW final water turbidity and daily counts of acute GI illness at lags of between 4 and 13 days, (3,5,8,9). The study that assessed the
association between daily fluctuations in raw (pre-treatment) water turbidity and daily counts of illness (4) recorded significant lags at between 3 and 29 days.

Heterogeneity between these studies may be explained by variation in the time between turbid water leaving the treatment plant and reaching the consumer (a characteristic of the water distribution system), variation in incubation periods in different settings (either due to differences in causative pathogens or in age groups affected), differences in the average time between illness and individuals seeking medical attention in different settings, or random variation. It should be noted that the lags found to be significant in the studies of WTW final water turbidity are consistent with the incubation period – the period from infection to onset of symptoms – for most known waterborne infectious gastrointestinal pathogens (1-14 days), especially *Cryptosporidium* species (19).

One study (6) tested one aggregate lag of 1-7 days, and thus could not assess the significance of lags shorter than one week. The final study in this group (7) did not find any significant lags among the 40 tested, which may suggest that, in this setting, WTW final water turbidity was not a good marker of microbiological contamination and/or that the quality of water treatment was superior.

Generally, relative risks reported were of the order of 1.2 – 3.0 (corresponding to an increased risk of between 20% and 200% per category of turbidity) (3-9). However, there was variation among the studies in the way in which this association was measured. Some studies examined the increase in GI illness for a 1 NTU increase in turbidity whilst others examined the effect of an interquartile change in turbidity levels. No studies reported a reduction in risk with increased
levels of turbidity, which might be expected if those exposed to obviously turbid water were less likely to drink it. Associations were found in different age groups – all ages, children and elderly – and among children and the elderly the association was stronger in the extremes of age.

Most of the studies did not report what proportion of acute GI illness in the population could be attributed to the exposure investigated. The study that assessed the effect of raw (pre-treatment) water turbidity (4) estimated that this exposure could account for 0.2-29% of daily admissions, physician visits and children’s A&E visits for acute GI illness, depending on level of turbidity and age; another study of WTW final turbidity (8) estimated that turbidity above 1 NTU was associated with 37.5 excess outcomes per 100,000 persons.

*Interpretation*

The ecological nature of time-series studies makes their interpretation difficult. As exposure data are not collected at individual level, the whole population is assumed to be exposed to the same extent. In addition, the temporal nature of the relationships studied means that any observed association might be due to coincident patterns in the seasonality of turbidity (or other time-varying factors) and acute GI illness, unless sufficient adjustment for these is made in the analysis. In assessing the strength of evidence for a causal association between turbidity and acute GI illness, the following possible alternative explanations should therefore be considered: measurement error of exposure, misclassification of exposure and/or outcome, residual confounding, and random error. We deal with each of these in turn below.
Measurement error of exposure

In all these studies, the instruments used to measure turbidity were those routinely used by water treatment plants to monitor quality as required by national guidelines. The Philadelphia studies (3,5) on which the methodology for all these studies is based, have previously been criticised on the basis that turbidity was found to have an effect on acute GI illness below the limit of detection for which turbidity meters were calibrated, making the results unreliable (20). The authors have countered this critique by arguing that readings below this level are routinely used to demonstrate compliance with legal requirements and, further, that measurement errors in readings below the limit of detection are random (i.e. not temporally associated with either turbidity or acute GI illness) and would, therefore, be expected to attenuate any observed association rather than be an alternative explanation for it (21). All studies used the mean of several daily readings of turbidity, increasing the accuracy and precision of exposure measurements, and all studies measured turbidity in the same units (NTU). Thus, measurement error of the exposure is unlikely to explain the association between turbidity and acute GI illness.

Misclassification of exposure

Misclassification of exposure could lead to a spurious association if a proportion of the study population was not actually exposed to the water supply studied (or was exposed to a different extent), and this difference in exposure was related to the risk of acute GI illness. For example, some individuals may have regularly consumed water from sources other than the local public water supply. In these studies, however, such misclassification would also have to be temporally related to both turbidity levels and GI illness. Most studies demonstrated a peak effect of turbidity on GI illness at specific lags, and there is no reason to believe that cases with misclassified exposure would systematically present to health services with GI illness on specific
lags following high turbidity days. Any potential misclassification of the exposure is thus likely to be random and have the effect of tempering rather than inflating the association between turbidity and GI illness.

**Misclassification of outcome**

In all these studies, the main outcome (hospital admission) was defined using ICD-9 codes plausibly related to acute GI illness. This method is superior to definitions based on, for example, patient self-report, as it is far less likely to be biased by individuals’ perception of their exposure level. Most of these studies (4-9) also used other sources of data on GI illness with less specific case definitions. In these cases, investigators carried out sensitivity analyses to assess the effect of including these less specific groups of cases. In general, these did not influence the results.

**Residual confounding**

Confounding could occur if insufficient adjustment for a time-varying factor was carried out or if important time-varying factors were omitted from the analysis. In particular, the degree of adjustment for seasonal patterns in both turbidity and GI illness is crucial. Insufficient adjustment could result in an association being observed simply because turbidity and GI illness are related in time and share similarities in their seasonal patterns, while over-adjustment could mean that no association is found. All analyses included some form of seasonal adjustment (usually locally-estimated sum of squares, or loess, smoothing) and, depending on the length of the time-series and the time-steps used, terms for long-term (year-on-year) trends and day-of-week effects. In general, however, the description of statistical analyses was not sufficiently detailed and the potential reproducibility of the methods based on the information provided in the papers was deemed to be poor. Few studies presented details of model diagnostics, such as residual plots to
assess the appropriateness of seasonal adjustment, and it was difficult to judge whether all or part of the observed association could be due to residual confounding. Some studies adjusted for climatic factors potentially associated with turbidity and GI illness, namely ambient temperature and precipitation.

Random error

The testing for associations between turbidity and GI illness over a number of time lags raises the issue of whether some associations could have occurred by chance alone. The studies reviewed tested between 1 and 40 lags. With a 0.05 level of precision it would be expected that, on average, 1 in 20 lags tested would be statistically significant simply by chance, assuming the effect of each individual lag was independent of all others. Most studies assessed the effect of turbidity at various lags using temporal exposure response surface (TERS) plots. For studies showing an association, these three-dimensional plots demonstrated that increases in GI illness clustered around specific lags following days of high turbidity levels at the treatment plants. If random error due to multiple testing were a likely explanation for the observed associations, increases in incidence would be expected to be distributed more evenly along the surface of the plots, with peaks also occurring following low turbidity days across all lags tested.

In summary, the associations found between daily variations in WTW final turbidity and subsequent increases in GI illness at particular lags are unlikely to be entirely explained by measurement error, exposure or outcome misclassification, or random error, although residual confounding due to insufficient seasonal adjustment or failure to account for other relevant time-varying factors could potentially explain all or part of the observed associations. However, the consistency of the results across different settings using different levels of adjustment argues
against this being the case. The time lags identified between increased levels of turbidity and subsequent increases in incidence of GI illness are consistent with the incubation periods of most known waterborne pathogens and Cryptosporidium in particular, which due to its high resistance to chlorine makes an association biologically plausible (19).

**Group 2** (10,11)

Of these two studies, one was set in Russia (10) and is not as applicable to the UK setting, as the water supply was considered to be unsafe by the local public health authorities and residents were advised to always boil drinking water; the remaining study (11), set in France, may be relevant to a rural UK setting.

**Appropriateness of design and analysis**

These two studies also employed time-series analysis, but differed from group 1 studies in that their definition of the outcome was less specific - one study (10) used self-reported illness, as recorded daily by study participants on diaries, and the other (11) used 3 and 7-day sales of prescription and over-the-counter anti-diarrhoeal medications.

In the analysis, one study (10) controlled for seasonality, day-of-week effects, consumption of unboiled tap water, attendance at summer houses and travel outside the study area; the other (11) controlled for long-term trends in drug sales, as well as seasonal and day-of-week effects.

**Results**

The Russian study (10) found a positive association between daily WTW final water turbidity and acute GI illness at a lag of 2 days. The French study (11) reported a 14% increase in sales of
anti-diarrhoeal medications per 10 NTU increase in raw (pre-treatment) water turbidity at a lag of 1 to 3 weeks; no significant association between WTW final water turbidity and sales of medications was found.

**Interpretation**

The following possible alternative explanations will again be considered: measurement error of exposure, misclassification of exposure and/or outcome, residual confounding, and random error.

**Measurement error of exposure**

As in group 1, these two studies measured turbidity with instruments used routinely by water treatment plants to monitor quality as required by national guidelines, and the mean of several daily readings of turbidity was used in the analysis. Errors in these measurements are likely to be random and the associations found in these two studies are unlikely to be entirely explained by measurement error of water turbidity at point of treatment.

**Misclassification of exposure**

Misclassification of exposure is more likely in these two studies. In one (10), visits to summer houses, where participants would have been exposed to private water supplies, were likely to be an important factor, but it was unclear how the investigators took this into account in the analysis. The other study (11) was prone to exposure misclassification, since medication sales could only be approximately linked to corresponding water supply zones. It is likely that a proportion of medications was sold to people who were not resident within the water supply zones studied; this could have affected the study’s ability to detect an association.
Misclassification of outcome

The Russian study (10) relied on participants’ self-reported illness in diaries – an ascertainment method prone to bias, particularly if individuals are aware of the hypothesis under study. Knowledge of the hypothesis may have made participants more likely to report symptoms of acute GI illness if they believed they had been exposed to turbid water. In the French study (11), sales of medications to individuals not exposed to the local water supply (up to 20% of sales according to the investigators) would have resulted in the inclusion of putative cases of acute GI illness who were not part of the population at risk. This could have contributed to a failure to find an association if one did indeed exist.

Residual confounding

As with group 1, adequate control for time-varying factors is an important issue in these two studies. Both studies controlled for long-term, seasonal and day-of-week effects. However, the extent to which this was done and the adequacy of these adjustments was more difficult to assess from the information presented in the reports. In the Russian study (10) information was collected on individual-level behavioural factors thought likely to influence risk of acute GI illness, but it was unclear how these were aggregated for inclusion in time-series analyses. In addition, the investigators reported considerable differences in behavioural factors by age (for example, younger participants were less likely to boil water before consumption), which was not adjusted for in the analysis. Exposure to private water supplies during visits to summer houses is also likely to have been an important confounder, but it was not clear whether this was adequately addressed.
Random error

Both studies had low power; one study (11) included medication data from only one pharmacy and the population studied was small. The Russian study included data from follow-up of only 100 families. The study failed to take into account the lack of independence of measures within families, which could potentially result in a spurious association being found.

In summary, both these studies presented some evidence for an association between temporal variations in turbidity and subsequent risk of acute GI illness. However, the studies had considerable limitations in terms of the precision with which exposure and outcome were ascertained, potential for bias, control of confounding factors and statistical power. We consider the evidence provided by these two studies to be far less convincing than that provided by group 1 studies.

Group 3 (12)

This study investigated changes in incidence of acute GI illness (as assessed by the number of stool specimens submitted for laboratory investigation of suspected GI pathogens) in a number of communities in Australia following the introduction of improvements to the treatment of the local public water supply. The improvements consisted of retrofitting disinfection and/or filtration processes to existing systems.

