Molecular detection and genotyping of pathogenic protozoan parasites in raw and treated water samples from southwest Colombia

Claudia Sánchez 1,2,3, Myriam Consuelo López 2, Luis Alejandro Galeano 1, Yvonne Qvarnstrom 4, Katelyn Houghton 4,5 and Juan David Ramírez 3*

Abstract

Background: Protozoan parasites such as Giardia duodenalis, Cryptosporidium spp., Toxoplasma gondii and Entamoeba histolytica represent a great challenge to the systems producing water for human consumption because their cystic forms are persistent in the environment and resist to the disinfection methods conventionally used for their control. In this study, we investigated the presence of these protozoan pathogens in both raw and treated water samples used for the production of drinking water in Nariño Department, southwest Colombia. We collected 110 water samples (10 lot each sample) and analyzed them with real-time PCR (qPCR). qPCR-positive samples were genotyped with PCR and DNA sequencing.

Results: Giardia duodenalis was detected in 35/110 (31.8%) of the samples and Cryptosporidium spp. in 9/110 (8.2%) of the samples; no sample was positive for T. gondii, E. histolytica or C. cayetanensis. Giardia duodenalis was detected in samples of both raw water (Drinking Water Treatment Plants (DWTP): 47.83%; Drinking Water Rural Plants (DWRP): 18.42%) and water collected either after conventional physicochemical treatment (26.09%) or after disinfection by chlorine (50%), whereas Cryptosporidium spp. were only detected in raw waters (DWTP: 17.39%; DWRP: 13.16%). The two pathogens were detected in both types of treatment plants supplying water to urban areas and to rural zones. Analysis of gdh and tpi markers identified assemblages AII, AII and H of G. duodenalis, while analysis of the small subunit rRNA and gp60 markers of Cryptosporidium-positive samples identified C. parvum (Subtype IIcA5G3c), C. galli, C. molnari, Cryptosporidium sp. genotype II of bats and Cryptosporidium sp. genotype VIII of birds.

Conclusions: The results obtained demonstrate the presence of protozoan parasites in the water of the study region, and the need to improve the surveillance systems for these pathogens and identify the corresponding sources of contamination.

Keywords: Protozoan parasites, Raw water, Treated water, PCR, Sequencing analysis

* Correspondence: juandramirez@urosario.edu.co
Background

Water is an essential resource for life and thus access to safe water is currently considered a fundamental human right [1]. However, more than a billion people currently lack access to drinking water worldwide, presenting a risk for public health in affected regions [2]. In Colombia, on average, 78% of the population has access to drinking water; however, there are large differences in coverage between urban and rural areas, and it has been reported that around 1300 children die each year from diarrheal diseases caused by unsafe quality water consumption [3]. Nariño is a department located in southwest Colombia that has a risk index of water quality for human consumption (IRCA) of 50.27, which places it as a department at high risk within the country. Among the health problems related to water in Nariño, acute diarrheal disease has an incidence rate of 65.8 cases per 1000 inhabitants and a mortality rate of 18.2 cases per 1,000,000 inhabitants [4]. Additionally, in this department, an outbreak was recorded in 2017 with 2560 cases of acute diarrheal disease, in which it is presumed that water was the main vehicle of transmission of the causal agent (not yet identified) [5].

The most common and widespread risk associated with water is contamination by pathogenic microorganisms, such as viruses, bacteria and helmint and protozoan parasites. *Giardia* and *Cryptosporidium*, the causative agents of giardiasis and cryptosporidiosis, respectively, are the protozoan parasites most commonly associated with transmission by water. These pathogens affect not only humans, but a wide range of domestic and wild animals. Similarly, protozoan parasites such as *Entamoeba histolytica*, *Toxoplasma gondii* and *Cyclospora cayetanensis*, responsible for amebiasis, toxoplasmosis and cyclosporiasis, respectively, may also be transmitted by contaminated water sources and affect global health [6]. These protozoan parasites have been responsible for large numbers of outbreaks worldwide, in the period 2011–2016 at least 381 outbreaks caused by the transmission of water-borne parasitic protozoa were reported [7–9]. Protozoan parasites also represent a challenge to the production of water suitable for human consumption because their transmissible forms (cysts and oocysts) are highly stable and persistent in the environment, they can cross the physical barriers used to remove contaminants, and are resistant to several conventional disinfectants widely used in the treatment system of drinking water, such as chlorine and chloramines [10–12]. Finally, it must be emphasized that these pathogens can cause infection at rather low concentrations [13].

Current regulations in Colombia recently included the monitoring of *Giardia* spp. and *Cryptosporidium* spp. in the parameters for the microbiological control of water quality in the treatment plants responsible for the distribution of drinking water [14]. In Colombia, the presence of *Giardia* cysts and *Cryptosporidium* oocysts is regularly checked with immunofluorescence microscopy, according to the method validated by the United States Environmental Protection Agency in municipal drinking water (USEPA 1623). However, this method has several disadvantages and limitations, described by several authors as extensive experience in the microscopic differentiation of cystic forms and rigorous laboratory staff, high costs, subject to interference resulting from the presence of ions in the sample (manganese, iron and calcium) and it is not capable to identify species or genotypes, which is important for the determination of public health significance [15, 16]. As far as we know, to date there are no validated methodologies for the detection of *T. gondii*, *E. histolytica* and *C. cayetanensis* in water samples. Molecular biological techniques offer a methodological alternative in the study of protozoan parasites because their sensitivity and specificity are greater than those of traditional methods [10, 17]. For this reason, various studies have used these techniques to detect protozoan parasites such as *Giardia* [11, 18], *Cryptosporidium* [11, 18–21], *C. cayetanensis* [20, 22, 23] and *T. gondii* [20, 24, 25].

One of the great advantages of molecular biological techniques is that they allow the discrimination of microorganisms at the species and genotype levels, information that may be relevant in evaluating the sources of infection in humans and in the study of the potential risks posed by protozoan parasites [26]. For example, several assemblages of *G. duodenalis* (A-H) and about 37 species of *Cryptosporidium* have been described as associated with different hosts. Of these, particular genotypes of *C. parvum*, *C. hominis*, *C. andersoni*, *C. meleagridis*, *C. ubiquitum*, *C. cuniculus*, *C. suis* and *G. duodenalis* assemblages A and B are of special interest because they have been reported in water sources and may also present zoonotic potential [27, 28]. The identification of the different genotypes of these protozoan parasites in water sources can be useful to determine the possible sources of contamination through their association with the type of host they parasitize.

