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A Comparative Analysis of Microscopy, Coproantigen Serology, and Nested Multiplex PCR in the Laboratory Diagnosis of Entamoeba histolytica Infection

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Abstract **Objectives** Amoebiasis is caused by the most common intestinal protozoan parasite Entamoeba histolytica. This parasite causes amoebic colitis, which is manifested by diarrhea, followed by dysentery. The laboratory diagnosis of intestinal amoebiasis in most cases is by microscopic examination of stool samples. Other nonroutine methods include coproantigen enzyme-linked immunosorbent assay (ELISA) from stool samples, serum ELISA for antibodies, stool culture, isoenzyme analysis, and polymerase chain reaction (PCR). The present study aimed to comparatively analyze the different diagnostic modalities used for the detection of *E. histolytica* from the stool sample of patients with intestinal amoebiasis. Materials and Methods This study was undertaken with 631 patients, during a period of 3 years, from January 2017 to December 2019. Stool specimen obtained from each patient was subjected to direct microscopic wet mount examination, coproantigen

ELISA, and nested multiplex PCR, respectively. **Results** Out of all the patients tested, 5.2% were positive for *E. histolytica*. Among the positive cases, stool microscopy was positive in 3.17%, coproantigen ELISA was positive

in 29 (4.6%) cases, and PCR was positive in 30 (4.75%) cases. Statistical Analysis The prevalence of E. histolytica infection was summarized as percentages. The three diagnostic tests done were statistically analyzed, taking microscopy as the gold standard. The agreement between techniques (microscopy, coproantigen ELISA, and PCR) was analyzed with kappa statistics. Sensitivity, specificity, positive predictive value, negative predictive value, and diagnostic accuracy were summarized as percentage with 95% confidence interval.

► amoebiasis

Keywords

- coproantigen testing Entamoeba histolytica
- amoebic colitis
- amoebic dysentery

Conclusion In all suspected amoebiasis cases, a combination of stool microscopy, coproantigen testing with molecular detection of the parasite offers the best approach to diagnosis of this parasitic infection.

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Introduction

Amoebiasis is caused by the intestinal protozoan parasite *Entamoeba histolytica*. Owing to the lack of adequate sanitation and increased fecal contamination of water supplies, this parasitic disease is more commonly seen in developing countries.¹ Apart from cases of *E. histolytica* infection, there have been other case reports of *E. dispar* and *E. moshkovskii* being isolated from patients with gastrointestinal symptoms, but there is no convincing evidence of a causal association between the isolation of these two species and the clinical manifestations in the cases reported.² Nowadays, it is also observed that *Entamoeba moshkovskii* is emerging as another protozoan parasite causing infantile diarrhea.^{3,4}

The fourth leading parasitic cause of mortality worldwide is attributed to amoebiasis and data reveal that it caused 11,300 deaths globally in the year 2013.⁵ Many people get infected with E. histolytica, making amoebic colitis a leading cause of diarrhea, which kills more than 55,000 people each year, owing to its severe complications.⁶ In India, approximately 15 to 20% of the population is affected by this protozoan parasite.⁷ The clinical features of amoebiasis range from asymptomatic colonization to extraintestinal invasive amoebiasis. Usually, the affected cases are asymptomatic, but invasive intestinal infection also manifests in certain patients, which has a range of clinical features like chronic abdominal cramps, tenesmus, watery diarrhea or dysentery, and loss of weight.⁸ A prompt diagnosis is crucial for the patients with amoebic dysentery and asymptomatic cases so as to interrupt the transmission of disease. The laboratory diagnosis of E. histolytica infection is challenging, and the currently available diagnostic methods lack sensitivity. In the developing countries, intestinal amoebiasis is usually diagnosed in laboratory by identifying parasite cysts or motile trophozoites by wet mount examination of the suspected stool samples. The drawbacks of this conventional method include its low sensitivity and specificity, with false positive results common, owing to the presence of other protozoans like E. dispar or E. moshkovskii. Ideally, diagnosis should be framed taking into consideration a combination of laboratory testing methods, namely the microscopic detection of parasite in stool specimens, serodiagnosis by detection of E. histolytica-specific coproantigen, and stool polymerase chain reaction (PCR) for detection of parasite DNA.

