RESEARCH ARTICLE

The relationship between environmental sources and the susceptibility of *Acanthamoeba* keratitis in the United Kingdom

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Abstract

**Purpose**

To determine whether *Acanthamoeba* keratitis (AK) patients have higher rates of *Acanthamoeba* and free-living amoeba (FLA) colonising domestic sinks than control contact lens (CL) wearers, and whether these isolates are genetically similar to the corneal isolates from their CL associated AK.

**Methods**

129 AK patients from Moorefield Eye Hospital, London and 64 control CL wearers from the Institute of Optometry were included in this study. The participants self-collected home kitchen and bathroom samples from tap-spouts, overflows and drains using an instructional kit. The samples were cultured by inoculating onto a non-nutrient agar plate seeded with *Escherichia coli*, incubated at 32˚C and examined for amoebae by microscopy for up to 2 weeks. Partial sequences of mitochondrial cytochrome oxidase genes (*coxA*) of *Acanthamoeba* isolates from four AK patients were compared to *Acanthamoeba* isolated from the patient's home. The association between sampling sites was analysed with the chi-square test.

**Results**

A total of 513 samples from AK patients and 189 from CL controls were collected. The yield of FLA was significantly greater in patients' bathrooms (72.1%) than CL controls’ bathrooms (53.4%) (*p*<0.05). Spouts (kitchen 6.7%, bathroom 11%) had the lowest rate of *Acanthamoeba* isolation compared to drains (kitchen 18.2%, bathroom 27.9%) and overflow (kitchen 39.1%, bathroom 25.9%) either in kitchens or bathrooms (*p*<0.05). There was no statistically significant difference between the average prevalence of *Acanthamoeba* in all three sample sites in kitchens (16.9%) compared to all three sample sites in bathrooms (21.5%) and no association for *Acanthamoeba* prevalence between AK patients and CL controls. All four
corneal isolates had the same \textit{coxA} sequence as at least one domestic water isolate from the patients’ sink of the kitchen and the bathroom.

**Conclusion**

The prevalence of \textit{Acanthamoeba} and FLA was high in UK homes. FLA colonisation was higher in AK patients compared to controls but the prevalence of \textit{Acanthamoeba} between AK patients and CL controls domestic sinks was similar. This study confirms that domestic water isolates are probably the source of AK infection. Advice about avoiding water contact when using CL’s should be mandatory.

**Introduction**

Free-living amoebae (FLA) are unicellular eukaryotic organisms that can grow independently in different environments, including natural and man-made bodies of water; lakes, ponds, swimming pools, and even treated water supplies [1–3]. Some genera of FLA such as \textit{Acanthamoeba}, \textit{Vahlkampfia}, \textit{Naegleria} and \textit{Hartmannella} are opportunistically pathogenic to humans [1, 4].

The term \textit{Acanthamoeba} keratitis (AK) refers to infection of the cornea by \textit{Acanthamoeba}. However, other FLA such as \textit{Vahlkampfia} and \textit{Hartmannella} are also known causative agents of keratitis [4, 5]. AK and other amoebal keratitis are increasingly being recognized as a severe ocular infection worldwide that occurs most often among contact lens (CL) wearers and can lead to blindness [6–10]. Water contamination has been recognized as the most important risk factor for CL-associated AK [11–14].

\textit{Acanthamoeba} keratitis was first reported in 1974 as an extremely rare disease [15]. With the increased population of CL wearers, the incidence of AK has significantly risen [16, 17]. The reported incidence of AK in developed countries is up to 149 cases per million per year for contact lens wearers but it is less than 2 per million per year for non-contact lens wearers [18, 19]. In an outbreak in England and Wales during 1997–1999, the annual incidence of AK was 1.13 (in general adults) to 21.14 per million (in CL wearers) [18]. The latter study also found that the incidence of AK was much greater in areas supplied with hard water, which enhances limescale formation on pipes and so increases colonisation of \textit{Acanthamoeba} [18]. Furthermore, distance from water purification plants, use of stagnant water (for example cisterns), and warmer air temperature were found can be associated with higher incidence of AK [14, 19–26]. In a more recent outbreak that started in the UK in 2010–2011 a three-fold increase in the incidence of AK was reported compared to the outbreak in 2004–2009 [27]. The increased number of AK cases in the UK has been linked to increased use of disposable contact lenses in case control studies [28, 29] and improper lens hygiene [29].