In this context, the aim of this study was to investigate the presence of protozoan parasites such as *G. duodenalis*, *Cryptosporidium* spp., *C. cayetanensis*, *E. histolytica* and *T. gondii* in samples of raw and treated water from water treatment plants that supply water for human consumption to urban areas and rural areas in the department of Nariño (southwest Colombia), using real-time PCR. We also identified the *Cryptosporidium* species and *G. duodenalis* assemblages with PCR and DNA sequencing.
Methods

Study area

This study was performed in three Drinking Water Treatment Plants (DWTP) currently supplying drinking water to urban areas and 11 Rural Plants currently supplying water to rural areas (DWRP), in the municipalities of Pasto (1° 12’52”N, 77°16’41”W; altitude: 2527 meters above sea level (masl); average temperature: 12 °C), Ipiales (0°49’44”N, 77° 38’26”W; altitude: 2900 masl; average temperature: 12 °C), Túquerres (1°05’14”N, 77°37’08”W; altitude: 3104 masl; average temperature: 11 °C), and Tumaco (1°48’24”N, 78°45’53”W; altitude: 1 masl; average temperature: 26 °C), all located in the Department of Nariño, southwestern Colombia (Fig. 1). The three DWTPs collect the surface waters from rivers as their supply source and use a conventional physicochemical treatment to produce drinking water, including the steps of coagulation, flocculation, sedimentation, filtration and disinfection by the addition of chlorine. The DWRPs of the municipality of Ipiales use surface waters of streams as their sources of supply (DWRP-IA, DWRP-IB, DWRP-IC, DWRP-ID and DWRP-IE) and the DWRPs of the municipality of Tumaco use the surface waters of rivers (DWRP-TE) and water from underground wells (DWRP-TA, DWRP-TB, DWRP-TC, DWRP-TD and DWRP-TF) as their sources of supply. In general, DWRPs undertake minimum water treatment before consumption, consisting mainly of the addition of chlorine. However, during the sampling period, only one of the 11 DWRPs (DWRP-TF) in the study applied some type of treatment.

Sampling

In total 117 water samples of 10 leach sample were collected (Additional file 1: Table S1). Of these, 72 water samples were collected in DWTPs at three different points: (i) 24 samples of raw water at the inlets to the plants; (ii) 24 samples after the physicochemical treatment; and (iii) 24 samples after disinfection with chlorine. The remaining 45 DWRP samples collected consisted of only raw water, because these plants did not apply any type of treatment at the times of sampling. The samples were collected on two occasions in 2016, one in March (rainy season) and the other in August-September (dry season). The sampling months were selected for convenience, based on the average monthly precipitation in the Department of Nariño, provided by the Institute of Hydrology, Meteorology, and Environmental Studies of Colombia.

Sample processing

Recovery of protozoan parasites

The raw water samples were filtered through cellulose membranes with 3 μm pore size (47 mm in diameter). The water samples taken after the physicochemical treatment and after the disinfection process were also filtered through cellulose ester membranes with 1.2 μm pore size (47 mm in diameter). The membranes used in the filtration processes were washed twice with 5 ml of elution solution (0.01% Tween 80 and antifoam). To concentrate the samples, the volumes recovered from the washes were transferred to sterile polystyrene tubes and centrifuged at 1500x g for 15 min. Afterwards, the supernatant was discarded, leaving 1 ml of it on the pellet, which was transferred to a new sterile tube.

DNA extraction

The DNA was extracted from 300 μl of the previously obtained cell suspensions, using the FastDNA® SPIN Kit for Soil (MP Biomedicals, Santa Ana, USA), according to
the manufacturer’s instructions. This was preceded by two cycles in a Mini-BeatBeater cell disruptor (California, USA) for 3 min and incubation on ice for 2 min to preferentially break the cystic forms of the protozoans. The DNA was eluted in 50 ml of elution buffer and stored at -20 °C until analysis. All water samples were spiked with a known concentration of a recombinant plasmid containing a sequence of an aquaporin of *Arabidopsis thaliana* as the internal control of amplification (IAC). This allowed the detection of PCR inhibitors in the samples [29].

**Detection of parasites using real-time PCR**

The DNA of *G. duodenalis*, *Cryptosporidium* spp., *T. gondii* and *E. histolytica* was detected with real-time PCR (qPCR) with endpoint detection, using primers and TaqMan probes previously reported for each of the protozoans [17, 19, 25]. The sequences of the primers and probes used are listed in Table 1. The qPCR assays were performed independently for each parasite in 96-well MicroAmp plates (Applied Biosystems, Foster City, USA), with a final reaction volume of 9 μl containing 3.5 μl of FastStart Universal Probe Master (Roche, Basel, Switzerland), 1.0 μl of each forward and reverse primer (10 μM) and 0.2 μl of TaqMan probe (5 μM) specific for each parasite or for the internal control (IC), 0.3 μl of water, and 2.0 μl of DNA. We used the sequence of an aquaporin of *Arabidopsis thaliana* cloned into a plasmid as the internal control, as reported elsewhere [29, 30]. To detect *Cryptosporidium* spp., the final reaction volume was 10 μl because 3.0 μl of DNA was included. The samples were processed in duplicate in

Table 1 Primers and probes used for the molecular detection of the protozoan parasites under study. In bold the fluorophores and quenchers.