Appropriateness of design and analysis

This study had a conventional ecological design. The introduction of improvements in water treatment in a number of communities was assessed for its effect on the mean annual turbidity
and the rate of submission of faecal specimens for microbiological investigation during a 12-month period before and after the changes to the local water supply. The rate of submission of midstream urine samples (MSU) for laboratory investigation was used as a control to rule out the possibility that changes in incidence were due to changes in patterns of specimen submission for laboratory testing. This study design had several limitations. The nature of the improvements in water treatment varied between communities and some communities had no treatment system prior to the study. Thus, any effect on the incidence of acute GI illness is likely to have differed by community. Turbidity values were averaged over a 12-month period and this may have obscured correlations present only at higher turbidities, which might be transient. The study did not take into account trends in acute GI illness in the different communities prior to the introduction of water treatment upgrades, making any before and after comparisons of incidence of acute GI illness difficult to interpret. Due to the non-specific nature of the outcome (submission of faecal specimens for investigation), the study is likely to have measured relative changes in incidence of acute GI illness rather than actual incidence. In addition, the treatment plants studied did not serve the entire postcode from which faecal samples originated, and the precision of exposure definition varied between communities. This means that a varying proportion of the various populations studied would not have been expected to be affected by changes in the water supply. The use of MSU samples is unlikely to have been a suitable control, as the age distribution of conditions resulting in the submission of MSU samples is very different to that of acute GI illness.

Results

No significant correlation was found between the implementation of improvements to the water treatment and the rate of faecal specimen submission. Although in most cases, treatment upgrades
resulted in a decrease in mean turbidity, in some cases an overall increase in turbidity was observed, and no correlation between change in turbidity after retrofit and change in the rate of faecal specimen submission after retrofit was found.

Interpretation

This study addresses a different question to studies in groups 1 and 2, namely, do improvements in treatment of the water supply, which are associated with overall changes in the mean turbidity level, have an effect on the overall incidence of acute GI illness. Given the methodological problems highlighted above, we did not find this study to be informative in terms of the putative role of discolouration/turbidity on risk of acute GI illness.

Group 4 (13)

This Russian study was also set within a context in which residents were advised to boil water all year round and so is not as directly relevant to a UK setting.

Appropriateness of design and analysis

This study employed a cross-sectional design and investigated the association between change in colour, turbidity, free residual chlorine and iron in drinking water (from plant to point of use) and self-reported GI illness. Families were recruited into the study from households within the city’s water distribution zone. Participants recorded details of acute GI symptoms in a diary and additionally provided information on a number of demographic, household, behavioural and other relevant factors. Water quality at point of use was measured at regular intervals from households within the distribution zone (but not necessarily those of participants). Regression
analyses were conducted to determine whether any associations existed between water quality indicators and incidence of self-reported acute GI illness.

**Results**

No significant associations were found between turbidity or iron content and GI illness, but some effect of decreased free chlorine on acute GI illness was observed. The effect of colour on illness was not assessed.

**Interpretation**

A number of limitations were apparent in the design and implementation of this study. Firstly, no sample size calculations were presented, making it difficult to assess whether the study had sufficient power to detect an association between the various water quality indicators and acute GI illness. The measurement of water quality parameters is likely to have been subject to considerable error; water quality was measured in several sites for each study area, but only two measurements per site separated by three weeks were taken, which is likely to have negatively influenced the accuracy and precision of these parameters.

Misclassification of exposure is likely to have been a problem in this study as water quality indicators were not measured frequently and were measured at sites other than participants’ homes. No details were given as to how sites for water sampling were selected, and it is unclear how representative these sites were of the water distribution zone as a whole. The study did not account for water consumed outside the home, which could have introduced a bias if, for example, participants consumed water from other sources when water quality was particularly poor.
The information supplied by one of the interviewers was excluded from the analysis (due to breach of study protocols). This would have decreased the power of the study to detect an association and could have introduced bias if participants excluded from analysis differed systematically in important ways from the rest of the study population.

In summary, although no association between turbidity and acute GI illness was found, it is unclear whether the study had sufficient power to actually detect an association and failure to detect an association could have resulted from the considerable potential for bias in the study design, most notably through misclassification of participants’ exposure.

**Group 5 (14)**

This study involved a telephone survey of Washington, DC residents and retrospective review of health data following a failure of filtration in the local water supply that resulted in an increase in turbidity levels above legal limits. The two-week period before the filtration failure was compared with the two-week period following the failure to determine whether an increase in incidence of acute GI illness was apparent.

*Appropriateness of design and analysis*

This study employed an ecological before and after design, but did not use a control area as a comparison. For the telephone survey, participants were asked to give details of acute GI illness and water consumption for varying lengths of recall (two weeks for the period after the filtration failure compared with four weeks for the period before the failure). As the quality of recall varies with time, it is likely that information for the period before the filtration failure was less accurate.
Results

No significant increase in incidence of acute GI illness (as measured by the telephone survey or various health data sources) was found following the filtration failure.

Interpretation

It is likely that the study did not have sufficient power to detect a significant increase in incidence in the period following the filtration failure, as it included a single incident taking place over a very short time period. No attempt to exclude the possibility of random error was made (for example, by using a control area, or making comparisons with data for previous years for the same period). In addition, the exposed population was not precisely defined (e.g. by linking affected postal codes to data on health outcomes). This is likely to have affected the analysis of emergency hospital visits, for which 30% of cases were not local residents and, thus, may not actually have been exposed to the local water supply. It is unclear to what extent the results of the telephone survey could have been biased by differential recall.

Non-microbiological (15,16) (Table 2)

These studies investigated the effect of contamination of the water supply at point of use with copper and aluminium - which cause changes to the normal colour of water – on the risk of acute GI illness. The studies were set in the US and UK.

Appropriateness of design and analysis

The US study was a descriptive report of an investigation into contamination of drinking water as a result of newly-installed copper pipes. The study reported the frequency of symptoms of copper
poisoning in the affected households following the incident, but no comparison group was used and no correlations between measurements of actual copper levels in tap water and acute GI illness were investigated. The UK study was a retrospective cohort analysis following an incident of aluminium contamination. The frequency of acute GI illness in the affected area was compared to that in a control area served by a different treatment plant. No quantification of actual exposure to aluminium was performed.

Results

Evidence of an association between copper contamination and acute GI illness was anecdotal, as no analytical investigations were conducted. In the aluminium study, there were significant differences in the proportions of people reporting an observed change in water colour, drinking habits and illness following the contamination incident (see Table 2). The calculated relative risk of diarrhoea was 5.1 (95% CI: 3.3-7.9).

Interpretation

Chemical poisonings due to copper and aluminium are recognised conditions. Such poisoning through the water supply is, however, very rare and occurs under very specific conditions (such as through inadequate installation of new pipes or accidental contamination of the water supply, as in the above studies). Their effects are thus likely to occur in highly unusual circumstances that may not be as relevant to the proposed study. Incidents in which water appears ‘brown’ or where discoloration is caused by the disturbance of iron sediment in the distribution pipes are likely to be more common forms of chemical discoloration, and the effects of these on acute GI illness are less well established. This review did not find any studies addressing these more common forms of chemical contamination.
**Discussion**

*Principal findings of the review*

In this review, we found no studies that specifically addressed the issue of water discolouration incidents and their effect on the risk of acute GI illness. We have found no evidence that water discolouration at point of use is associated with GI illness. Only one study (13) identified in this review actually measured water quality at point of use and, although this study found no association, it was deemed to be insufficiently rigorous to provide conclusive evidence. The studies relating to copper and aluminium exposure are not typical of routine discolouration incidents.

Most studies identified in this review investigated the effect of water turbidity at point of treatment (either pre-treatment or WTW final water). It should be noted that causes of discolouration or high turbidity at the treatment plant are likely to differ from causes of discolouration/turbidity in the distribution zone and that, therefore, any potential effects on the risk of GI illness are likely to be different. Moreover, most of these studies investigated the effect of temporal variations in turbidity at point of treatment under normal operating conditions (that is, where turbidity levels were within limits regarded as acceptable). However, most discolouration incidents reported to the DWI involve increases in turbidity levels well in excess of legal limits. Thus, results from these studies cannot be used to inform the specific question of whether discolouration incidents result in increased risk of acute GI illness, determine the likely magnitude of any such effect, or estimate its potential population impact.
In order to establish whether there is an association between discolouration incidents and risk of acute GI illness in the UK setting, as well as determine the magnitude and population impact of any such association, an epidemiological study specifically designed to address this question would be needed.

Weaknesses of review

This review may be biased toward published, peer-reviewed literature, which is more comprehensively catalogued and easier to retrieve than grey literature. The problem of publication bias should be considered in any systematic review, as studies reporting positive associations may be more likely to be published than those finding no association. This review may also be biased towards papers dealing with outcomes with infectious aetiology, either because more such studies have been performed or because the search strategy employed was less sensitive to outcomes caused by non-infectious agents.

Meaning of review’s findings

Although this review comprised a group of studies of varying hypotheses and designs, the most common hypothesis investigated was that of the putative association between variations in turbidity levels within acceptable limits at point of treatment and subsequent risk of acute GI illness. The findings of this review suggest that a causal association is likely and biologically plausible. The potential limitations of these studies, which generally use ecological time-series methodology, have been extensively discussed in the literature, particularly with regard to measurement errors in turbidity levels, misclassification of exposure (because no individual level exposure information is collected), and the possibility of finding spurious chance associations due to the testing of multiple lags. We have discussed these concerns above and concluded that,
on balance, the design of group 1 studies was scientifically rigorous and that these factors are unlikely to explain the significant associations found. In general, these studies provided insufficient detail regarding the degree of adjustment for time-varying factors, which may be important confounders in such time series analyses. In particular, the level of seasonal adjustment appeared to involve a degree of subjectivity and the reproducibility of these analyses based on the information provided in the reports was felt to be poor. Further methodological work investigating the most appropriate methods for seasonal adjustment would help clarify this issue. However, the general agreement between these studies, carried out in different settings, in terms of the direction, magnitude of association and significant lags between exposure and disease lends weight to the argument for a causal association. The lags identified were consistent with the incubation periods of likely causal pathogens, most notably *Cryptosporidium*, which due to its high chlorine resistance makes such an association biologically plausible. The time-series design does not lend itself to a meta-analysis for obtaining a combined estimate of effect across all studies, as it is unclear how individual studies should be weighted. A measure of population impact (that is, the proportion of acute GI illness attributable to variations in turbidity within normal limits) is also difficult to estimate from these studies, as they used hospitalised and emergency cases, which are likely to represent more severe disease and constitute a minority of all cases occurring in the community. The context of these studies was relevant to the UK situation, suggesting that a similar association might be found in the UK setting. However, because of the differences in organisation of healthcare and, potentially, important time-varying factors (e.g. temperature, precipitation), the findings of these studies cannot be readily extrapolated to inform the likely magnitude of effect and population impact in the UK. Despite the consistency of their findings, these studies did not draw strong conclusions, nor make strong recommendations.
Conclusions

Questions not answered by this review

Due to a lack of research in this area, neither the effect nor the population impact of water discolouration at point of use on the risk of acute GI illness can be determined from the existing literature. Although there appears to be consistent evidence of a positive association between increases in turbidity at point of treatment and subsequent incidence of acute GI illness, turbidity is only one of many contributing factors to discoulouration. Moreover, data on water quality at point of treatment cannot be extrapolated to the situation at point of use, as the causes of increases in turbidity and discoulouration (and, indeed, other water quality indicators) at the treatment plant and the distribution zone are likely to be different.