Kilvington et al. have found 89% of patients with culture-positive AK contained FLA including \textit{Acanthamoeba} in tap water from their kitchens, or bathrooms, and water storage tanks were implicated as promoting this colonisation [11]. \textit{Acanthamoeba} were cultured from 30% of all homes, and 75% of isolates from domestic water and isolates from the corneas of AK patients had identical mtDNA profiles [11]. However, that study did not examine water samples from CL wearers who were not AK patients. Such sampling may help the understanding of the CL-wearing population’s risk of developing AK. In the current study, samples from both AK patients and control CL wearers and from different areas of their kitchen and bathroom sinks were cultured to understand whether AK patients have higher rates of
Acanthamoeba sink colonisation than control CL wearers. The current study also aimed to
determine the differences in the prevalence of Acanthamoeba and FLA in spouts, overflows
and drains of kitchens and bathrooms, whether Acanthamoeba colonisation of domestic water
systems remained constant over time and whether domestic waterborne Acanthamoeba iso-
lates were similar to those isolated from cases of AK.

Methods

Sample collection
A total of 129 AK patients from Moorfield’s Eye Hospital, London and 64 control CL wearers
from Institute of Optometry, London were included in this study. The research protocol
received approval from the National Research Ethics Service Committee London-Hampstead
(REC reference 13/LO/0032) and the Moorfields Eye Hospital Research governance commit-
tee. Written informed consent was received from participants before initiation of the study.
Each participant was provided with a sampling pack containing six sterile polyester-tipped
applicators and sterile screw-cap test tubes, written instructions (Supplementary information
S1 Data) and a questionnaire which included questions on the suburb, date and time of sample
collection, and the date and time that samples were returned to researchers. Participants were
requested to swab the inside of their bathroom and kitchen spouts, sink drains and overflows
for 10 seconds with the applicator, place the swab into the test tube, fill the tube with 5mL cold
tap-water, then fasten the test tube cap tightly. A total of 23 repeat samples at least one month
apart were collected from patient’s kitchen and bathroom.

Culture and microscopy
Upon receipt at the laboratory, tubes were vortexed and 500 μL of water was inoculated onto
1.4% non-nutrient agar (NNA) plate pre-seeded with 100μL of viable Escherichia coli. Each
agar plate was then incubated at 32°C in sealed polythene bags. After incubation for 3–4 days,
plates were examined daily for up to 10 days using an inverted light microscope for the pres-
ence of FLA and Acanthamoeba. Isolates were identified by morphologic examination of the
trophozoite and cyst forms [30]. Samples identified with Acanthamoeba were classified as
Acanthamoeba positive and those with Acanthamoeba or FLA were classified as FLA in cur-
rent analysis.

PCR assay, coxA sequencing and sequence homology analysis
Nucleotide sequence of the mitochondrial cytochrome oxidase subunit-1 and -2 (cox1/2) of
corneal isolates were compared with isolates from the patients’ homes. DNA was extracted
using Chelex resin (MB Chelex-100 resin; Bio-Rad Laboratories, Hercules, CA, USA) follow-
ning the method described by Kilvington et al.[31]. Cox1/2 was amplified by PCR using previ-
ously established primer and cycle conditions [11]. The amplified products were sent for
Sanger sequencing and DNA sequences were aligned using ClustalW and a phylogenetic tree
constructed using MEGA 7 [32].