| Parasite                      | Primer   | Sequence (5’-3’)   | Target       | Reference |
|-------------------------------|----------|--------------------|--------------|-----------|
| *Cryptosporidium* spp.        | CF       | GTTTTCATTATCAAGAAGCGAGTTAGGG | 18S rRNA     | [19]      |
|                               | CR       | GAGTAAGGAAACCCTTCAATCTCTAG  |              |           |
|                               | CP       | 6FAM/TCAGATACCGCTGTAGTCTTAACCATAAATGAATC/TAMRA |              |           |
|                               | SSU/rRNA | AGTGCAAGAAACACAAATACAGG   | 18S rRNA     | [34]      |
|                               | SSU/RNR  | CTCGTTTTAAGCATGCTTTCAG   |              |           |
|                               | GpF      | GCCGTTCCACTGAGGAAC        | gp60         |           |
|                               | GpR      | CCACATTCAAATGAAAGTGCCGC  |              |           |
| *Giardia duodenalis*          | GdF      | CATGCAATGCAGGCTCTCA      | 18S rRNA     | [17]      |
|                               | GdR      | AGCGGTGTCCGGCTAGC         |              |           |
|                               | GdP      | 6FAM/AGGACAAACGGTTGCAC/MGB |              |           |
|                               | GDHeF    | TCAACGTYYAYGGGCTTCCGT    | gdh          | [33]      |
|                               | GDHiF    | AGTACACACTGGGCTTCCG      |              |           |
|                               | GDHiR    | GTRRTCTTGTCCATCTGCC      |              |           |
|                               | AL3543   | AAATATGGCTGCTGTCG        | tpi          | [32]      |
|                               | AL3546   | CACACTTTTACGCGGACTA      |              |           |
|                               | AL3544   | CCCCCTTATCGGCGATCTTGT    |              |           |
|                               | AL3545   | GTGCGCAACCGCTGGGCGC      |              |           |
| *Entamoeba histolytica*        | EhF      | GTTTTGATTAGTACACAAATGCGCAATTC | 18S rRNA     | [17]      |
|                               | EhR      | TCCTGCGCCTCAATACCTTATCAC |              |           |
|                               | EhP      | 6FAM/CAATGAATCCGAATGAC/A/MGB |              |           |
| *Toxoplasma gondii*            | TgF      | TCCCCCTGCTGCGGAAAAAGT    | B1 gene      | [25]      |
|                               | TgR      | AGCGGTCTGCGTCTCAATCTGAT  |              |           |
|                               | TgP      | 6FAM/TCTGCGACATCTTATCGTGAC/MGB |              |           |
| *Cyclospora cayetanensis*      | CcF      | TAGTAAAGGCAACGATGTTTAC   | 18S rRNA     | [31]      |
|                               | CcR      | AAAT GCC ACC GTA GCC CAA TA |              |           |
|                               | CcP      | HEX/CACCGCATATCCATGCTGCGC/2DABCYL |              |           |
| Internal control (Aquaglyceroporine) | CIF      | ACCGTCAATGGAGCAGCAGTGA   | AQGP         | [30]      |
|                               | CIR      | CTCCGGGACAAACACCTTATAA   |              |           |
|                               | CIP      | VIC/AGCATCTGTCCTGAAGGT/NFQ-MGB |              |           |
the Applied Biosystems 7500 system, using the default parameters and 40 cycles of amplification, except for Cryptosporidium spp., for which 50 cycles were used. The results of qPCR were considered negative if the cycle threshold (Ct) values were > 38. This cycle threshold was determined in a previous study by our research group. The qPCR results were considered negative if the cycle threshold values (Ct) were > 38. To corroborate the Ct value, we conducted experiments to establish the dynamic range of our assay using standards from 10,000 fg/μl to 1 fg/μl. For quantification, plasmids containing the target sequences were cloned into the pGEM-T Easy Vector System I (Promega, Madison, USA), according to the manufacturer’s instructions, and transformed into XL1-Blue Escherichia coli (Agilent Technologies, California, USA). The transformed colonies containing the plasmids were extracted by using the QIAprep Spin Miniprep Kit (Qiagen, Valencia, CA). The purified plasmid DNA was quantified by using a Nanodrop and diluted to have a concentration range of 10,000 fg/μl to 1 fg/μl. The dynamic range established that the limit of detection was the proposed by Sanchez et al. [29]. We used DNA from G. duodenalis, Cryptosporidium spp., T. gondii and H. histolytica provided by the Parasitology Laboratory of the National University of Colombia as the positive controls in the corresponding reactions, and type I water as the negative control. We detected C. cayetanensis in collaboration with the Division of Parasitic Diseases and Malaria of the Centers for Disease Control and Prevention (Atlanta, USA), using a previously described method [31].

Identification of G. duodenalis assemblages and Cryptosporidium species

Samples positive on qPCR for G. duodenalis and Cryptosporidium spp. were genotyped with conventional PCR and DNA sequencing. To identify the G. duodenalis assemblages, we used the following molecular markers: gdh (glutamate dehydrogenase) amplified with the primers GDHeF, GDHiF and GDHiR and tpi (triose phosphate isomerase) using the primers AL3543, AL3546, AL3544 and AL3545, as previously described [32, 33]. To identify the Cryptosporidium species the small subunit (SSU) RNA marker was used, using the primers SSUrRNAF and SSUrRNAR, and subtyping of C. parvum was based on sequence analysis of gp60 genes as previously reported [29, 34]. The sequences of the primers are listed in Table 1. The samples were processed in a MultiGene OptiMax Thermal Cycler (Labnet, California, USA). The amplification products were verified on 2% agarose gels stained with the SYBR® Safe Stain (California, USA). The gel was visualized in an E-Gel Imager (Life Technologies, Carlsband, USA).

The PCR products were sequenced by the dideoxy-terminal method in an automated capillary sequencer (Applied Biosystems, Foster City, USA). The sequences were verified and edited in the MEGA 6.0 program [35] and then aligned with sequences from the GenBank database using the NCBI BLAST tool [36] and with reference sequences in the program ClustalW v.1.8. Phylogenetic reconstruction was performed using a maximum likelihood analysis in MEGA v6.0, using the Tamura 3-parameter method with 1000 bootstrap replicates [35]. The reference sequences considered for the gdh marker of G. duodenalis were: assemblage AI (M84604.1), assemblage AII (AY178737.1), assemblage BIII (AF069561.1), assemblage BIV (AY178739.1), assemblage C (U60982.2), assemblage D (U60986.2), assemblage E (AY178741.1), assemblage F (AB569384.1), assemblage G (AF069582.1) and assemblage H (GU176089.1). The sequence of G. ardeae (AF069062.0) was used as the outgroup. For the G. duodenalis tpi marker, the following reference sequences were used: assemblage AI (AF069556.1), assemblage AII (AF069557.1), assemblage BIII (AF069561.1), assemblage BIV (AF069560.1), assemblage C (AF069563.1), assemblage E (AY228645.1) and assemblage F (AF069585.1). The sequence of G. microti (AY228649.1) was used as the outgroup. For Cryptosporidium, the reference sequences used were: C. andersoni (AF093496.1), C. baileyi (L19068.1), C. bovis (AY741305.1), C. canis (AF112576.1), C. fayeri (AF159112.1), C. felis (AF108862.1), C. fragile (EU162751.1), C. galli (AF316624.1), C. hominis (AF108865.1), C. macrocopodum (AF13227.2), C. meleagris (AF112574.1), C. cf. molnari (AY524773.1), C. moris (AB089284.1), C. parvum (AF112571.1), C. ryanae (AY587166.1), C. cf. scophthalmi (KR340588.1), C. serpentis (AF151376.2), C. suis (AF115377.1), C. varanii (AF112573.1) and C. wrairi (AF113578.1). The sequences obtained were deposited in GenBank under the accession numbers MH730625–MH730644.