Implications for the design of the proposed study

1. The proposed study will use incidents of water discoulouration (rather than quantified turbidity) as the exposure. Discoulouration can have different causes and thus be indicative of exposure to many factors (both chemical and microbiological) that may or may not be associated with acute GI illness, the definition of exposure will be less precise than, for example, in studies investigating only the effect of turbidity. The study should have sufficient power to enable analyses stratified by the most relevant components of discoulouration.

2. Adequate information on possible confounding factors should be collected. Depending on the study design, these may include age, sex, socioeconomic factors, as well as time-varying factors such as temperature, precipitation and seasonality of exposure and outcome if relevant.
3. Hospital admissions, the outcome of interest in the majority of papers in this review, are likely to represent less common, more severe cases of acute GI illness. Data from these cases may not necessarily be applicable to the wider spectrum of acute GI illness. More common outcomes comprise milder disease, but are less likely to be captured by routine sources of health data. A specific sub-study to investigate the effect of water discoloration on milder disease may be necessary.
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<td>All residents of Philadelphia, US over 65 years of age</td>
<td>Ecological study; Jan 1992-Dec 1993</td>
<td>Mean daily WTW final water turbidity (NTU) measurements from each of three water treatment plants</td>
<td>Hospital admissions for ICD-9 codes plausibly related to GI illness separately for each plant service area</td>
<td>Poisson Generalised Additive Modelling (GAM) time-series correlating daily counts of hospital admissions with daily water turbidity (with lags)</td>
<td>Increase in turbidity 0.035 NTU associated with 9% (5.3-12.7) increase in admissions over 3 plants combined at lag of 9-11 days controlled for time trends, seasonal patterns and temperature; also 9.1% increase (5.2-13.3) at lag of 4-6 days for one plant; effect greater in those over 75 years (p&lt;0.0001)</td>
<td>Design Unclear whether some areas served by more than one plant Not clear if all cases in exposed area were ascertained No details as to sample size/power Analysis Insufficiently detailed methods section No measure of impact</td>
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<td>Aramini (2000)</td>
<td>Population of Greater Vancouver, Canada</td>
<td>Ecological study, January 1992-December 1998</td>
<td>Mean daily raw turbidity (NTU) for three watersheds provided by Greater Vancouver Regional District</td>
<td>Daily hospital admissions, physician visits, and BC Children’s Hospital A&amp;E visits for ICD-9 codes association with gastro illness and respiratory outcomes (controls for binomial modelling)</td>
<td>Poisson GAM time-series correlating daily counts outcome with daily water turbidity (with lags) Also, binomial Generalised Linear Modelling (GLM) regression modelling for individual risk analysis</td>
<td>Strongest association between turbidity and illness at lags of 3-6, 6-9, 12-16 and 21-29 days. Relative risks between 1.2 and 2.0 for different watershed/age combinations. The authors estimated that 0.2 to 29% of the GI illness (measured in this way) in Greater Vancouver could be accounted for by raw water turbidity, depending on level of turbidity and age.</td>
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<td>Mean daily WTW final water turbidity (NTU) across three treatment plants</td>
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<td>Morris (1996)</td>
<td>Inhabitants of Wisconsin (US) who would seek treatment at Medical College of Wisconsin hospitals</td>
<td>Ecological study; January 1992-April 1993 (including outbreak at end of study period)</td>
<td>Daily effluent turbidity (NTU) for each of two water treatment plants in Milwaukee obtained from water dept of city</td>
<td>Historical records from Medical College of Wisconsin of admissions, A&amp;E and outpatient visits for ICD-9 codes plausibly associated with GI illness</td>
<td>Poisson GAM time-series correlating 2-weekly counts of admissions/visits with 2-weekly water turbidity (with lags)</td>
<td>Including outbreak: For 0.5 NTU increase in turbidity, RR outpatient gastro event 1.53 (0.92-2.55) at Linwood and 1.36 (0.99-1.87) at Howard Ave for 0-18 years. Emergency gastro events: 2.82 (1.44-5.52) Linwood and 1.73 (1.19-2.50) Howard Ave for same agegroup. Effect was lower and non-significant for &gt;18 years for Linwood and higher and sig. for this agegroup at Howard Ave plant. Excluding outbreak: Effect unchanged for age 0-18 for Linwood plant and all other effects were non-significant.</td>
<td>Design Unclear if multiple readings of turbidity taken per day Unclear whether cases matched to a treatment plant and what proportion of all cases of the outcome are treated in these hospitals The use of 3.5 day time step, instead of 1 day, suggests low power as this was likely done to increase the number of observations per time step Analysis Insufficiently detailed methods section Only one lag tested (aggregate of 1-7 days) Unclear presentation of results Conclusions not appropriate to results No measure of impact</td>
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<td>Lim (2002)</td>
<td>Population of Edmonton, Canada supplied by one of two treatment plants, prior to refurbishment carried out on the 10 December, 1997</td>
<td>Ecological study, 1993-1999</td>
<td>Mean daily effluent turbidity (NTU) data from one of two plants serving Edmonton (via EPCOR water services, inc.)</td>
<td>Daily hospital admissions, A&amp;E, physician and long-term care visits; latter three come under physician-billing records for ICD-9 codes associated with gastro illness and respiratory outcomes (controls for binomial modelling)</td>
<td>Poisson GAM time-series comparing daily counts of GI illness cases with daily water turbidity (with lags) Also, binomial GLM and GAM regression modelling</td>
<td>No significant lags identified between finished water turbidity and gastroenteritis. Odds ratios comparing level of GI illness before and after refurbishment ranged from 0.99 to 1.22 (no confidence intervals)</td>
<td>Design No details as to sample size/power Analysis Large number of tested (1-39 days) Insufficiently detailed methods section Overcomplicated analysis and unclear presentation of results</td>
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<td>Naumova (2003)</td>
<td>People aged 65 and older in Milwaukee county (US)</td>
<td>Ecological study; January 1, 1992-April 24, 1993 (pre-outbreak and outbreak)</td>
<td>Daily maximum effluent turbidity (NTU) at South plant</td>
<td>Hospital admissions and A&amp;E visits for ICD-9 codes plausibly related to GI illness</td>
<td>Poisson GAM time-series comparing daily counts of hospital admissions/ER visits with maximum daily water turbidity (with lags)</td>
<td>1 NTU increase in turbidity associated with RR’s of GI illness within the 95% CI 1.54-4.48. Strongest association (TERS plot) at lag of 6 (primary) and 13 days (secondary spread). Turbidity over 1 NTU associated with 37.5 excess GI cases per 100,000 people</td>
<td>Design No details of sample size/power Analysis No information given on how account taken of seasonality/time trends Insufficiently detailed methods section Unclear presentation of results Interpretation Analysis done after known that Crypto caused outbreak – circular arguments</td>
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<td>Morris (1998)</td>
<td>Inhabitants of Wisconsin (US) who would seek treatment at Medical College of Wisconsin hospitals</td>
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<td>Poisson GAM time-series comparing daily counts of hospital admissions/ER visits with daily water turbidity (with lags)</td>
<td>During outbreak: GI illness most strongly associated with turbidity (south plant) at lag of 7 days in children (correlation coefficient 0.34) and 8 days in adults (correlation coefficient 0.41). Pre-outbreak: North plant had highest turbidity and turbidity was most strongly associated with gastro illness at lag of 8 days in children (correlation coefficient 0.12) and 9 days in adults (0.09)</td>
<td>Design Insufficient data on the proportion of all cases in Wisconsin that are seen at a Medical College Hospital No details as to sample size/power Analysis Insufficiently detailed methods section Unclear and incomplete results No measure of effect calculated. Unclear and incomplete results Interpretation Incomplete discussion of plausibility of results. No discussion bias/confounding</td>
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<td>Egorov (2003)</td>
<td>100 randomly chosen families (367 individuals) in Cherepovets, Russia</td>
<td>Ecological study from June to November 1999</td>
<td>Mean daily effluent turbidity (mg/l standard kaolinite suspension) at sole water treatment plant</td>
<td>At least one of liquid stool, vomiting, severe intestinal or stomach cramps during 1 day after 1 week of being symptom-free ascertained via daily self-report diaries</td>
<td>Poisson GAM time-series comparing daily counts of GI illness with daily effluent water turbidity (with lags) GLM analysis for trend</td>
<td>Increase in turbidity of 0.8 NTU associated with RR 1.47 (1.16-1.86) at lag of 2 days after control for non-boiled tap water, behavioural covariates, day of week and seasonality. Subgroup analyses showed no association among those who always boil water, and associations at lags of 1, 2 and 7 days among those who drank non-boiled tap water</td>
<td>Year-round boil water advisory Design Self-reported illness – lack of specificity Unclear whether participants blinded to the study hypothesis – response bias No details as to sample size/power Analysis Insufficiently detailed methods section Potential household clustering not taken into account Interpretation Insufficient discussion of results No discussion of bias</td>
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<td>Beaudeau (1999)</td>
<td>Inhabitants of lower city of Le Havre, France living near one participating pharmacy</td>
<td>Ecological study; April 1993 – September 1996</td>
<td>Mean daily effluent turbidity (NTU) from two of three treatment plants in Le Havre</td>
<td>Daily sales of prescription and over the counter treatments for gastrointestinal illness ascertained from pharmacies</td>
<td>Time-series analysis (Box and Jenkins 1976); mean of 3-day or 7-day drug sales and mean raw or effluent water turbidity over same time steps</td>
<td>No correlation found between mean daily effluent water turbidity and drug sales. An increase of 10 NTU raw water turbidity corresponded to an increase in sales of 14% at a lag of 1 to 3 weeks. Breakdown in chlorination system associated with 19% increase in sales 3 to 8 days later</td>
<td>Design Unclear extent of overlap between outcome and exposure source populations – 20% sales to residents from other communities 3 and 7 day time step suggests low power Analysis Insufficiently detailed methods section Unclear and incomplete results Interpretation Insufficient discussion of results Insufficient discussion of bias</td>
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| Group 3 McConnell (2001) | 17 communities in 2 states in Australia (Victoria, South Australia) | Ecological Before and After analysis of improved water treatment | Upgrade of water treatment system | Requests for GI-related faecal specimens and midstream urine (MSU) samples to test for urinary tract infections as control from Australian Health Insurance Commission | Change in turbidity compared with change in risk of faecal specimen using Spearman rank test for correlations | No significant correlations found. | Design  
Low power (only 17 communities)  
Imprecise measure of turbidity and GI outcome  
No details as to sample size/power  
Inappropriate control group as age distribution different  
Analysis  
No analytical techniques presented  
No control for confounding  
Incomplete presentation of results  
No measure of effect or impact.  
Interpretation  
Conclusions not justified by results (trends in GI before upgrade and type of upgrade not taken into account.  
No discussion of bias  
No discussion of causality |
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<td>Egorov (2002)</td>
<td>15 areas of Cherepovets, Russia 22 October 1998 – 31 December 1999; 50 families per area were interviewed</td>
<td>Cross-sectional study</td>
<td>Water quality parameters such as colour (unspec. units), turbidity (formazin turbidity units FTU), free chlorine (mg/l) and iron (mg/l) measured at point of use and at sole water treatment plant</td>
<td>Self-reported diarrhoea or other GI symptoms such as vomiting or cramps lasting at least one day after at least two week symptom-free</td>
<td>Poisson GLM modelling using Generalized Estimating Equations (GEE) to account for clustering of observations from same household (give more robust standard errors) for association between water quality parameters and self-reported GI illness</td>
<td>Relative risk of GI illness for IQR change in turbidity 1.25 (0.93-1.68) or iron content 1.07 (0.84-1.37); exposure to colour not assessed. RR of GI illness for IQR change in free chlorine was 1.42 (1.05-1.91)</td>
<td>Note: primary objective was to assess association between residual chlorine levels and illness. Design: Unclear definition of exposure. Self-reported illness – lack of specificity. Unclear whether participants blinded – response bias. No details as to sample size/power. Analysis: Insufficiently detailed methods section. No analysis on colour carried out. No measure of impact calculated. Interpretation: Recommendations too strong given borderline results.</td>
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<tr>
<td><strong>Group 5</strong></td>
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<tr>
<td>Anon (1994)</td>
<td>Washington D.C. (US) November 22 – December 26</td>
<td>Before and After analysis of increased effluent turbidity at DC treatment plant 7 December</td>
<td>Being served by treatment works that experienced episode of increased turbidity</td>
<td>Self-reported loose/watery stools; A&amp;E diagnosis records; Nursing home survey of bowel movements and antidiarrheal medications; Microbiology lab survey</td>
<td>Calculated risk ratios for various outcomes using period before 8 December as baseline</td>
<td>All four surveys showed no statistically significant differences between the two periods</td>
<td>No discussion or interpretation.</td>
</tr>
<tr>
<td>First author, setting (year)</td>
<td>Participants</td>
<td>Study Design</td>
<td>Exposure</td>
<td>Outcome</td>
<td>Analysis</td>
<td>Results</td>
<td>Reviewers’ comments</td>
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<tr>
<td>Knobeloch, US (1998)</td>
<td>24 families in mobile home park, Wisconsin, US, 1996</td>
<td>Survey of health and copper levels initiated after complaints of blue water</td>
<td>“First draw” and “flushed” tap water copper levels (mg/l)</td>
<td>GI related illnesses ascertained through written questionnaire</td>
<td>Descriptive.</td>
<td>58% respondents reported “new” illness, i.e. after copper pipes fitted. Mean “first draw” copper level 1.2 mg/l; three samples exceeded federal action level. Mean “flushed” copper level 1.9; six samples exceeded action level.</td>
<td>Design Small study (38 respondents) No analysis of correlation between water copper levels and illness. No info on number of illnesses reported before copper pipes fitted</td>
</tr>
<tr>
<td>Rowland, UK (1990)</td>
<td>Residents and visitors in North Cornwall, UK around time of aluminium contamination of treatment works (6 July, 1988)</td>
<td>Retrospective cohort study including unexposed control area</td>
<td>Living in area served by Lowermoor treatment plant.</td>
<td>Self-reported symptoms (e.g. diarrhoea) in July and August 1988 (after contamination)</td>
<td>Descriptive and univariable analysis of association between living in exposed area and different types of self-reported symptoms</td>
<td>49.4% respondents in exposed area changed drinking habits around time of incident vs. 2.4% in control area 63% in exposed area noticed change in water, 51% of whom said colour changed; this was significantly higher than proportion of those in the control area who reported a colour change (p&lt;0.001) Relative risk of any symptom 4.2 (95% CI 3.3-5.4) in exposed compared to control area Relative risk of diarrhoea 5.1 (3.3-7.9) Non-significant relative risks when was restricted to those with symptoms (Relative proportional morbidity) Insufficient numbers for dose-response analysis with amount of water drunk</td>
<td>Design Self-reported illness – lack of specificity Unclear whether participants blinded – response bias Analysis No measure of impact calculated Significant confounding and bias in results likely.</td>
</tr>
</tbody>
</table>
Acknowledgements
The reviewers would like to thank the following people for responding to requests for information: Eugene Cloete, Lorna Fewtrell, Jamie Bartram, Arie Havelaar, Julian Dennis, Marty Allen, Donald Reid, Mark LeChevallier and Andrey Egorov. The reviewers would also like to thank Karen Stark for clerical assistance. Finally, the reviewers would like to thank Tony Lloyd and the Drinking Water Inspectorate of England and Wales for putting out the call for proposals, and accepting the application of this group.