The present study aimed to comparatively analyze the currently used different diagnostic modalities for the detection of *E. histolytica* from stool sample of patients clinically diagnosed with intestinal amoebiasis.

Materials and Methods

This cross-sectional study was conducted from January 2017 to December 2019 in a tertiary care hospital, Puducherry, south India. Ethical approval for the study was obtained from the Jawaharlal Institute of Postgraduate Medical Education & Research (JIPMER) Institute ethics committee. Considering a prevalence of *E. histolytica* as 4.5% with 1.35 absolute precision, the calculate sample size for the study was 734. We have included all eligible patients visited outpatient department (OPD) during the study period.

A group of 631 participants included in the study and their results were analyzed. All the subjects who reported to the hospital OPD with complaints of abdominal pain, vomiting, diarrhea, dysentery, indigestion, flatulence, dehydration, and weight loss were included. The cases with suspected bacterial causes of gastroenteritis and noninfectious etiology were excluded from the study. About 5 to 8 g of stool sample was collected in a 25 mL, sterile, screw-capped container, on 3 consecutive days from each patient. A written informed consent was obtained from all the study participants. A detailed history of prior treatment with any antiamoebic drugs or antibiotics was taken at the time of collection of stool samples from the clinically suspected patients. The unpreserved stool samples were transported to the laboratory for processing within 1 hour of collection. After reaching the laboratory, macroscopic examination of stool sample for noting the color, consistency, presence of blood, mucus, etc. was done. Then, each stool sample was divided into three parts. The first part was used for direct wet mount examination and formol ether sedimentation concentration method, according to Garcia et al.⁹ The second and third part of stool sample were stored at -20°C for coproantigen detection by enzyme-linked immunosorbent assay (ELISA) and PCR, respectively.

Direct Stool Microscopy

Fresh unpreserved stool samples were subjected to a wet mount examination in 0.9% saline and Lugol's iodine. Each wet mount was microscopically examined for pus cells, mucus flecks, red blood cells and for the presence of E. histolytica quadrinucleate cysts and trophozoites. The trophozoites of a typical E. histolytica are approximately 20 to 30 µm in diameter. It has a nucleus containing a central endosome, peripheral chromatin deposition, and radial achromatic fibrils, which gives the appearance of "cart-wheel" morphology. The serial stepwise motility of E. histolytica trophozoite from a patient sample is demonstrated in Fig. 1A. Mostly, the mature parasite cysts are spherical, quadrinucleate, and measure approximately 10 to 15 μ m in diameter (**>Fig. 1B**). The stool samples were also concentrated by formol-ether sedimentation technique, by the procedure given by Garcia et al,⁹ where wet mount was inconclusive or negative. Apart from a

Fig. 1 (A,B,C) The serial stepwise locomotion images of a motile *Entamoeba histolytica* trophozoite found in a positive stool sample of a case. (D) The quadrinucleate cyst of *E. histolytica* found in the stool sample of a positive case.

standard wet mount, the stool specimens were subjected to a modified iron-hematoxylin staining.

Coproantigen Detection by ELISA

All the second part of patient's stool samples was batched and E. histolytica ELISA was performed once a week. Each stool sample was subjected to coproantigen detection ELISA according to the manufacturer's instructions (Savyon Diagnostics Ltd, Israel). For detection of antigen in each stool sample, 100 µL of diluted stool specimen was added to one of the antibody-coated microtiters well of the plate. It was incubated for 1 hour at 37°C and washed. Then, 100 µL of horseradish peroxidase-conjugate was added and incubated for 1 hour at 37°C and washed. After that, 100 µL of 3,3',5,5'tetramethylbenzidine-substrate was added, incubated for 15 minutes at room temperature. The reaction was terminated by addition of 100 µL of stop solution. Finally, the absorbance was read at 450 nm using an ELISA reader. According to the manufacturer's instructions, a test was considered positive when the optical density reading of a sample was > 0.15 at 450 nm. The kit had a sensitivity of 100% and specificity of 95.4%.