Data analysis

Descriptive statistics was used to describe the main events of interest; the results are reported as percentages and frequencies. Statistically significant associations between the detection of the parasites examined and variables, such as the place of sampling, type of water and time of sampling, were determined by logistic regression analysis, using the statistical program Statgraphics Centurion XVII (Royal Technologies, Bogota, Colombia). Differences were considered significant at P < 0.05.

Results

Detection of protozoan parasites in water samples with qPCR

A total of 117 water samples were collected, 110 of which were analyzed and seven were excluded from the
study because the internal control was not amplified in any of the qPCRs. The seven samples excluded were four water samples collected in DWTPs and three water samples taken from DWRPs. Of the 110 samples analyzed, 31.82% (35/110) were positive for *G. duodenalis* and 8.18% (9/110) for *Cryptosporidium* spp. We detected both protozoan parasites *G. duodenalis* and *Cryptosporidium* together in 2.73% (3/110) of the water samples analyzed. None of the processed samples were positive for *T. gondii* (0/110), *E. histolytica* (0/110) or *C. cayettesensis* (0/110).

*Giardia duodenalis* was detected in the three DWTPs studied and in five of the 11 DWRPs (DWRP-IA, DWRP-IC, DWRP-ID, DWRP-TA and DWRP-TB). Most positive samples were found in the DWTP in Ipiales, followed by the DWTPs in Túquerres and Pasto, indicating contamination was prevalent in water from plants supplying urban areas. *Cryptosporidium* was detected in two of the three DWTPs and in four of the 11 DWRPs (DWRP-IE, DWRP-TA, DWRP-TE and DWRP-TF). We detected the greatest numbers of positive samples in the DWRPs of Tumaco, followed by the DWTP of Pasto; *Cryptosporidium* was most frequently found in treatment plants supplying rural areas. However, we found no statistically significant correlation between the sampling site and the detection of *G. duodenalis* (logistic regression, \( P = 0.1017 \)) or *Cryptosporidium* (logistic regression, \( P = 0.4780 \)) (Table 2).

Of the two parasites detected, *G. duodenalis* was found in both raw and treated water samples from DWTPs (raw water: 11/23, 47.83%; after physicochemical treatment: 6/23, 26.09%; and after chlorination: 11/22, 50%) and in raw water samples from DWRPs (7/38, 18.42%). However, we detected no statistically significant relationship between the type of water and presence of this microorganism (logistic regression, \( P > 0.6835 \)). In the DWTP of the municipality of Pasto, *G. duodenalis* was detected with greater frequency in raw water (3/7, 42.9%) than in either water collected after physicochemical treatment (2/7, 28.6%) or after chlorination (1/7, 14.3%). In the DWTPs of the municipalities of Ipiales and Túquerres, this protozoan was more frequent in the water collected after chlorination (Ipiales DWTP 6/7, 85.7%; Túquerres DWTP 4/8, 50%) than in the raw water (Ipiales DWTP 5/8, 62.5%; Túquerres DWTP 3/8, 37.5%). The fewest positive samples of *G. duodenalis* were detected in water collected after the physicochemical treatment (Ipiales DWTP 2/8, 25%; Túquerres DWTP 2/8, 25%). *Cryptosporidium* spp. were only found in raw water samples in both the DWTPs (raw water: 4/23, 17.39%) and DWRPs (5/38, 13.16%), and there was a statistically significant relationship between the type of water evaluated and this microorganism (\( P < 0.0097 \)) (Fig. 2). *Giardia duodenalis* was detected more frequently in samples collected in the dry months of August-September (22/52, 42.31%) than in the rainy month of March (13/58, 22.41%; \( P < 0.0078 \)). However, the detection of *Cryptosporidium* did not vary significantly between the samples collected in August-September (3/52, 5.77%) and March (6/58, 10.34%; \( P > 0.3768 \)) (Fig. 3).

**Identification of *G. duodenalis* assemblages and *Cryptosporidium* species**

To identify the assemblages of *G. duodenalis* from the qPCR-positive samples (35/110, 31.82%), we determined the nucleotide sequences of 17 PCR products with the *gdh* marker and 28 PCR products with *tpi* marker using Sanger sequencing (the remaining samples showed only faint bands or no bands and could not be sequenced). Nucleotide sequences were determined satisfactorily for 13 samples (five with the *gdh* marker and eight with the *tpi* marker), and identified assemblages A1 (1/5, 20%), AII (1/5, 20%) and H (1/5, 20%) (particularly in Tumaco) with the *gdh* marker and A1 (1/8, 12.5%) and AII (4/8, 50%) with the *tpi* marker. The other samples had multiple nucleotide sequences in the same sample and therefore could not be analyzed. The same results were obtained when the samples generating these aberrant sequences were re-amplified and the PCR products sequenced again.

*Cryptosporidium* species present in the qPCR-positive water samples (9/110, 8.18%) were identified by sequencing 9 PCR products with the SSU rRNA marker. The nucleotide sequences were determined satisfactorily for seven samples, and detected *C. parvum* (1/9, 11.1%), *C. galli* (1/9, 11.1%), *C. molnari* (1/9, 11.1%), *Cryptosporidium* sp. genotype II of bats (1/9, 11.1%) and *Cryptosporidium* sp. genotype VIII of birds (3/9, 33.3%). The result of subtyping the only *C. parvum* sample showed the presence of genotype IIcASG3c. The sequences of the two remaining samples (2/9, 22.2%) only allowed their identification to the genus level.