Conflicts of interest
None declared.

References


Annex A  Sources searched and search strategies used.

Published Literature

Pubmed Search Strategy

Exposure:


Outcome:


Limit:

Human

Restrict to developed countries only:


Exclude studies stating developing, or middle-income, country (only) setting:


Embase Search Strategy

Exposure:

exp SEWAGE TREATMENT/ OR exp SEWAGE/ OR exp Microbiology/ OR exp Water Supply/ OR exp Water Management/ OR exp Water Pollutant/ OR exp NEPHELOMETRY/ OR exp Water Pollution/ OR exp CORROSION/ OR water treat$ OR nephelomet$ OR
turbid$ OR water source$ OR water filt$ OR boil$ water OR tap water OR water discolo$ OR water qualit$ OR water deteriorat$ OR potable OR corrosion by-product OR waste water

Outcome:

exp Gastrointestinal Disease/et, pc, ep [Aetiology, Prevention, Epidemiology] OR exp CRYPTOSPORIDIOSIS/et, pc, ep [Aetiology, Prevention, Epidemiology] OR exp CRYPTOSPORIDIIUM/ or exp CRYPTOSPORIDIIUM PARVUM/ OR exp GIARDIASIS/ep, et, pc [Epidemiology, Aetiology, Prevention] OR exp GIARDIA/ or exp GIARDIA LAMBLIA/ OR exp Virus Infection/et, pc, ep [Aetiology, Prevention, Epidemiology] OR exp BACTERIAL INFECTION/et, pc, ep [Aetiology, Prevention, Epidemiology] OR exp Intoxication/et, pc, ep [Aetiology, Prevention, Epidemiology] OR diarr$ OR stomach$ OR vomit$ OR waterborne OR water-borne OR water borne

Limit

Human

Restrict to developed countries only:

exp CANADA/ OR united states. OR exp United States/ OR exp JAPAN/ OR Japan OR Europe OR exp EUROPE/ OR exp "AUSTRALIA AND NEW ZEALAND"/ OR exp AUSTRALIA/ OR Australia OR new Zealand. OR exp New Zealand/

Exclude studies stating developing, or middle-income, country (only) setting:

Asia OR exp ASIA/ OR exp SOUTH ASIA/ OR Asia OR exp ASIA/ OR exp SOUTHEAST ASIA/ OR south America OR exp South America/ OR central America OR exp Central America/ OR exp AFRICA/ OR Africa OR Mexico OR exp MEXICO/

Aquatic Science and Fisheries Abstracts 3: Aquatic Pollution and Environmental Quality and Industrial and Applied Microbiological Abstracts (Microbiology A) combined search strategy
discolo* or turbid* or waterqualit* or drink*AND Gastrointestinal or diarr* (all free text terms)

**Unpublished Literature**

System for Information on Grey Literature in Europe (SIGLE) search strategy

discolo* or water quality or turbid* or drink* AND enteritis or gastrointestinal or diarr* or gastroenteritis or enteric

No structured search strategy was used for the following sources:

- Water Intelligence Online: (2001-2004 Reports and Conferences from International Water Association (IWA) and its Water Environment Research Foundation (WERF) and the American Water Works Association (Awwa))
- Health Canada
- Theses.com (UK/Ireland)
- Theses Canada
**Annex B** Form used to critically assess the papers included in the review.

**Water discolouration and GI illness**

**Aspects to consider in reviewing papers**

Paper ID #:

First Author:

**Design Issues**

1. Relevance to UK setting

2. Exposure definition:
   a. Clear definition of exposure (discolouration/turbidity)
   b. Appropriate definition of exposure

3. Outcome definition:
   a. Clear definition of outcome (GI)
   b. Appropriate definition of outcome

4. Study population:
   a. Clearly defined exposed (to discoloured water) population
   b. Appropriately chosen exposed population, eg. corresponds to population from which cases arise
c. Appropriate ascertainment of exposure

d. Clearly defined catchment population for outcome (GI)

e. Appropriate ascertainment of GI

f. Identical coverage of exposed and outcome population

5. Appropriate sample size/power (no. of people affected, analysis units, over time)

Where relevant to study design (e.g. exposed versus non-exposed areas, outbreak analysis):

6. Selection method of study (discolouration) population following event

7. Use of control area/group

8. Selection method of control area/group

9. Appropriateness of control/area group

10. Pre-event measurements of discolouration/turbidity (exposure baseline)

11. Pre-event measurements of GI (outcome baseline)

12. Extent of blinding (where appropriate) of subjects/investigators to study hypothesis
Analytical Issues

1. Appropriate statistical analyses

2. Account taken of delay between exposure and outcome (eg. incubation period or lags for time-series)

3. Account taken of seasonality/time trends

4. Account taken of other confounders (for time-series only time-varying factors)

5. Account taken of autocorrelation (eg. appropriate degree of smoothing, residual analysis)

6. Multiple testing

7. Account taken of baseline characteristics (eg. per-discolouration levels)

8. If relevant, appropriateness of dose-response analyses

9. Clear presentation of results

10. Crude analyses (eg. unadjusted/univariate)
11. Multivariate analyses (including interactions)

12. Details of missing data

13. Appropriate measure of effect/impact calculated

14. Appropriate confidence intervals/p-values

**Interpretation**

1. Conclusions justified by results

2. Consideration of bias/confounding

## Annex C Email responses received from authors and researchers in field of water research.

<table>
<thead>
<tr>
<th>Contact Name</th>
<th>Comment</th>
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<tbody>
<tr>
<td>Eugene Cloete</td>
<td>No readily available data or references at hand</td>
</tr>
<tr>
<td>Lorna Fewtrell</td>
<td>Not aware of anything on discoloured water</td>
</tr>
</tbody>
</table>
| Jamie Bartram         | Two recent headlines:  
2. Launch of third edition of WHO Guidelines for Drinking-water Quality (also accessible through the general WHO water and sanitation link above and “drinking water quality”)  
Also noted:  
Biological plausibility is self evident.  
Mark Lechevalier’s piece on correlations with turbidity.  
Would imagine an issue is how to make the association (temporal and spatial relationships) – there could be visibly apparent quality changes due to central (system-wide), in-distribution and household; the fact that many may be short lived; those of health significance versus those not; and a psychosomatic effect.  
Jack Colford and Lorna Fewtrell with World Bank and WHO completed exhaustive search on all health impact studies on water/sanitation/hygiene and proceeded to meta analysis. |
| Arie Havelaar         | No experience of this kind in the Netherlands                                                                                                                                                   |
| Julian Dennis         | Will circulate and see what sort of response they get.                                                                                                                                         |
| Martin Allen          | Forwarded message to Research Management Group                                                                                                                                                  |
| Donald Reid           | Never seen any plausible published evidence.                                                                                                                                                |
| Mark LeChevallier     | No data on discoloured water quality, although pressure changes could cause suspension of settled water. (Attached document entitled: The Potential for Pathogen Intrusion During Pressure Transients.pdf) |
| Andrey Egorov         | 1. Informed reviewers of two additional papers to do with Milwaukee cryptosporidiosis outbreak (1993)  
2. Forwarded email to Anne Seeley of the New York Bureau of Water Supply – who recently analyzed data on turbidity and gastroenteritis in the city of New York.  
3. Informed reviewers of current/forthcoming research he is conducting in this area. |
Proposal for a study investigating any potential relationship between the supply of discoloured water and gastrointestinal illness

Background

There has been some speculation regarding whether drinking water that meets legal requirements for quality can cause acute gastrointestinal illness (GI illness). A body of research exists addressing this issue, mainly investigating the effect of variations in turbidity at point of treatment and the use of filters at point of use on the incidence of GI illness. There has also been public health interest in whether events resulting in sudden obvious discolouration of the water supply could result in an increase in the incidence of GI illness among those exposed; the review phase of this study failed to find any studies that specifically addressed this issue. Discolouration of tap water can be caused by inadequately treated water entering the distribution system, but is more often associated with the disturbance of sediment, particularly corrosion products, within the distribution systems. The possibility of poor control within the distribution system may coincide with presence and survival of gastrointestinal pathogens such as Salmonella, Campylobacter, Cryptosporidium, Giardia or toxigenic strains of E. coli and Shigella that transiently pass through the system and are not detected during routine monitoring of grab samples.