Nested Multiplex PCR

The third part of each stool sample was subjected to nested multiplex PCR targeting the 16S-like rRNA gene for simultaneous detection and differentiation of E. histolytica, E. moshkovskii, and E. dispar, according to the method given by Khairnar and Parija.¹⁰ The parasite nucleic acid extraction from the stool specimens was performed by a cetyltrimethylammonium bromide extraction method. The extracted DNA was then quantified in a spin column and purified DNA extract from stool specimens was determined by ultraviolet (UV) absorbance using a double beam spectrophotometer. The genus-specific primers sequences used for the first PCR were E-1 5' TAAGATGCACGAGAGCGAAA 3' (forward primer) E-2 5' GTACAAAGGGCAGGGACGTA 3' (reverse primer). The species-specific primer sequences used in the second nested PCR were E. histolytica species EH-1 5' AAGCATTGTTTCTA-GATCTGAG 3' (forward primer) EH-2 5' AAGAGGTCTAACC-GAAATTAG 3' (reverse primer), E. moshkovskii species Mos-15' GAAACCAAGAGTTTCACAAC 3' (forward primer) Mos-2 5' CAATATAAGGCTTGGATGAT 3' (reverse primer), and E. dispar species ED-1 5' TCTAATTTCGATTAGAACTCT 3' (forward primer) ED-2 5' TCCCTACCTATTAGACATAGC 3' (reverse primer), respectively. For a reaction volume of 25 μ L, comprising 2.5 μ L of 10XPCR buffer (Biogene), 1.5 µL of 25 mM MgCl₂ (Bangalore Genei Ltd), 1.4 µL of deoxynucleoside triphosphate mix (5mM each dNTP, ABgene), 0.3 μ L (5 IU/ μ L) of Taq polymerase (Biogene Ltd), 0.3 µM of each primer (IDT) and 2.5 µL of template DNA was added in genus-specific and species-specific PCR. The PCR reaction tubes were then placed in a thermal cycler. The PCR mix underwent an initial denaturation at 96°C for 2 minutes, followed by 30 cycles of PCR. Each amplification cycle consisted of 92°C for 60 seconds (denaturation), 56°C for 60 seconds (annealing), and 72°C for 90 seconds (extension). Finally, one cycle of extension at 72°C for 7 minutes was performed. In the species-specific nested multiplex PCR (which

had multiple primer sets in the same tube), only the annealing temperature was modified. About 3 µL of the amplification products were subjected to electrophoresis through 1.8% agarose gel at 120 V for 45-minute duration, and were visualized by ethidium bromide dye staining under UV light, for bands of target DNA detected. The PCR product giving bright band at 439 bp was considered positive for E. histolytica, whereas all other bands were taken as negative for E. histolytica (**Figs. 2A, B**). Among the other bands, the PCR product giving bright bands at 553 and 174 bp was considered positive for E. moshkovskii and E. dispar, respectively. Positive and negative control reactions were included with each batch of samples analyzed by nested multiplex PCR. The positive control used for Entamoeba and the individual species were an inhouse designed positive controls, obtained after sequencing the highly conserved region. A negative sample, which was proven PCR negative, was used as the negative control. The nested multiplex PCR detected E. histolytica, E. dispar, and E. moshkovskii DNA at a rate of 1,000 parasites/0.05 g of feces. The detection limit of nested multiplex PCR for E. histolytica was 25 Entamoeba protozoa cells.

Statistical Analysis

The study data for analysis were entered into Microsoft Excel sheet. The three diagnostic tests done were statistically analyzed, taking microscopy as the gold standard. The agreement between techniques (microscopy, coproantigen ELISA and PCR) was analyzed with kappa statistics. Sensitivity, specificity, positive predictive value (PPV), negative predictive value (NPV), and diagnostic accuracy were summarized as percentage with 95% confidence interval. All the statistical analysis was performed with IBM SPSS Statistics 20 and Open Epiversion 3.01 software.

Results

A total of 631 patients participated in the study. The unpreserved stool samples from all the 631 patients were subjected to three different diagnostic modalities for diagnosis of Entamoeba spp. infections. The three tests used were stool microscopy, coproantigen ELISA, and nested multiplex PCR. A positive result obtained in any one of the above three diagnostic tests used was considered as a positive case.