Statistical analysis
The Pearson Chi-square test was used to assess whether there was a statistically significant dif-
fERENCE in the association between sampling sites. Odds ratios and their 95% confidence inter-
vals (95% CIs) were calculated to measure association between AK cases and detection rate of
Acanthamoeba and FLA in AK patients’ and CL controls’ homes.
Results

*Acanthamoeba* and free living amoeba colonisation

A total of 513 samples from 77 AK patients and 189 samples from 40 CL controls were retrieved and examined in this study. Samples were collected from water tap-spouts, sink over-flows and drains from the kitchen and bathroom. The proportion of samples from kitchens and bathrooms were broadly similar (Fig 1). Samples collected from the cloakrooms of AK patients were excluded from the current analysis because the number of samples were small (1.7%) and there were no cloakrooms samples from CL controls group.

A slightly higher proportion of *Acanthamoeba* were cultured from bathrooms (average 21.5%) compared to kitchens (average16.9%) (Fig 2) and there was no difference in this proportion between the patient and control group. However, bathrooms yielded a higher proportion of FLA positive samples compared to kitchens. Bathrooms from the AK cohort had a...
higher proportion (72.1%) of FLA than samples from the control’s bathrooms (54.3%) (p < 0.05) (OR 2.1; 95% CI 1.33–3.55). Furthermore, AK patient’s bathrooms had a significantly higher proportion of FLA than kitchens (p < 0.05) (Fig 2). Controls had a similar higher proportion of FLA from bathrooms, but this did not reach statistical significance.

Colonisation of *Acanthamoeba* in tap-spouts was lower in both bathrooms and kitchens (p < 0.05) than drains or overflows, but there was no statistical difference between patients and controls in the prevalence of *Acanthamoeba* in these sites (Fig 3). On the other hand, the rate of colonisation of FLA was higher in drains, overflows and tap-spouts of patients’ kitchens and bathrooms compared to these three sites from controls, and this reached statistical significance between bathroom’s overflow (controls 14/27 vs patients, 55/72, p = 0.02) and kitchen’s drain (controls 8/25 vs patients 40/66, p = 0.02) (Fig 4).

Twenty-three repeat samples at the two-sampling time-points for both bathrooms and kitchens spouts were examined to understand whether *Acanthamoeba* was consistently present at the same sites. *Acanthamoeba* isolation appeared to be sporadic from either bathrooms or kitchens (Table 1).

**Patients and environmental *Acanthamoeba* colonisation**

Four AK patients were selected to the study genetic relatedness between pathogenic and environmental isolates on the basis of association between case and detection of *Acanthamoeba* in water samples from their homes. Partial sequence of the mitochondrial cytochrome oxidase gene (*coxA 1/2*) was obtained from *Acanthamoeba* isolates of four different keratitis patients and those sequences were compared with isolates from the patient’s home. Fig 5 shows that
each pathogenic isolate had the identical coxA 1/2 sequence to at least one domestic water isolate including isolates from both the kitchen and the bathroom. The *Acanthamoeba* isolate from patient ID78 was similar to isolates from the patient’s bathroom spout, drain and overflow and their kitchen’s overflow suggesting a possible source of keratitis isolate. This isolate was also similar to the corneal isolate from another patient ID47 and an isolate from that patient’s kitchen spout, suggesting possibility of common source of infections. For patients ID81 and ID37, their corneal isolates were similar to those from their bathrooms. Water contact was reported by three of these four patients in the days prior to suffering AK.