There were 44 discolouration events in England and Wales during 2003 that were notified to the DWI and classified as ‘incidents’, affecting approximately 1.4 million consumers (1). A similar number of events were notified to DWI but classified as ‘non-incidents’ (for definition of incidents and non-incidents see ‘Definition of Exposure’ section below). Thus, water discolouration is a relatively common occurrence: even a small relative increase in the risk of GI illness may be of considerable public health importance. No studies have been conducted to investigate specifically whether events of water discolouration cause an increase in GI illness. There have been studies on the variation of parameters of water quality (such as turbidity), but these studies mostly measured parameters at the treatment plants and rarely in the distribution zone. The effect on GI illness of changes in turbidity or discolouration occurring after the water has left a plant has not been studied rigorously before. The
The vast majority of discolouration events in England and Wales (E&W) occur in the supply zones rather than at the treatment plants (personal communication, Annex 5). In order to address the question of whether water discolouration events cause increases in the incidence of GI illness, an epidemiological study is needed. We propose here a study to investigate the effect of water discolouration events (notified to the Drinking Water Inspectorate (DWI) and irrespective of whether they are subsequently classified as incidents or non-incidents) on the incidence of acute GI illness in the affected populations.

**Overview of project structure**

The study will consist of two separate components. The first component will be a study of water discolouration events (incidents and non-incidents) and their effect on the incidence of acute GI illness as assessed by routinely collected health data. This analysis will be performed using historical data from past discolouration events. Consideration was given to whether this part of the study would need to be extended to include future events if, after examination of reports of all past events, it was apparent that existing data are not of sufficient quality for analysis, or that the number of past events with good quality information is too small to provide sufficient statistical power. However, following examination of a number of incident and non-incident reports and discussions with six water companies, it was concluded that the data held on historical events would be of sufficient quality to address the primary aims of the study. The proposed retrospective study also includes an analysis of events of depressurization, for which the data availability and quality are similar to events of discolouration.

In the second study component, a series of telephone surveys is proposed in areas affected by new water discolouration events and appropriately selected unaffected areas. The difference in incidence of reported acute GI illness between affected and unaffected communities will then be estimated. Because the telephone surveys will be conducted soon after the event, it will be possible to assess awareness of the event and consequent behavioural changes in water consumption.
Part I: Study of discolouration events using routine health databases

Aims and objectives

The main aims of the study are to determine (1) whether events of water discolouration lead to an increase in the incidence of acute GI illness in the affected population, and (2) whether specific features of events (for example chemistry, turbidity) and of affected water supply networks influence the risk of acute GI illness following an event.

Primary study objectives

1. To determine whether events of water discolouration in a water supply zone lead to an increase in the incidence of acute GI illness in the affected population.

2. To determine whether events of water depressurisation in a water supply zone lead to an increase in the incidence of acute GI illness in the affected population.

3. To determine if any characteristic of water discolouration (e.g. cloudy vs. brown discolouration, presence of iron/manganese/aluminium, turbidity level) or characteristics (e.g. age of the system, material of pipes, type of soil, renewal efforts, rural/urban/metropolitan setting, social deprivation) of the water supply zone affected by an event influence any subsequent risk of acute GI illness.

4. To determine whether any characteristics of the water supply zone affected by a depressurisation event influence any subsequent risk of acute GI illness.

5. To determine the extent to which events of discolouration are associated with depressurisation of the water supply, and if possible, examine the separate and combined effects of these on the risk of GI illness.

6. To estimate the proportion of acute GI illness in England and Wales, as reported by routine health data sources, that is attributable to water discolouration and depressurisation events, i.e. the population attributable fraction (PAF).
Secondary study objectives

7. To assess whether there is a characteristic delay (lag) between exposure to discoloured (or depressurised) water and any subsequent peak in the incidence of GI illness, which might point to specific aetiologies.

8. To explore whether the effect of a water discolouration and/or depressurisation event on acute GI illness is more pronounced in certain age groups, in particular children and the elderly.

Study design and methods

The study aims will be investigated using a retrospective cohort drawn from the population of England and Wales during the years 1999 to 2003. The cohort will comprise residents of households in areas affected by water discolouration (and depressurisation) events (all events notified to the DWI, irrespective of whether DWI classify them as incidents or non-incidents) during this time period. The incidence of acute GI illness in these areas in the two weeks following an event will be compared with that in appropriately selected unaffected (control) areas over the same time period. In addition, the change in incidence of acute GI illness at various time points following an event will be estimated in order to identify characteristic lag times between exposure and illness.

Choice of outcome

Episodes of acute GI illness will be ascertained from two different sources: hospital admissions and calls to the NHS Direct Telephone Service. Data on hospital admissions will be obtained from the Hospital Episode Statistics (HES) Database (England) and the Patient Episode Database Wales (PEDW). Data on calls to NHS Direct will be obtained from the NHS Direct database. These data sources were selected because they both provide widespread geographic coverage, geographic location at seven-digit postcode level, and precise timing. The two databases capture different aspects of GI illness - calls to NHS Direct are likely to include the most common, milder episodes, whereas hospitalisations are likely to result from more severe disease. We outline the characteristics of these data sources below.
HES/PEDW

HES records details of all admissions to National Health Service (NHS) hospitals in England and Wales according to the ICD-10 code of an illness or symptom/sign. Its Welsh equivalent is PEDW. Information available on admissions includes patient age and gender, postcode of residence, primary diagnosis, date of admission, and length of hospitalisation. For the purposes of the study, the following ICD-10 codes will be considered, as they are the most likely to capture acute GI illness: A00-A09.9 (diarrhoea of infectious or presumed infectious origin); K52.9 (non-infective gastroenteritis and colitis, unspecified); R11 (nausea and vomiting). Cases will be included in the study if they are hospitalised with any of these conditions as the primary diagnosis and if they resided in an affected area (as determined by their postcode) in the two weeks following an event or in a pre-defined control area during the same time period.

HES/PEDW data have good geographic resolution and good resolution in time. Of particular interest, they are robust to bias; hospitalised cases are more severe, making it less likely that hospitalisation could be influenced by patients’ perception of risk if, for example, they were aware that their water supply was discoloured. This is in contrast to NHSdirect (see below).

NHS Direct

NHS Direct is a telephone service provided by the NHS since May 2001 with the aim of providing advice to individuals prior to seeking medical advice in healthcare facilities. Information is available on calls related to the occurrence of 10 specific conditions, including diarrhea and vomiting. The age, sex and seven-digit residential postcode of callers who do not wish to remain anonymous are also available. Thus, it is possible to use NHS Direct as a measure of disease occurrence (for which health advice is sought) in the community. NHS Direct data have good geographic and temporal resolution. A limitation of NHS Direct data is the potential for bias: cases recorded by NHS Direct are a self-selected group of cases, in that they represent those cases who have GI illness and have decided to call
NHS Direct. If people who know that they have been exposed to discoloured water are more likely to call NHS Direct, this could lead to an observed increase in GI illness that did not necessarily reflect a true increase in disease incidence.

Although neither of these data sources provide complete ascertainment of all GI illness in the community - they only capture GI illness either resulting in hospitalisation or for which telephone advice was sought - they should be sufficiently consistent to be good indicators of changes in incidence of mild and severe GI illness over time.

Other sources of routine data on GI illness have been assessed for use in the proposed study, but have been deemed to be inadequate, either due to poor geographic resolution or limited geographic coverage. These include the General Practitioner Research Database (GPRD), over-the-counter sales of diarrhoea-related drugs, laboratory reports of pathogens routinely isolated from stool samples, notifications of food poisoning and reports of disease outbreaks. The grounds on which these data sources were excluded are summarised in Annex 1.

**Definition of exposure**

The main exposure under study is discolouration classified as an event occurring within geographic areas defined by affected water supply zones. Under the Water Undertakers (Information) Direction 1998, water companies are required to report as soon as possible to the Drinking Water Inspectorate (DWI) any event which gives rise, or is likely to give rise, to a significant risk to the health of consumers. All events must be confirmed in writing within 72 hours and an assessment of this initial report is carried out by the DWI within five days of receipt. Events of a particular nature, including adverse water quality changes (e.g. discolouration) as perceived by consumers, are classified by the DWI as an ’incident’ and a full report must be submitted by the water company to the DWI within 30 days. We propose that in this study all events reported to the DWI since 1999 (the first full year in which the Direction came into force defining an incident) and affecting at least 1000 people (for statistical power and applicability/generalisability of results) are included, irrespective of whether they
are classified as an incident or non-incident. Events not classified by DWI as incidents (non-incidents) may include changes to the water supply that are less perceivable (e.g. less visible) to the consumer. A potential advantage of including such events is that the potential for bias (resulting from people modifying their water consumption behaviour) could be reduced. Although only a ‘72-hour’ report is available for each non-incident the quality of data relevant to the primary objectives of the proposed study is comparable to that of incidents (a number of non-incidents may have insufficient data on the characteristics of the event (objective 3) because a full report is not available but this is likely to be a small proportion of all non-incidents). See annex 6 for more details regarding the inclusion of non-incidents.

A database of all incidents and non-incidents under study will be produced. The variables in the database will include parameters characterising each event, e.g. cloudy or brown discolouration, level of iron, manganese, aluminum and turbidity, ground or surface water, number of consumer complaints. Data regarding individual events will be abstracted from the 72-hour and 30-day reports and entered into the database. Geographic information on postcodes affected by the event will be extracted from the geographical information system (GIS) held by each company (see annex 6). An exposed area will be defined as a group of postcodes affected by an event at a certain time as identified by the water company together with a dedicated researcher. The geographic boundaries of the exposed areas will be defined by mapping the postcodes affected by an event to census Output Areas (OAs), the smallest geographical area for which basic demographic data have been aggregated. These are statistical units of geography largely defined on the basis of geographic proximity and social homogeneity (2). All 2001 census data are available at the level of census OAs with a population size of approximately 300 individuals. Thus, the 2001 census data on the exact population size, age structure, female/male ratio, social deprivation and ethnicity can be linked to OAs. Although these data are only available for the 2001 census year, we are confident that they can be used to characterise affected and control areas throughout the study period, as socio-demographic measures are unlikely to have changed materially over such a short time. Since tables specifying the OA to which a postcode belongs are available, routine health data at 7-digit postcode level can be linked to
each OA, numbers of cases of GI illness from routine sources can be related to the population in the areas, and the incidence of acute GI illness in exposed and unexposed areas estimated and compared. The misclassification introduced by the use of OAs rather than 7-digit postcode is likely to be minimal, particularly for large events, and can be examined by conducting sensitivity analyses (e.g. by excluding smaller events from the analysis). See Annex 6 for further discussion of exposure misclassification.