**Discussion**

The methods used in this study allowed the recovery and detection of protozoan parasites from water samples. In the case of membrane filtration, this

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**Table 2** Number and percentage of positive samples for *G. duodenalis* and *Cryptosporidium* spp. for each sampling site

| Site (n) | *G. duodenalis* (%) | *Cryptosporidium* spp. (%) |
|---------|---------------------|--------------------------|
| DWTP Tumaco (n = 25) | 4 (16.0) | 4 (16.0) |
| DWTP Ipiales (n = 24) | 9 (37.5) | – |
| DWTP Pasto (n = 21) | 6 (28.6) | 3 (14.3) |
| DWRP Tumaco (n = 25) | 4 (16.0) | 4 (16.0) |
| DWRP Támesis (n = 20) | 4 (20.0) | 4 (20.0) |
| DWRP Guárico (n = 20) | 4 (20.0) | 4 (20.0) |
| DWRP Túquerres (n = 23) | 13 (56.5) | 1 (43) |
| DWRP Ipiales (n = 17) | 3 (17.6) | 1 (59) |

Abbreviations: DWTP Drinking Water Treatment Plants, DWRP Drinking Water Rural Plants, n number of samples
Methodology has been used in some Latin American studies, mainly in countries such as Brazil for the recovery of protozoan parasites from various environmental samples [37]. Among its advantages are its low cost, shorter processing time in the laboratory and allowing the processing of samples with complex physicochemical composition (oils, fats, organic matter). However, a high turbidity can limit the use of this methodology due to the obstruction of the filter pores, making the use of several membranes necessary for the filtration of a single sample [15]. One limitation of our study was the use of membrane filtration. Further studies must consider the cartridge filters used in Method 1623 [1].

Regarding the qPCR, this is a technique with great potential in the detection of protozoan parasites due to its high sensitivity, a feature of great importance in environmental samples in which this type of microorganism can occur at low concentrations [11, 38]. Nevertheless, this molecular technique also has some limitations, due to the presence of PCR inhibitors that can be found in water samples and their dependence on the quality of the DNA obtained [6, 39]. In order to avoid false negatives due to the presence of inhibitors in the sample, an internal control was used in this study; this internal control identified samples that did not amplify the target DNA during the qPCR, due to the presence of inhibitors.

**Fig. 2** Molecular detection of *G. duodenalis* and *Cryptosporidium* spp. a In Drinking Water Treatment Plants currently supplying drinking water to urban areas from the municipalities of Pasto, Ipiales and Túquerres. Abbreviations: Raw water, water samples collected before treatment; After pch, water samples collected after the physicochemical treatment; After Cl, water samples collected after disinfection with chlorine. b In Rural Plants currently supplying water to rural zones from the municipalities of Ipiales and Tumaco. Abbreviations: Raw w, water samples collected before treatment; After T, water samples collected after treatment. DWRP Ipiales: A (Yaramal), B (La Orejuela), C (Charandu), D (Loma de Zuras) and E (Chaguaipe). DWRP Tumaco: A (Km 36), B (Cajapi), C (El Ceibito), D (Inguapí el Carmen), E (Bajo Jagua) and F (Pueblo Nuevo).

**Fig. 3** Detection of *G. duodenalis* and *Cryptosporidium* spp. in the two seasons of sampling: March (rainy season) and August-September (dry season). Abbreviations: DWTP, Drinking Water Treatment Plant; DWRP, Rural Plant.
substances that can act as PCR inhibitors, and therefore should be excluded from the study. In future studies it would also be important to consider the use of PCR enhancers, which could also improve the results.

With respect to the results obtained, the detection of *G. duodenalis* and *Cryptosporidium* by qPCR in water samples in this study was consistent with reports worldwide [12, 40]. *Giardia duodenalis* and *Cryptosporidium* spp. can be detected in water samples because they are widely distributed in the environment and affect both humans and a wide range of domestic and wild animals [41, 42], facilitating their transmission to water resources that are exposed to contamination [13, 18, 43]. In the DWTPs that supply water to urban areas in Nariño Department, the sources of supply are the surface waters of rivers, which are exposed to the impact of anthropogenic activities before their capture by the treatment plants. Contaminants mainly include discharges of domestic and agricultural origin, industrial effluents, runoff from waste, or the products of livestock, agriculture and human activities [44, 45].

In the DWTP of Pasto, reductions in the numbers of positive samples of *G. duodenalis* and *Cryptosporidium* were observed as the treatment of the drinking water progressed. However, this trend was not observed in the DWTPs of Ipiales and Túquerres. *Giardia duodenalis* was detected in water samples taken after the physicochemical treatment and disinfection with chlorine, at considerably higher frequencies than those in the untreated water, indicating that the procedures used in these plants must be reviewed; these findings may indicate, e.g. poor maintenance of the disinfection units, accumulation of biofilms within the pipelines or lack of optimization in these treatment facilities. One of the key factors that should be reviewed in the three DWTPs examined in this study is the physicochemical treatment (coagulation, flocculation, sedimentation and filtration), because the elimination of protozoan parasites correlates strongly with the adequate operation of each stage of this process [46]. Special attention should be paid to the steps of coagulation and flocculation [47].

In the case of *G. duodenalis*, its detection in post-treatment water samples may also be related to the resistance of the cystic forms to oxidative conventional disinfectants, such as chlorine [43, 46], which is the method applied in the plants studied. Likewise, it could be due to post-treatment contamination by cysts of this protozoan parasite during the production process of drinking water, caused by infiltrations in the treatment system through leaks, open or crossed connections, manipulation of system elements, repair of pipes, or the formation of biofilms of *Giardia* in the pipes, representing a possible source of secondary contamination of water, due to the accumulation and concentration of cysts that may occur during periods of low flow. Studies conducted in Colombia by other authors also report the presence of *Giardia* in drinking water [6, 48, 49]. However, it is important to note that in this study there was no information on the viability or infectivity of the protozoan parasites detected and therefore no conjecture could be made about the risk associated with the detection of *G. duodenalis* in the post-treatment water.