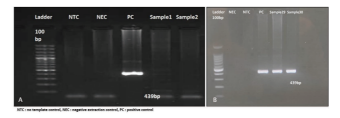


Fig. 2 Amplified products of PCR were analyzed by agarose gel electrophoresis. The size of the amplification product is indicated on the right (in base pairs). Samples 1 and 2 are negative for *Entamoeba histolytica* and the PCR products showing the bright band at 439 bp (samples 29 and 30) are positive for *E. histolytica*, along with the positive and negative controls used for the nested multiplex PCR. NTC, no template control; NEC, negative extraction control; PC, positive control; PCR, polymerase chain reaction.

Among the 631 patients, 33 (5.2%) patients were positive for *E. histolytica* infection. The multiplex PCR results revealed 30 (4.7%) cases to be positive for *E. histolytica*, 9 (1.4%) cases to be positive for *Entamoeba dispar*, and 3 (0.4%) cases to be positive for *E. moshkovskii* cysts in stool sample of patients. These two species, that is, *E. dispar* and *E. moshkovskii*, were not included for analysis.

Among the *Entamoeba histolytica* infection cases, analysis of the different diagnostic techniques used showed that, stool microscopy was positive in 20 (3.17%) cases and negative in 611 (96.83%) cases, coproantigen ELISA was positive in 29 (4.6%) cases and negative in 602 (95.4%) cases, and multiplex PCR was positive in 30 (4.75%) cases and negative in 601 (95.25%) cases (**-Table 1**). It was also observed that the most commonly affected age group was between 26 and 45 years and the disease was more prevalent among the male gender (**-Table 2**). The individual *E. histolytica* positive cases were tabulated according to the results obtained by the diagnostic method results used (**-Table 3**). Further, the percentage shows the % positivity among the three stool samples collected from each positive case (**-Table 3**).

The statistical analysis was done, taking microscopy as the gold standard. The statistical agreement between microscopy and coproantigen ELISA was found to be 0.79, using kappa statistics with 95% confidence interval (**-Table 4**). The validity indicators of coproantigen ELISA against microscopy showed a sensitivity of 90%, specificity of 98.2%, PPV of 62.07%, NPV of 99.67%, and a diagnostic accuracy of 97.94% (**-Table 5**). The statistical agreement between microscopy and PCR was found to be 0.72, using kappa statistics with 95% confidence interval (**-Table 6**). The validity indicators of PCR against microscopy showed a sensitivity of 100%, specificity of 98.36%, PPV of 66.67%, NPV of 100%, and a diagnostic accuracy of 98.42% (**-Table 7**).

Discussion

The present study detected 5.2% cases to be positive for *E. histolytica* among a total of 631 patients tested over a duration of 3 years. This study showed male gender was more commonly affected, belonging to the age group 26 to 45 years. Similar findings were also noted by Tharmaratnam et al and Kantor et al in their study.^{4,11}

From the very old days, the laboratory diagnosis of *E. histolytica* has always been based on direct microscopic examination of stool samples showing distinct protozoan morphology.¹² Currently, however, usage of the direct microscopy-based identification techniques to differentiate among protozoa with similar morphological features like *E. coli, E. moshkovskii, E. dispar*, and *E. hartmanni* is seemingly