**Selecting a control area for each exposed area**

In order to obtain potential control areas of sufficient size, England and Wales will be divided into areas resulting from the combination of adjacent OAs, comprising between 1,000 and 50,000 individuals (between 3 and 160 OAs). For events affecting more than 50,000 people, OAs will be aggregated further. Control areas for each individual event will be selected from these pre-defined areas. In order to control for differences in the baseline (pre-event) level of acute GI illness, control areas will be matched to exposed areas with respect to the incidence of acute GI illness in a defined period before the event. For each half-year (Jan-Jun or Jul-Dec) between July 1998 and December 2003, the incidence of GI illness will be calculated for every pre-defined control area. These estimates will be used to construct sets of 11 separate maps of England and Wales (one per half-year) categorising control areas by incidence of GI illness. A separate set of GIS maps will be developed for each data source used (hospitalisations and NHS Direct calls).

The matching of a control area to an exposed area will be based on the incidence of GI illness before the event. For each exposed area, the incidence of acute GI illness in the preceding six full calendar months (Jan-Jun or Jul-Dec) will be calculated. For example, for an event taking place in September 2000, the cumulative incidence of acute GI illness will be calculated for January to June 2000 separately for hospitalisations and calls to NHS Direct.

Each exposed area will then be matched to a pre-defined control area based on geographic proximity and baseline incidence of acute GI illness using the map appropriate to the time period of the incident
in the exposed area. In order to avoid exposure misclassification in control areas, a safety boundary will be constructed around each exposed area. A potential control area must lie outside this boundary to minimise the likelihood that (a) the control area will be served by the same water supply zone as the exposed area, and (b) that individuals residing in the control area will not consume water in the exposed area (e.g. which might occur if, for example, their place of employment were in an exposed area).

**Estimating incidence of acute GI illness in pairs of exposed and control areas after an event**

Incidence of acute GI illness in each pair of exposed and control areas will be estimated separately for hospitalisations and NHS Direct calls for the two weeks following an event. This will be done by linking cases to areas using their postcode of residence and dividing the number of cases by the population of the area. Incidence estimates will be calculated overall, as well as by sex and age group.

**Statistical analysis**

The analysis will compare the incidence of acute GI illness in affected areas and in control areas; the estimate produced by this analysis is the rate ratio (RR) - the relative increase in rate of acute GI illness. The analysis will control for incidence of acute GI illness in the affected and control areas before the event and any relevant census characteristics of the areas. Several analytical approaches would be appropriate to compare rates of acute GI illness in exposed and control areas following an event (Objectives 1 and 2). A simple meta-analysis, Poisson model, or Random Effects Model (3) would all provide an estimate of the rate ratio (RR), while taking account of the fact that each exposed area is individually paired with a control area. The adequacy of each model will be assessed by examining the goodness of fit and the choice of which approach to adopt will be based on this formal evaluation. Each of these approaches can be extended to the multivariate situation in which the confounding and modifying effects of co-factors, (Objectives 3, 4, and 5) and the age of the population affected (Objective 8) are examined. Ninety-five percent confidence intervals (95% CIs) and significance tests will be produced. The proportion of cases (the population attributable fraction, or PAF) of acute GI illness occurring in England and Wales that can be attributed to discoloured (or
depressurised) water would be calculated on the basis of the RR calculated from objective 1 and 2 and the estimated proportion of people affected each year by an event (Objective 6). If an increase in acute GI illness is found in areas exposed to a water discolouration (or depressurisation) event (Objective 1), then analyses will be carried out to examine whether there is a characteristic time lag after which a peak in incidence occurs (Objective 7).

**Sample size calculation**

The following sample size calculations refer to events of water discolouration, but apply equally to depressurisation events, since an equal number of these would be available for analyses.

According to the DWI Website (1), a total of 310 water discolouration incidents occurred between 1999 and 2003 (the annual number of incidents has decreased steadily from 95 in 1999 to 44 in 2003). Approximately 90% of these incidents affect more than 1000 people, so the maximum number of discolouration incidents available for a retrospective analyses is likely to be approximately 280. Approximately 50% of discolouration events reported to the DWI are classified as non-incidents, so a similar number (280) of non-incidents (affecting at least 1000 people) would be available, providing a total of 560 events in total. However, as stated in the ‘Choice of outcome’ section earlier, NHS Direct has only been in operation since May 2001. Therefore, only incidents and non-incidents since that date can be included in an analysis of NHS Direct calls, of which there have been approximately 240 discolourations in total.

Assuming the number of incidents and non-incidents continues to decline to a plateau over forthcoming years, then the number of future events per year is likely to be, on average, around 60 (30 incidents and 30 non-incidents) in total.

Based on the 1999-2003 incident and non-incident data, the mean number of people affected per incident is approximately 18,000.
**Background incidence in HES/PEDW**

According to the 2000 and 2001 HES/PEDW databases the number of NHS hospital admissions in England and Wales due to acute GI illness (as defined by the ICD-10 codes described earlier) is around 70,000 per year. This equates to approximately 5 admissions per 100,000 individuals during a two-week period.

**Background incidence in NHS Direct**

Based on NHS Direct calls from 1st Aug 2003 to 31st July 2004, at least 14 calls due to diarrhoea and 20 calls due to vomiting can be expected in a two-week period.

The table in Annex 2 shows the number of events that would be required in order to detect a range of specified rate ratios, reflecting the increase in risk of GI after an event. The following assumptions are made: (i) the coefficient of variation between clusters is $k=0.5$, (ii) a power of 80% and significance level of 0.05 is required.

If all (560 for HES/PEDW and 240 for NHS Direct) retrospective incidents and non-incidents were available for analysis, there would be 80% power to detect a rate ratio of approximately 1.2 for HES/PEDW and 1.18 for NHS direct. A rate ratio of 1.2 for HES/PEDW represents a 20% increase in the number of admissions in exposed areas during the two weeks following an event. Thus, if the baseline rate of hospitalisations for acute GI illness in control areas is 5 per 100,000 (as stated above), then there would be approximately 504 expected admissions in all of the control areas combined (from 560 areas of 18,000 people = 10,080,000 people) and 605 admissions (an extra 20% = 101 admissions) in areas exposed to discoloured water.

Up to 20% of retrospective events may not be available for analysis because of practical limitations of collecting the data or difficulties in identifying the affected population. If only 80% of events during 1999 to 2003 were available for analyses then the retrospective study would still have 80% power to
detect an increase in hospital admissions of approximately 22% (compared to 20% if all events were available for analysis) and an increase in NHS Direct calls of 20% (compared to 18%).

The extent to which only readily available retrospective events could be used to examine whether specific characteristics of water discolouration events are responsible for increases in acute GI illness (Objective 3) is dependent on the frequency with which the characteristic occurs. For example, approximately 70% of discolouration events are classified as “brown”, so there would be 392 retrospective events of “brown” discolouration available for HES/PEDW analysis and 168 events for NHS Direct. This would provide 80% power to detect a 25% increase (RR=1.25) in hospital admissions and a 22% increase (RR=1.22) in NHS Direct calls.

Although the inclusion of prospective events over future years would increase the statistical power of the study, the detectable rate ratios would be reduced only slightly. Continuation of the study for a further 5 years would add approximately 300 extra events into the analysis, and an increase in hospital admissions of 16% (rather than 20%) and NHS Direct calls of 12% (rather 18%) would be detectable if all 300 prospective events were analysed together with the retrospective events.

**Decision to include future incidents into the study**

The inclusion of events occurring after the start of the study would be necessary if the number of past events (or the population affected by past events) with adequate geographic information (i.e. information of sufficient quality to obtain a precise definition of exposed areas) is too small or the information regarding the events themselves as provided by DWI/water companies is insufficiently detailed to provide adequate statistical power to detect the desired magnitude of effect on GI illness. Detection of smaller increases in risk require larger sample sizes. In this case, the study can be extended to include future events. More precise geographic and other information for future events may be obtainable by increasing the degree of sampling following an event and increasing the degree of customer contact. However, the practical limitations (e.g. costs) of increased sampling and increased customer contact, together with the modest increase in statistical power (as stated in the
sample size section above) suggest that inclusion of prospective events over a five-year period may be of limited value. Inclusion of prospective events may be of greatest relevance for analyses of NHS Direct calls (since only 240 retrospective events are available) and examination of whether specific characteristics of events are responsible for increases in acute GI illness (objective 3). For example, in a group of 300 prospective events there are likely to be approximately 200 in which the water is classified as “brown”. The analysis would then have 80% power to detect a 20% increase in hospital admissions of acute GI illness related to “brown” water events (compared with 25% if only retrospective events were included).

The methods of selecting exposed and control areas would be the same as used in the retrospective analysis. Since the annual number of future events is uncertain, but likely to decline, the timeframe of the prospective assessment of events is not entirely certain (see sample size calculation).

### Part II: Survey of the effect of new discolouration incidents on water consumption and on community-level acute GI illness

**Aims of the study**

The first part of the study described above will investigate whether water discolouration events have an effect on the incidence of acute GI illness - as measured by hospitalisations and calls to NHS Direct - that is detectable at a population level. Although geographic information will be used to define exposed areas with the greatest possible precision, the study does not explore whether an absence of increased risk after an event is associated with the discoloured water being safe or an avoidance of water consumption by consumers because of awareness of discolouration. Data on consumption of water are not routinely available at individual level. The proposed second phase of the study will address these issues by using telephone surveys of exposed and control areas following an event in order to collect information on changes in water consumption. This information will be used to interpret findings arising from Part I of the project. This second phase will also investigate whether discolouration events have an effect on the incidence of milder GI illness as recalled by individuals. A
limitation of such telephone surveys is that they are subject to reporting bias as awareness of the
discolouration event can influence whether the respondent will consider as diarrhoea mild changes in
frequency and/or consistency of stools that are unrelated to acute GI illness. Thus, data from both the
telephone surveys and NHS Direct are more susceptible to bias than data from hospitalisations. An
advantage of using telephone surveys, however, is that they allow for investigation of any effect of
discolouration events on mild acute GI illness (as opposed to severe illness obtained using
hospitalisation data).

Study objectives

1. To estimate the effect of discolouration events on mild acute GI illness in the community.
2. To calculate the population attributable fraction (PAF) of mild acute GI illness caused by water
discolouration events
3. To study the effect of water discolouration on water consumption behaviour and to determine
whether changes in drinking behaviour can explain any effect on frequency of diarrhoea.

Study design and methods

Because each survey would need to be conducted a short time after the event, the timeliness of
notification and information exchange is crucial. This will require close collaboration between the
water companies, the DWI and the study team. Water companies will be requested, through the
Drinking Water Inspectorate, to provide accurate geographic information on the area affected by an
event within 10 days of it being declared. Once geographic information is obtained, exposed and
control areas will be determined by the same methods as used in the retrospective component.

Data collection: telephone surveys

Two weeks after each event a telephone survey in the exposed and control areas will be conducted for
a duration of seven days. During this time, a number (sample size calculations are shown for 500 and
1000) of randomly selected households in each exposed and control area will be contacted. In the UK,
approximately 95% of private households have a fixed line\(^4\). Established procedures exist for using
postcode information to carry out random digit dialing of residential telephone numbers within a geographical area (MORI). One option is to subcontract the telephone survey to an organisation such as MORI who are established in carrying out such large volume telephone surveys over a short time. Mobile telephone numbers will not be included. Individuals selected for an interview will be asked questions including those on symptoms of acute GI illness (using procedures established for the assessment of diarrhoea in the community (4)) following an event, frequency and recent changes in tap water consumption and whether they noticed anything unusual about their water in the last few weeks.