In the DWRPs of the municipalities of Ipiales and Tumaco, both *G. duodenalis* and *Cryptosporidium* spp. were detected, indicating the contamination of the natural water sources captured for human consumption in rural areas. In the DWRPs of Ipiales, the water sources used for supply could be exposed to domestic discharges, residues from crops, and the care or breeding of animals, because different anthropogenic activities are undertaken in this area. In Tumaco’s DWRPs, the groundwater wells could be contaminated because they are directly exposed to the environment and the passage of domestic and wild animals, and they are inadequately maintained. The sizes of both *Giardia* cysts and *Cryptosporidium* oocysts could also allow them to leach into the groundwater, which has previously been reported as a threat to groundwater [50, 51]. In the treatment plants that take water from rivers, the sources of contamination could be the various activities frequently performed in the rivers, including recreation, personal hygiene and laundry, and the general residual wastes of domestic origin.

Regarding the *G. duodenalis* assemblages in the qPCR-positive samples, we identified assemblage A in the DWTPs of Pasto and Ipiales. This assemblage has been reported previously in humans, livestock, companion animals, some species of marine animals and wild mammals; it is essential to consider its zoonotic potential [33, 52]. In the DWTP of Pasto, the sub-assemblage AI was identified, and in the DWTP of Ipiales, the sub-assemblages AI and AII were identified. Of these, the sub-assemblage AI has been reported predominantly in livestock and pets, and AII in humans [53]. Therefore, the sources of contamination are probably associated with these hosts. We also detected a sample containing assemblage H, so far only reported in seals and gulls [27, 52]. This represents the first description of assemblage H in Colombia or South America. Currently, the animal reservoirs of assemblage H remain undefined. Interestingly, this sample was detected in Tumaco, located at the pacific coast of the country. One plausible explanation might be that the *Giardia* assemblage H came from a seabird or marine vertebrates. However, this premise is too speculative and future studies to determine the exact frequency of this assemblage in the country and the region, as well as its hosts, are needed. Unfortunately, it was only possible to sequence a limited
number of samples to identify the *G. duodenalis* assemblages, but similar results have been obtained in other studies [13, 39, 54]. The failure to amplify all qPCR-positive samples of *G. duodenalis* with nested PCR may be attributable to that the molecular marker used in the qPCR (18S rRNA) has a higher number of copies than the *gdh* and *tpi* molecular markers used in the nested PCR. However, it is also possible that some qPCR results were false positive results, due to the complexity of the environmental samples and the presence of other DNA that could cross-prime with the qPCR assays of *Cryptosporidium* spp. and *G. duodenalis* in the water samples collected and possibly with the excess of PCR cycles reported for this assay [39].

Of the *Cryptosporidium* species detected in the DWTPs, we identified *C. parvum*, *C. molnari*, *C. galli* and *Cryptosporidium* sp. genotype II of bats. *Cryptosporidium parvum* is considered one of the most widely distributed *Cryptosporidium* species, reported in more than 150 mammalian hosts, including humans, mice, cows, horses, sheep, goats, pigs and deer, and is one of the main causal agents of outbreaks of water-borne cryptosporidiosis [34, 55]. Particularly, we found the genotype IIcA5G3c, which is considered zoonotic and has been previously reported in Colombia [29]. The other species identified are considered to be host specific; *C. molnari* has mainly been described in fish, *C. galli* in birds such as chickens and finches [41], and *Cryptosporidium* sp. genotype II corresponds to the sequence of an isolate obtained from bats. It is important to remember that within the genus *Cryptosporidium*, there are several species and genotypes for which a definitive classification is still pending, so the genus is in continuous review [19, 56].

Little information is available on the relationship between the protozoan parasites detected in water resources across seasons in Latin America, where countries experience only rainy and dry seasons [16, 57]. In the present study, the number of samples positive for *G. duodenalis* was greater in August-September, which are dry months. However, there was no significant variation in the detection of *Cryptosporidium* in the dry and rainy months sampled. To better analyze this issue at a regional level, we recommend that future studies monitor these types of microorganisms over longer periods, taking into account the effects of phenomena such as El Niño and La Niña, which can alter the climatic patterns considered typical or normal for a specific region.

**Conclusions**

We used molecular methodologies to determine the presence of protozoan parasites, such as *G. duodenalis* and *Cryptosporidium*, in water samples collected from the Department of Nariño. Our results provide important insights into the transmission dynamics of these protozoans in water resources. They emphasize the need for continued research and monitoring for the presence of these types of microorganisms in the water sources for human consumption, and the measures and precautions that must be considered to mitigate water contamination before its arrival at treatment plants.

**Additional file**

**Additional file 1**: Table S1. Metadata information of the water samples collected and submitted to molecular detection of *Giardia duodenalis*, *Cryptosporidium*, *Entamoeba histolytica*, *Toxoplasma gondii* and *Cyclospora cayetanensis*. (DOCX 24 kb)

**Abbreviations**

DWRP: Rural plants currently supplying water to rural areas; DWTP: Drinking Water Treatment Plants currently supplying drinking water to urban areas; qPCR: Real-time PCR; SSU rRNA: Small subunit ribosomal ribonucleic acid; USEPA: United States Environmental Protection Agency; gdh: Glutamate dehydrogenase; tpi: Triose phosphate isomerase; masl: Meters above sea level

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**Availability of data and materials**

The data supporting the conclusions of this article are included within the article and its Additional file.

**Authors’ contributions**

CS wrote the manuscript, analyzed the data and carried out the sampling and experimental tests of molecular biology. YQ and RH performed the detection of *Cyclospora cayetanensis*. AG, MCL and JDR analyzed the data, and CS undertook her MSc study. The authors thank the Dirección de Investigación e Innovación de la Universidad del Rosario and Universidad de Nariño.

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

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1Grupo de Investigación en Matemáticas Funcionales y Catálisis (GIMFC), Departamento de Química, Facultad de Ciencias Exactas y Naturales, Universidad de Nariño, 520002 Pasto, Colombia. 2Departamento de Salud Pública, Facultad de Medicina, Universidad Nacional de Colombia, 111321 Bogota, Colombia. 3Grupo de Investigaciones Microbiológicas-UR (GIMUR), Programa de Biología, Facultad de Ciencias Naturales y Matemáticas,
References

1. UN-Water. UN-Water Technical Advisory Unit. In: Annual Report 2014. Switzerland: Geneva; 2014. http://www.unwater.org/publications/un-water-annual-report-2014/.