unreliable.² There is a vast diversity in the morphological appearances of the cysts and trophozoites of Entamoeba species. Therefore, the confirmed identification of these intestinal parasites requires observation by a skilled microscopist. Moreover, various studies prove that the sensitivity and specificity of conventional microscopy, on a single stool specimen, for distinguishing the various species of Entamoe*ba* are far less than reliable.^{13,14} In fresh wet mount preparation, the locomotion of *E. histolytica* in fresh preparations usually occurs linearly, with the clear hyaline ectoplasm flowing to form blunt-ended pseudopodia, then leading the granular endoplasm containing the nucleus to flow in the same direction.¹⁵ At times, when a fresh stool specimen cannot be examined immediately, it may be preserved with a fixative like polyvinyl alcohol or maintained at 4°C temperature. Stool specimens can be examined either unstained or stained with Lugol's iodine. Iodine increases the contrast and improves the clarity of internal structures like nucleus of the parasite. Wheatley's trichrome staining or modified iron hematoxylin stains for permanent smears have been suggested as better stains for regular use in the diagnosis of *E. histolytica* or *E. dispar*.^{16–18} There are several factors that adversely affect the results of microscopy. These include lack of trained microscopists; delayed transport of sample to the laboratory; difficulty in differentiation between nonmotile trophozoites and polymorphonuclear leukocytes or macrophages; inappropriate sample collection; interfering substances such as antibiotics, purgatives, antacids, cathartics, antidiarrheal preparations (kaolin or bismuth), or enemas; inadequate number of specimens collected (at least three specimens are needed due to intermittent shedding nature of the parasite cyst); lack of preservation of stool specimens with fixatives; and presence of other commensal amoebae.¹⁹ CDC guidelines say that multiple stool samples (at least 3) should be tested before a negative result is reported, and stool samples in formalin, or other fixatives, need to be concentrated prior to microscopic examination²⁰ and finally the choice of diagnostic techniques should depend on available equipment and reagents, experience, and considerations of time and cost.

In the present study, it was observed that coproantigen ELISA was positive in 29 (4.6%) cases and negative in 602 (95.4%) cases. The validity indicators of coproantigen ELISA against microscopy showed a sensitivity of 90%, specificity of 98.2%, PPV of 62.07%, NPV of 99.67%, and a diagnostic accuracy of 97.94%. This is similar to the study findings by el-Hamshary et al and Bayoumy et al, where they have concluded that coproantigen assay using ELISA is more sensitive and specific than microscopy even when the parasitic count is low, thus reducing the chances of missing

Table 1 Result showing the comparative performance of different diagnostic methods (n = 631)

Result	Microscopy	%	Serology by coproantigen ELISA	%	PCR	%
Positive	20	3.17	29	4.60	30	4.75
Negative	611	96.83	602	95.40	601	95.25

Abbreviations: ELISA, enzyme-linked immunosorbent assay; PCR, polymerase chain reaction.

Table 2	Age and	sex distri	bution of	t the pos	itive cases	(n = 33))

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Age	Male	%	Female	%
0-14	1	3.03	0	0
15–25	3	9.09	1	3.03
26-35	5	15.2	4	12.2
36-45	8	24.2	2	6.06
46-55	1	3.03	3	9.09
> 55	5	15.2	0	0
Total	23	69.7	10	30.3

Table 3 Matrix showing the result of testing method of each positive cases with % positivity among the 3 stool samples of each case (n = 33)

Sl. no.	Microscopy	ELISA	PCR
1	+ (100%)	+ (100%)	+ (100%)
2	+ (100%)	+ (100%)	+ (100%)
3	+ (100%)	+ (100%)	+ (100%)
4	+ (66.6%)	-	+ (66.6%)
5	+ (66.6%)	+ (66.6%)	+ (66.6%)
6	-	+ (66.6%)	+ (66.6%)
7	-	+ (66.6%)	-
8	-	-	+ (66.6%)
9	+ (100%)	+ (100%)	+ (100%)
10	+ (100%)	+ (100%)	+ (100%)
11	+ (100%)	+ (100%)	+ (100%)
12	+ (66.6%)	-	+ (66.6%)
13	+ (100%)	+ (100%)	+ (100%)
14	-	+ (66.6%)	-
15	-	+ (66.6%)	+ (66.6%)
16	-	-	+ (66.6%)
17	+ (100%)	+ (100%)	+ (100%)
18	-	+ (66.6%)	-
19	+ (100%)	+ (100%)	+ (100%)
20	+ (100%)	+ (100%)	+ (100%)
21	+ (100%)	+ (100%)	+ (100%)
22	+ (100%)	+ (100%)	+ (100%)
23	+ (100%)	+ (100%)	+ (100%)
24	+ (100%)	+ (100%)	+ (100%)
25	+ (100%)	+ (100%)	+ (100%)
26	-	+ (66.6%)	+ (100%)
27	-	+ (66.6%)	+ (100%)
28	-	+ (100%)	+ (100%)
29	+ (100%)	+ (100%)	+ (100%)
30	+ (100%)	+ (100%)	+ (100%)
31	-	+ (66.6%)	+ (66.6%)
32	-	+ (100%)	+ (100%)
33	-	+ (100%)	+ (100%)

Abbreviations: ELISA, enzyme-linked immunosorbent assay; PCR, polymerase chain reaction.