Statistical analysis

Analysis similar to that proposed in Part I of the project will be carried out to compare incidence of mild acute GI illness in exposed and control areas (Objective 1) and also to calculate the proportion of mild acute GI illness in the community that is attributable to events of water discoulouration (Objective 2). Descriptive analyses will be carried out of reported changes in water consumption, reported awareness of the discoulouration event and reported frequency of mild acute GI illness in the two weeks preceding the interview (Objective 3). A comparison of the incidence of mild acute GI illness in exposed and control areas will then be conducted taking into account awareness of the event and water consumption.

Sample size

The incidence rate of diarrhoea in the community is estimated to be around 194 per 1000 person years (6). This approximately corresponds to a cumulative incidence risk of 8 cases per 1000 individuals in a two-week period.

Table 2 in Annex 2 shows the number of prospective events (incidents and non-incidents) that would be required to detect specified rate ratios using a telephone survey. Five years of prospective events (i.e. 300 in total) would provide 80% power to detect rate ratios between 1.15 and 1.2 depending on the number of successful calls made per event. If only two years of prospective events were analysed...
(120 events) then a rate ratio of 1.3 would be detectable with 80% power (assuming 500 calls were made in exposed and control areas) for both HES/PEDW and NHS Direct.
Timeframe of study

Retrospective part:

<table>
<thead>
<tr>
<th>Preparatory phase</th>
<th>Months</th>
</tr>
</thead>
<tbody>
<tr>
<td>Planning and recruitment</td>
<td>3</td>
</tr>
<tr>
<td>Ethic and confidentiality approval PEDW , HES, NHSdirect databases</td>
<td>3</td>
</tr>
<tr>
<td>Acquiring GIS system and training</td>
<td>4</td>
</tr>
<tr>
<td>Acquiring HES, PEDW, NHSdirect, and census data</td>
<td>6</td>
</tr>
<tr>
<td>Liaison with DWI and water industry to identify all potential past events and acquire postcode data of areas affected</td>
<td>11</td>
</tr>
<tr>
<td>Abstracting information on past events (including characteristics of events) from 7 day and 30 days reports and creating the database on past events, and entering data on database</td>
<td>9</td>
</tr>
<tr>
<td>GIS mapping of England and Wales into potential control areas based on SOAs (3 months)</td>
<td>3</td>
</tr>
<tr>
<td>Linking incidence of acute GI (separately for hospital data and from NHS direct data) to all potential control areas and estimation of incidence by age and sex in 6 months periods in control areas</td>
<td>3</td>
</tr>
<tr>
<td>Linking postcodes of areas affected by each event to geographical areas in the GIS system. Definition of exposed areas. Estimate of incidence of GI in the 6 months prior to event. Transfer to the event database</td>
<td>5</td>
</tr>
<tr>
<td>Selection of a control area for each exposed area.</td>
<td>2</td>
</tr>
<tr>
<td>Linking incidence of acute GI (separately for hospital data and from NHS direct data) to each pair of exposed area and control area and estimation of incidence by age and sex in the 2 weeks after the event. Transfer to the event database.</td>
<td>2</td>
</tr>
<tr>
<td>Abstracting census data on characteristics of exposed areas and control areas and transferring to the event database.</td>
<td>2</td>
</tr>
</tbody>
</table>

Analytical Phase

| Analysis of primary study questions (for both discolouration and depressurisation events) | 10     |
| Analysis of secondary study questions (for both discolouration and depressurisation events) | 7      |
| Interpretation and writing up phase: Writing report to DWI, discussion with DWI, preparation of final report and writing manuscripts for publications | 6      |
Prospective part:

<table>
<thead>
<tr>
<th>Preparatory phase</th>
<th>Months</th>
</tr>
</thead>
<tbody>
<tr>
<td>Planning and recruitment</td>
<td>3</td>
</tr>
<tr>
<td>Ethic and confidentiality approval PEDW, HES, NHSdirect databases</td>
<td>3</td>
</tr>
<tr>
<td>Acquiring GIS system and training</td>
<td>4</td>
</tr>
<tr>
<td>Acquiring HES, PEDW, NHSdirect, and census data</td>
<td>6</td>
</tr>
<tr>
<td>Liaison with DWI and water companies to establish procedures for identifying</td>
<td>6</td>
</tr>
<tr>
<td>events and notifying LSHTM</td>
<td></td>
</tr>
<tr>
<td>Liaison with telephone survey company to establish procedures for responding to</td>
<td>6</td>
</tr>
<tr>
<td>incident.</td>
<td></td>
</tr>
<tr>
<td>GIS mapping of England and Wales into potential control areas based on SOAs</td>
<td>3</td>
</tr>
<tr>
<td>(3 months)</td>
<td></td>
</tr>
<tr>
<td>Linking incidence of acute GI (separately for hospital data and from NHS direct</td>
<td>3</td>
</tr>
<tr>
<td>data) to all potential control areas and estimation of incidence by age and sex</td>
<td></td>
</tr>
<tr>
<td>in 6 months periods in control areas</td>
<td></td>
</tr>
<tr>
<td>Accrual of prospective events. Within 2 weeks following event: (I) Notification</td>
<td>24</td>
</tr>
<tr>
<td>of event by DWI to LSHTM, (ii) selection of a control area, (iii) notification by</td>
<td></td>
</tr>
<tr>
<td>LSHTM to telephone survey company of postcodes affected by event and postcodes</td>
<td></td>
</tr>
<tr>
<td>of potential control households, (iv) conduct of telephone interview by company.</td>
<td></td>
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<td></td>
<td></td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Analytical Phase</th>
</tr>
</thead>
<tbody>
<tr>
<td>Analysis of primary study questions</td>
</tr>
<tr>
<td>Analysis of secondary study questions</td>
</tr>
<tr>
<td>Interpretation and write up phase: Writing report to DWI, discussion with DWI,</td>
</tr>
<tr>
<td>preparation of final report and writing manuscripts for publications</td>
</tr>
</tbody>
</table>
Ethical considerations

HES and PEDW require approval by the Security and Confidentiality Advisory Groups responsible. This process will take approximately 2 to 3 months. The NHS Direct Clinical Governance Committee gives the final decision on approval of any studies hoping to use NHS Direct data. This will have to be achieved separately for England and Wales. NHS Direct also requires approval by an appropriate Research Ethics Committee, before research activity may commence. Part II of the project will also require approval by a Research Ethics Committee.
Annex 1: Data sources excluded from the proposal and the reasons for exclusion:

- The General Practitioner Research Database (GPRD). Only 5% of the population of England and Wales is covered by this data source. The geographic resolution is poor (cases are allocated to large areas for confidentiality reasons). Detailed information of acute GI cases can only be obtained with the approval of all GPs responsible for providing care. This would be unfeasible in the time frame required.

- Sales of over-the-counter medications. These data can be acquired only at regional level. Sales from wholesalers to pharmacies are available for 4 to 5 letter postcodes, but the resolution in time is poor (sales per calendar month only).

- Laboratory reports of isolates in routine stool samples. These represent only a small fraction of incident acute GI illness and do not include non-microbiological causes. Temporal resolution is not very good as date of disease onset is rarely available, and geographic resolution is poor.

- Notifications of food poisoning are of limited use for this study, as the geographic information provided is imprecise and date of illness onset is not available at national level.

- Reports of disease outbreaks contain information about the characteristics of outbreaks, but no information at the individual level. Thus, geographic information on cases affected is of insufficient quality.
Annex 2: Sample size calculation

The following formula (Bennett and Hayes) has been used to determine the number of incidents required to detect specified rate ratios:

\[ c = \frac{2 + f[p_0(1 - p_0)/m + p_1(1 - p_1)/m + k^2(p_{0}^2 + p_{1}^2)]}{(p_0 - p_1)^2} \]

where \( c \) is the number of water discolouration events (clusters) required (an equal no. of unexposed areas is also required), \( p_0 \) is the cumulative incidence risk in the unexposed area, \( p_1 \) is the incidence risk in the exposed area, \( m \) is the number of individuals in each cluster, and \( k \) the coefficient of variation between clusters; \( f \) equals 7.84 for a power of 80% and alpha of 0.05.

Table 1. Sample size (number of events) required to detect specified Rate Ratios using HES/PEDW, NHS Direct.

<table>
<thead>
<tr>
<th>Relative Risk</th>
<th>HES</th>
<th>NHSdirect</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.1</td>
<td>3480</td>
<td>704</td>
</tr>
<tr>
<td>1.2</td>
<td>558</td>
<td>192</td>
</tr>
<tr>
<td>1.3</td>
<td>263</td>
<td>151</td>
</tr>
<tr>
<td>1.5</td>
<td>176</td>
<td>93</td>
</tr>
<tr>
<td>2.0</td>
<td>58</td>
<td>16</td>
</tr>
</tbody>
</table>
Table 2. Sample size (number of events) required to detect specified Rate Ratios using a telephone survey.

<table>
<thead>
<tr>
<th>Relative Risk</th>
<th>500 calls*</th>
<th>Years required</th>
<th>250 calls*</th>
<th>1000 calls *</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.1</td>
<td>843</td>
<td>10+</td>
<td>1251</td>
<td>639</td>
</tr>
<tr>
<td>1.2</td>
<td>228</td>
<td>5-10</td>
<td>300</td>
<td>174</td>
</tr>
<tr>
<td>1.3</td>
<td>110</td>
<td>3-4</td>
<td>143</td>
<td>85</td>
</tr>
<tr>
<td>1.5</td>
<td>47</td>
<td>2</td>
<td>60</td>
<td>37</td>
</tr>
<tr>
<td>2.0</td>
<td>17</td>
<td>1</td>
<td>21</td>
<td>15</td>
</tr>
</tbody>
</table>

*Number of calls required in both the exposed and control areas
Annex 3: Water Discolouration and GI Illness: Timetable of tasks for retrospective analyses

- Planning and recruitment
- Ethics and confidentiality
- Acquisition of GIS
- Acquisition of HES, PEDW, and NHSdirect
- Acquisition of census data
- Liaison with DWI and water companies
- Creation of incident database
- Use of GIS to divide E&W into potential control areas
- Linkage & calculation of 6-month GI incidence in all control areas
- Use of GIS to identify geographical area of each incident
- Linkage & calculation of 6-month GI incidence in exposed areas
- Selection of control areas (matched to exposed areas)
- Linkage & calculation of 2-week GI incidence (exposed & control areas)
- Linkage to census database
- Analysis of primary study questions (discolouration and depressurisation)
- Analysis of secondary study questions (discolouration and depressurisation)
- Interpretation and writing up
Annex 4: Water Discolouration and GI Illness: Timetable of tasks for prospective part

1 January 2006

31 January 2006

2 March 2006

1 April 2006

2 May 2006

1 June 2006

1 July 2006

31 July 2006

31 August 2006

30 September 2006

30 October 2006

29 November 2006

30 December 2006

29 January 2007

28 February 2007

30 March 2007

30 April 2007

30 May 2007

29 June 2007

29 July 2007

29 August 2007

28 September 2007

28 October 2007

27 November 2007

28 December 2007

27 January 2008

26 February 2008

27 March 2008

27 April 2008

27 May 2008

26 June 2008

26 July 2008

26 August 2008

25 September 2008

25 October 2008

24 November 2008

25 December 2008

24 January 2009

23 February 2009

25 March 2009

25 April 2009

25 May 2009

24 June 2009

24 July 2009

24 August 2009

23 September 2009

23 October 2009

22 November 2009

23 December 2009

Planning and recruitment
Ethics and confidentiality
Acquisition of and training in GIS
Acquisition of HES, PEDW, and NHSdirect
Acquisition of census data
Liaison with DWI and water companies
Liaison with telephone survey company
Use of GIS to divide E&W into potential control areas
Calculation of 6-month GI incidence in potential controls areas
Accrual of all prospective events
Analysis of primary study questions
Analysis of secondary study questions
Interpretation and writing up
Annex 5: Personal communication with Mr Peter Marsden

1) The vast majority of discoloured water incidents will be identified through customer complaints not through the spot sampling done to comply with the regulations.