2. WHO. Guidelines for Drinking-Water Quality. Volume 1. 3rd ed. Geneva, Switzerland: World Health Organization; 2008. http://www.who.int/water_sanitation_health/dwq/fulltext.pdf.

3. Pérez A, Lozada P, Cruz C. Water safety plans. Fundamentals and prospects for implementing them in Colombia. Ing Inv. 2009;29:79–85 (In Spanish).

4. Ministerio de Salud y Protección Social. National Report on the Quality of Water for Human Consumption Based on the IRCA. Bogotá D.C. Colombia; 2016. https://www.minsalud.gov.co/sites/rd/Lists/BibliotecaDigital/RDEVS/PP/SA/irca-2015_reducido.pdf. Accessed 08 Apr 2018.

5. Instituto Departamental de Salud de Nariño. Epidemiological Bulletin. Ipiales, Colombia. 2017. p. 1–27. http://www.idns.gov.co/images/documentos/epidemiologia/boletines/bepidemiologico.pdf. Accessed 08 Apr 2018.

6. Triviño-Valencia J, Lora F, Zuluaga JD, Gomez-Marin JE. Detection by PCR of pathogenic protozoa in raw and drinkable water samples in Colombia. Parasitol Res. 2016;115:1789–97.

7. Efstatiou A, Ongerth JE, Karanis P. Waterborne transmission of protozoan parasites: Review of worldwide outbreaks - an update 2011–2016. Water Res. 2017;1:14–22.

8. Baldursson S, Karanis P. Waterborne transmission of protozoan parasites: review of worldwide outbreaks - an update 2004–2010. Water Res. 2011;45:6603–14.

9. Karanis P, Kourienti C, Smith H. Waterborne transmission of protozoan parasites: a worldwide review of outbreaks and lessons learnt. J Water Health. 2007;5:1–38.

10. Ali MA, Al-Herrawy AZ, El-Hawaary SE. Detection of enteric viruses, Cryptosporidium spp. and Giardia duodenalis in surface water: a health risk for humans and animals. Water Environ Res. 2004;76:3931–9.

11. Kumar T, Majid MAA, Onchichandan S, Jaturas N, Andiappan H, Salibay CC, et al. Presence of Cryptosporidium parvum and Giardia lamblia in water samples from Southeast Asia: towards an integrated water detection system. Infect Dis Poverty. 2016;5:3.

12. Castro-Hermida JA, Gonzalez-Warleta M, Miez O, Cryptosporidium spp. and Giardia duodenalis as pathogenic contaminants of water in Galicia, Spain: the need for safe drinking water. Int J Hyg Environ Health. 2015;218:132–8.

13. Castro-Hermida JA, Garcia-Preso JD, Almeida A, Gonzalez-Warleta M, Correia Da Costa JM, Miez O. Detection of Cryptosporidium spp. and Giardia duodenalis in surface water: a health risk for humans and animals. Water Res. 2009;43:4133–42.

14. Ministerio de la Protección Social. Resolución 2115: The resolution indicates the characteristics, basic instruments and frequencies of the control and monitoring system for the quality of water for human consumption. Colombia. 2007. p. 1–23. http://www.minambiente.gov.co/images/GestionIntegralldeRecursosHidrico/pdf/Legislativa%283%29_del_agua_Resolucion%282%29_2115.pdf. Accessed 08 Apr 2018.

15. Bueno R, Hachich E, Zanolli M, Hoffmann L, Cauchie HM. Two-year monitoring of Cryptosporidium parvum and Giardia lamblia occurrence in a recreational and drinking water reservoir using standard microscopic and molecular biology techniques. Environ Monit Assess. 2011;179:163–75.

16. Burnet JB, Ogorzaly L, Tissier A, Penny C, Cauchie HM. Novel quantitative TaqMan real-time PCR assays for detection of Cryptosporidium at the genus level and genotyping of major human and cattle-infecting species. J Appl Microbiol. 2013;114:1211–22.

17. Lakonde LF, Gadjadhur AA. Detection and differentiation of cocccidian oocysts by real-time PCR and melting curve analysis. J Parasitol. 2011;97:275–30.

18. Li N, Neumann NF, Rucker N, Alderisio KA, Sturbaum GD, Vilegas EN, et al. Development and evaluation of three real-time PCR assays for genotyping and source tracking Cryptosporidium spp. in water. Appl Environ Microbiol. 2015;81:5945–54.

19. Kitajima M, Hara-Moto E, Iker BC, Gerba CP. Occurrence of Cryptosporidium, Giardia, and Cyclospora in influent and effluent water at wastewater treatment plants in Arizona. Sci Total Environ. 2014;484:129–36.

20. Varma M, Hester JD, Schaefer FW, Ware MW, Lindquist HDA. Detection of Cryptosporidium cayetanensis using a quantitative real-time PCR assay. J Microbiol Methods. 2003;53:27–36.

21. Shapiro K, Mazet JAK, Schriewer A, Wuerthz S, Fritz H, Miller WA, et al. Detection of Toxoplasma gondii oocysts and surrogate microsporidians in water using ultrafiltration and capsule filtration. Water Res. 2010;44:893–903.

22. Lin MH, Chen TC, Kuo TT, Tseng CC, Tseng CP. Real-time PCR for quantitative detection of Toxoplasma gondii. J Clin Microbiol. 2003;41:4121–5.

23. Xiao L, Feng Y. Food and waterborne parasitology molecular epidemiologic tools for waterborne pathogens Cryptosporidium spp. and Giardia duodenalis. Food Waterborne Parasitol. 2017;8:9–14.

24. Monis P, Caccio S, Thompson A. Variation in Cryptosporidium: a worldwide review of outbreaks and lessons learnt. J Water Health. 2017;8:145–60.

25. Sánchez A, Muñoz M, Gómez N, Tabares J, Segura L, Salazar Á, et al. Molecular epidemiology of Giardia, Blastocystis and Cryptosporidium among Indigenous children from the Colombian Amazon basin. Front Microbiol. 2017;8:248.