**Discussion**

Domestic tap water is a major risk factor for contact-lens associated *Acanthamoeba* keratitis [11, 18, 20] and is a reservoir for *Acanthamoeba* and other FLA [11]. Based on coxA 1/2 gene sequence [33], this study identified that isolates from water sources were very similar to isolates from the corneas of AK patients, suggesting that domestic waterborne *Acanthamoeba* could be associated with AK. The current study examined water samples from CL wearers who were not AK patients and compared these to samples from AK patients. There was no variation in the rate of colonisation of *Acanthamoeba* in kitchens and bathrooms of AK patients compared to that of CL wearers controls, suggesting that all contact lens wearers are at risk of developing AK and it is not that AK patients have bathrooms or kitchens that are more...
frequently colonised. Patient’s bathrooms yielded a higher proportion of FLA compared to the bathrooms of control CL wearers or the kitchens of patients. Poorly maintained sinks may allow microbial biofilms to form, and this is believed to facilitate amoebal colonisation of water outlets and drains [11], not least because amoebae can graze of the bacteria in biofilms [34].

The prevalence of Acanthamoeba observed in this study (21.5%) was slightly lower than the prevalence reported by another UK based study, in which Kilvington et al. isolated Acanthamoeba from 30% of homes [11]. In the current study tap spouts tended to have filters on them making it difficult to swab the inside of spouts and this could be the reason for these differences [11]. In addition, spouts have the fastest water flow and large volume of water regularly flows through it, which self-flushes the spouts and may prevent colonisation of FLA.

Table 1. Repeatability of samples.

|          | Bathroom | Kitchen |
|----------|----------|---------|
|          | First Sample |          | First Sample |          |
|          | Positive | Negative | Positive | Negative |
| Second samples | 1 2 | 3 17 |
|           | Positive | 0 3 |
|           | Negative | 3 17 |

Fig 4. Colonisation of free-living amoeba (FLA) in samples for spout, drain and overflow from bathroom and kitchen in patients and control groups. * denotes level of significance (*p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, ****p ≤ 0.0001).

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Furthermore, there is a wide variation in the rate of isolation of *Acanthamoeba* in previous reports. Studies based in Hong Kong have shown that household tap water yielded 10% [35], 8.3% [36] and 7.7% [36] *Acanthamoeba* positive samples. In addition, studies from Scotland (12% bathrooms, 2% kitchen) [37], Jamaica (36%, tap-water) [38] and Florida (2.8%, domestic water) [39] have reported various isolation rates in water samples. One of the reasons for these variations may be due to use of different methods for identification of *Acanthamoeba*, for example morphological [39] versus molecular [38] techniques. Given that PCR-based methods are more sensitive than culture for detection of *Acanthamoeba* [40], further studies based on molecular identification will be required to validate rate of isolation of *Acanthamoeba* observed in the current study.

The domestic water supply system of the UK, which uses water storage cisterns as well as often having hard water, is believed to be a cause for the higher prevalence of *Acanthamoeba* in the tap water in this region [11]. *Acanthamoeba* and other microbes proliferate in the cistern water. Also, there is a seasonal trend with the colonisation of *Acanthamoeba* increasing in warmer months [22–24].

A limitation of this study is that the genus-level identification of FLA by molecular methods was not performed. FLA such as *Hartmannella* and *Vahlkampfiid* amoebae [4, 5], although rare compared to *Acanthamoeba*, are also causative agents of keratitis. In addition, the significantly higher FLA positive samples in the patient cohort suggests that their identification in the genus-level will help to better understand the risk of keratitis associated with domestic tap water. Further studies regarding the pathogenicity of FLA and *Acanthamoeba* will also require assessing the source of transmission as well as the severity of AK.

**Fig 5. Cytochrome oxidase gene (coxA 1/2) sequence comparison of patient and environmental *Acanthamoeba* isolates.**
In conclusion, this study suggested that *Acanthamoeba* colonisation in UK water supply is high and occurs in CL-wearers and AK patients. Domestic water isolates were similar to isolates from the cornea of AK patient, confirming that domestic waterborne *Acanthamoeba* is still associated with keratitis. Advice about avoiding domestic water contact when using CL’s should be mandatory.

**Supporting information**

S1 Data. Water sample instruction. (DOCX)

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