2) The vast majority of discoloured water incidents are associated with distribution rather than treatment works.

3) We do not hold postcode data for the areas affected by incidents, just the maps/descriptions from the company 30 day reports. However I am reasonably confident we can get companies to provide this information should a full study go ahead. If necessary we can revise the requirements. Do not contact the companies at this stage, DWI will do so should a full study go ahead.

4) We discussed the need to include "non-incidents". There are arguments on both sides. For many non-incidents water quality will not have been visibly affected. In those where quality has been affected it may be only mildly so resulting in increased likelihood of exposure (ie failure to reject the water). On the other hand, they are probably of a relatively small scale and so may not add to the power of the study. The data on area affected may be less good, since no 30 day report will be available. You should also be aware we have also given companies a commitment not to report non incidents against individual companies though this should not prevent including in this analysis. In conclusion, and contrary to my previous advice, I consider your report should give consideration to factors described above and the merits or otherwise of including or excluding non-incidents from any study design including effect both on the power and the cost of the study. I will
forward data on "non-incidents" to inform your judgement.

5) I mentioned that from 1 Jan 2004 DWI will hold individual compliance sample data that is geocoded or postcoded. We think your report should also give consideration to the merits or otherwise of comparing these data against any health data.

6) In my preliminary review of our database I have noticed a significant number of notifications that relate to loss of supply or reduced pressure, as you would expect these are almost all classified as non incidents. During such episodes the distribution system will be vulnerable to potential ingress which in turn could cause illness. Many discoloured water events will also be related to loss of pressure which may be a confounding factor. Consequently you should give consideration to how we can establish in the main study a relation between loss of pressure and illness and hence eliminate this from any relation between dirty water and illness. I will forward the relevant information
Annex 6: Modifications to the study design to address geographic resolution and the inclusion of ‘non-incident’ and ‘depressurisation’ events.

Following receipt of the draft Study Design Proposal in December 2004, the DWI commissioned an extension to the original contract to address several further issues.

Specifically the DWI requested:

“The additional objective is to:

Discuss with six selected water companies (names and contact details to be provided by DWI) the practical aspects of providing company data on the geographical area affected by event in, and/or converting into, a format that can be used in the proposed retrospective study.

1) The discussions should cover availability of data for both the discolouration and depressurisation events for the relevant period and irrespective of whether they are classified as incidents or non-incidents.

2) Discussion should identify any scope for misclassification of the exposed group both in term of the data available from the company and the proposed use of SOAs and the resulting reduction in the power of the study the such misclassification may cause.

3) The output should inform a more detailed discussion in the final report of how any possible confounding of association of illness with depressurisation may be quantified and eliminated from any possible association of illness with discolouration.”

These additional issues were addressed by visiting six water companies selected by the DWI (United Utilities, Three Valleys, Dwr Cymru, Northumbrian, Wessex, and Yorkshire) and discussing the
following questions with them. A brief conclusion is given for each question and a more detailed commentary is presented later.

1) How accurately can the area affected by an incident be defined, what is the likely degree of misclassification of exposure, and what are the practical implications of making the geographical data available?

Conclusion: For the majority of incidents, the affected area (defined as area from where complaints were received) can be identified at the 7-digit postcode level and analysed at the level of census Output Area. The area affected might be larger than the one identified, but the use of an exclusion boundary when selecting control areas will ensure that exposure misclassification will not be a significant problem. No major difficulties are envisaged in transferring the data.

2) Could events classified by the DWI as ‘non-incidents’ be included in a study?

Conclusion: Yes, the quality of data relating to non-incidents is adequate for addressing the primary aims of the study.

3) Could events of depressurisation be included in a study and could the separate effects of depressurisation and discolouration be examined?

Conclusion: Yes, the quality of data relating to depressurisation events is adequate for examining their separate effects on risk of GI illness.

A detailed commentary relating to each of the above questions is given below together with the proposed amendments to the study protocol.
1. Geographical definition of area affected by an ‘incident’

All six water companies visited use a geographic information system (GIS) to map customer complaints and water supply systems. In general, at least the first 50 customer complaints of each incident are held on the system (with a 7-digit postcode) and can be mapped and overlaid onto district metering areas (DMAs) and water supply systems. Subsequent calls, which may be recorded by an automated answering service, may be stored on other media and would need to be retrieved and uploaded onto the GIS system. This information, together with supplementary information (e.g. sampling measurements) from the ‘30-day’ paper report, could be used to identify groups of 7-digit postcodes affected by an incident and so define the affected area. According to the water companies, this task is likely to take an average of approximately 1 to 2 hours per incident and would require both a dedicated researcher familiar with the paper report and a GIS person within the company.

For some incidents (estimated at 20%) during 1999-2003, either none or only a small proportion of the customer complaints are currently available on the GIS system. In order to include these incidents in a study, it would be necessary to extract the data from either magnetic tape or the paper report. In this case, an additional 1 to 2 hours (estimated by the water companies) would be required per incident, so 20% of the incidents may require up to 4 hours of labour to obtain 7-digit postcode data.

The health data (HES/PEDW and NHSdirect) are also available at 7-digit postcode. However, it is proposed that analyses be conducted at the level of census Output Area (OA), the smallest (approximately 120 households) geographical area for which basic demographic data (e.g. exact population size, age structure) have been aggregated.

Two forms of exposure misclassification are possible: (i) that houses truly affected by water discolouration are not included in the exposed group, (ii) that houses truly unaffected by water discolouration are included in the exposed group. The first of these would only bias results if the affected house were included in the control (comparison) group, but since the proposed design
excludes adjacent Output Areas as potential controls this is unlikely to occur. In fact, both forms of misclassification will be minimised by the identification of affected and control areas at the level of Output Area. However, for certain incidents it may be more difficult to achieve 7-digit postcode (and hence OA) accuracy (e.g. if not all customer complaints are available or only a small proportion of customers complain), so the degree of misclassification may vary by incident. Larger incidents are likely to be less susceptible to bias due to misclassification, as a smaller proportion of households will be misclassified. Thus, the effect of misclassification on the results could be assessed by conducting sensitivity analyses (e.g. by excluding smaller incidents from the analyses). Any differences found following the exclusion of smaller incidents will, of course, need to be interpreted with caution given that smaller incidents may differ from larger incidents in factors other than size (e.g. cause).

All six companies reported that transferral of the postcode data from their GIS system would be relatively straightforward. One company was reluctant (for data protection reasons) to provide exact addresses of customer complaints.

**Summary**: For the majority of incidents, the area affected can be identified at the 7-digit postcode level. This will take an average of 1 to 2 hours per incident and involve both a researcher and a GIS person within the company. For approximately 20% of incidents, an extra 1 to 2 hours per incident will be required to transfer data from other sources. For all 280 incidents from 1999 to 2003, a total of 84 days (up to 4 reports per day) would be required. Analyses will be conducted at the level of census Output Area, which will improve the accuracy of population denominators whilst preserving a minimal level of misclassification error.

2. **Inclusion of “non-incidents”**

The information held by each company on non-incidents is broadly similar to that of incidents (i.e. customer complaint postcodes, DMA boundaries, and water supply systems on a GIS system) and the data availability, quality and transferability are likely to be comparable with that for incidents. Data
from incidents and non-incidents can therefore be pooled in the analysis, which will increase the power of the main study objectives. Only a 72-hour report is available for non-incidents, but the information necessary to address the primary objectives of the study should be available within these.

For a small number of non-incidents, the availability and quality of the data necessary for an analyses of the characteristics of the event (objective 3) may be inferior to that recorded in the 30-day incident reports.

As with incidents, there is a proportion (estimated to be 20%) of non-incidents between 1999 and 2003 for which the geographical data are currently not on the GIS system and would need to be transferred from other media. A similar process of examining the GIS data and paper report of each non-incident would enable identification of the affected area at 7-digit postcode level. The person-time required for this would be 84 days.

Inclusion of both incidents and non-incidents in the study would approximately double the number of discolouration events available for the analyses (560 in total) and increase the power of the study such that the minimum detectable rate ratios would be 1.2 (compared to 1.3 for 280 incidents only) for HES/PEDW and 1.1 (compared to 1.2) for NHS Direct.

**Summary:** The primary aims of the study can be addressed by including non-incidents in the analyses without compromising data quality. The contribution of non-incidents to an analyses of the characteristics of an event (objective 3) may be limited because there is no 30-day report on these. A further 84 days of labour would be required to extract the data on non-incidents from the water companies. The number of events available for analyses would double and allow detection of a 20% increase in GI-related hospital admissions and a 10% increase in NHSdirect calls over a two-week period in affected compared to control areas.

3. **Inclusion of depressurisation events (incidents and non-incidents)**
All six water companies reported that depressurisation events reported to the DWI invariably relate to a total loss of supply. They are recorded by the water companies in the same way as events of discolouration. Thus, the geographical data relating to them (e.g. customer complaints within the GIS system) are the same, making an analysis of GI illness following events of depressurisation possible.

The process of defining the geographical area affected would be similar to discolouration events above and would require similar person-hours of labour (168 days for both incidents and non-incidents).

Approximately half of all events (incidents and non-incidents) reported to DWI are classed as depressurisation events, so from 1999 to 2003 there are likely to be approximately 560 of these available for analysis. The study would, therefore, have 80% power to detect an increase of 20% (RR=1.2) in the rate of hospital admissions for GI illness over a two-week period.

A proportion of discolouration events is associated with a loss of pressure, so their effects can be examined separately and in combination. Interpretation of apparent effects will, however, be difficult given the potentially complicated causal pathways that may exist (e.g. depressurisation may lead to GI illness by causing discolouration, which could in turn results in GI illness; by mechanisms independent of discolouration; or both). Thus the causal factor may not be easily identifiable. The paper reports of each discolouration event can be examined to determine whether there was an associated loss of pressure during the event. This would take approximately 20 minutes per report, 30 days of labour in total (for 560 discolouration events).

**Summary:** The data available on depressurisation events are of equal quality to that of discolouration and separate analyses can be conducted on these. Where a discolouration report indicates a loss of pressure, then it may be possible to examine their combined effects but cautious interpretation will be required.
Summary of data extraction time

Discolouration

Identifying geographical area:

224 incidents @ 90 minutes each. 4 per day = 56 days.
56 incidents @ 180 minutes each. 2 per day = 28 days
224 non-incidents @ 90 minutes each. 4 per day = 56 days.
56 non-incidents @ 180 minutes each. 2 per day = 28 days

Identifying whether associated with depressurisation

560 events @ 20 minutes each. 18 per day = 30 days.

Depressurisation

Identifying geographical area:

224 depressurisation incidents @ 90 minutes each. 4 per day = 56 days.
56 depressurisation incidents @ 180 minutes each. 2 per day = 28 days
224 depressurisation non-incidents @ 90 minutes each. 4 per day = 56 days.
56 depressurisation non-incidents @ 180 minutes each. 2 per day = 28 days

Total number of days = 366
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