26. Duffy T, Cura CI, Ramirez JC, Abate T, Cayo NM, Parrado R, et al. Analytical performance of a multiplex real-time PCR assay using TaqMan probes for quantification of Tryptosoma cruzi satellite DNA in blood samples. PLoS Negl Trop Dis. 2013;7:e2000.

27. Luanström Y, Benedict T, Märcet PL, Wiegand RE, Herwaldt BL, da Silva AJ. Molecular detection of Cyclospora cayetanensis in human stool specimens using UNEX-based DNA extraction and real-time PCR. Parasitology. 2018;145:865–70.

28. Sulaeman IM, Fayer R, Bern C, Gilman RH, Trout JM, Schantz PM, et al. Triosephosphate isomerase gene characterization and potential zoonotic transmission of Giardia duodenalis. Emerg Infect Dis. 2003;9:1444–52.

29. Read CM, Monis PT, Thompson RCA. Discrimination of all genotypes of Giardia duodenalis at the glutamate dehydrogenase locus using PCR-RFLP. Infect Genet Evol. 2004;4:125–30.

30. Hunter PR, Hadfield SJ, Wilkinson D, Lake IR, Harrison FCD, Chalmers RM. Subtypes of Cryptosporidium parvum in humans and disease risk. Emerg Infect Dis. 2007;13:82–8.

31. Tabara K, Stecher G, Peterson D, Filipski A, Kumar S. MEGA6: Molecular Evolutionary Genetics Analysis version 6.0. Mol Biol Evol. 2013;30:2725–9.

32. Autschbl SF, Gish W, Miller W, Myers EW, Lipman DJ. Basic local alignment search tool. J Mol Biol. 1990;215:403–10.

33. Bueno M, Bianco N, Guiguet A. Methods of concentration and detection of Cryptosporidium spp. and Giardia spp. in water samples. Rev Patol Trop. 2012;41:119–35.

34. Carey C, Lee H, Tveirys J. Biology, persistence and detection of Cryptosporidium parvum and Cryptosporidium hominis oocyst. Water Res. 2004;38:818–62.

35. Plutzer J, Karanis P, Domokos K, Törökné Á, Márialigeti K. Detection and characterisation of Giardia and Cryptosporidium in Hungarian raw, surface and sewage water samples by IFT, PCR and sequence analysis of the SSU rRNA and gdh genes. Int J Hyg Environ Health. 2008;211:524–33.

36. Sato MIZ, Galvani AT, Padula JA, Nardocci AC, Lauretto M de S, Razzolini MTP, et al. Assessing the infection risk of Giardia and Cryptosporidium in public drinking water delivered by surface water systems in Sao Paulo State, Brazil. Sci Total Environ. 2013;442:389–96.
41. Fayer R. Cryptosporidium: a water-borne zoonotic parasite. Vet Parasitol. 2004;126:37–56.
42. Plutzer J, Ongerth J, Karanis P. Giardia taxonomy, phylogeny and epidemiology: facts and open questions. Int J Hyg Environ Health. 2010;213:321–33.
43. Castro-Hermida JA, García-Presedo I, Almeida A, González-Warleta M, Correia Da Costa JM, Mezo M. Presence of Cryptosporidium spp. and Giardia duodenalis through drinking water. Sci Total Environ. 2008;405:45–53.
44. CORPONARÍN. Plan for the Regulation of the Main Stream of the Rio Pasto. Pasto, Nariño; 2011. http://corpoganono.gov.co/expedientes/descontaminacion/poehriopasto.pdf. Accessed 08 Apr 2018.
45. CORPONARÍN. Plan for the Regulation and Management of the Hydrographic Basin of the Tio Blanco. Ipiales, Nariño; 2011. http://corpoganono.gov.co/expedientes/descontaminacion/poehrioblanco.pdf. Accessed 08 Apr 2018.
46. Schoenen D. Role of disinfection in suppressing the spread of pathogens with drinking water: possibilities and limitations. Water Res. 2002;36:874–88.
47. Betancourt WQ, Rose JB. Drinking water treatment processes for removal of Cryptosporidium and Giardia. Vet Parasitol. 2004;126:219–34.
48. Alarcón M, Beltrán M, Cárdenas M, Campos M. Presence and viability of Giardia spp. and Cryptosporidium spp. in drinking water and wastewater in the high basin of Bogotá river. Biomédica. 2005;25:353–65 (In Spanish).
49. Lora-Suárez F, Rivera R, Triviño-Valencia J, Gómez-Marin JE. Detection of protozoa in water samples by formalin/ether concentration method. Water Res. 2016;100:377–81.
50. Elfadaly HA, Hassanain NA, Hassanain MA, Barakat AM, Shaapan RM. Evaluation of primitive ground water supplies as a risk factor for the development of major waterborne zoonosis in Egyptian children living in rural areas. J Infect Public Health. 2018;11:203–8.
51. Silfko TR, Smith HV, Rose JB. Emerging parasite zoonoses associated with water and food. Int J Parasitol. 2000;30:1379–93.
52. Feng Y, Xiao L. Zoonotic potential and molecular epidemiology of Giardia species and giardiasis. Clin Microbiol Rev. 2011;24:110–40.
53. Ryan U, Paparini A, Monis P, Hijjawi N. It’s official - Cryptosporidium is a gregarine: what are the implications for the water industry? Water Res. 2016;105:305–13.
54. Nguyen TT, Taub R, Pham PD, Nguyen HV, Nguyen KC, Phung CD, et al. Prevalence and molecular characterization of Cryptosporidium spp. and Giardia spp. in environmental samples in Hanam Province, Vietnam. Food Waterborne Parasitol. 2016;3:13–20.
55. Abeywardena H, Jex AR, Nolan MJ, Haydon SR, Stevens MA, McNulty RW, et al. Genetic characterisation of Cryptosporidium and Giardia from dairy calves: discovery of species/genotypes consistent with those found in humans. Infect Genet Evol. 2012;12:1984–93.
56. Navarro-I-Martinez L, Del Águila C, Bornay-Llinares FJ. Cryptosporidium: a genus in revision. The situation in Spain. Enferm Infeccc Microbiol Clin. 2011;29:135–43 (In Spanish).
57. Neto RC, Dos Santos LU, Sato MIZ, Franco RMB. Cryptosporidium spp. and Giardia spp. in surface water supply of Campinas, Southeast Brazil. Water Sci Technol. 2010;62:217–22.