Table 4 Agreement between gold standard (microscopy) andELISA

ELISA	Microscopy		Карра	
	Positive	Negative	(95% CI)	
Positive	18	11	0.79 (0.72–0.87)	
Negative	2	600		
Total	20	611		

Abbreviations: CI, confidence interval; ELISA, enzyme-linked immunosorbent assay.

Table 5 Validity indicators of ELISA against microscopy

Parameter	Estimate (95% CI)
Sensitivity	90
Specificity	98.2
PPV	62.07
NPV	99.67
Likelihood ratio positive test	49.99
Likelihood ratio negative test	0.1018
Diagnostic accuracy	97.94

Abbreviations: CI, confidence interval; NPV, negative predictive value; PPV, positive predictive value.

Table 6 Agreement between gold standard (microscopy) and PCR

PCR	Microscopy		Карра
	Positive	Negative	(95% CI)
Positive	20	10	0.72 (0.65–0.80)
Negative	0	601	
Total	20	611	

Abbreviations: CI, confidence interval; PCR, polymerase chain reaction.

Table 7 Validity indicators of PCR against microscopy

Parameter	Estimate (95% CI)
Sensitivity	100
Specificity	98.36
PPV	66.67
NPV	100
Likelihood ratio positive test	61.1
Likelihood ratio negative test	0.0
Diagnostic accuracy	98.42

Abbreviations: CI, confidence interval; NPV, negative predictive value; PCR, polymerase chain reaction; PPV, positive predictive value.

positive cases even in the asymptomatic cases and widespread use of this technique has allowed for revising the epidemiology of the true pathogenic *E. histolytica*, thus eliminating the need for unnecessary treatment.^{21,22} These tests may be helpful from a diagnostic perspective in developed nations, where infections due to *E. histolytica* are not very commonly seen. The coproantigen detection ELISA is more useful as an adjunct to microscopic diagnosis in detecting intestinal parasites, than antibody detection ELISA. Recent studies reveal regarding the high sensitivity and specificity of fecal antigen assay with the use of polyclonal or monoclonal antibodies that can successfully detect Entamoeba infections, which are negative by stool microscopy.^{23,24} These antigenbased detection tests have a sensitivity approaching that of stool culture.² A good sensitivity and specificity have been observed for tests detecting of E. histolytica antigen in stool specimens, especially in patients with amoebic colitis and asymptomatic cases.²⁵ Importantly, among the routinely used diagnostic methods, that is, antigen detection, antibody detection, microscopy, and isoenzyme analysis, PCR., only coproantigen detection using ELISA is technically simple to perform and can be used in resource poor settings that do not have molecular facilities, thus making it appropriate for use in the developing world, where amoebiasis is more commonly seen.²⁶ The only drawback of this coproantigen testing is that it can be performed only on unpreserved fresh or frozen stool samples, since fecal antigens get destroyed by fixative agents.²⁷ Also, sensitivity of coproantigen tests is lower than molecular methods of detection.²⁸

Due to the many limitations of the conventional techniques, molecular methods of detection are emerging as a handy tool in the diagnosis of various infections including amoebiasis. The accurate identification of pathogenic E. histolytica is essential in the treatment of the affected patients as well as studying the epidemiology of amoebiasis outbreaks. The World Health Organization also recommends the application of PCR technology for the accurate detection and differentiation of E. histolytica from other commensal amoebae, directly from stool samples.² Blessmann et al developed a closed tube real-time PCR to detect E. histolytica directly from stool samples. They concluded in their study that the two sets of primers are actually species specific and that PCR is not influenced by the presence of considerable amounts of other Entamoeba species.²⁹ In the current study, the nested multiplex PCR technique was used for the detection of E. histolytica directly from stool sample, by the method given by Khairnar and Parija.¹⁰ This nested multiplex PCR detected E. histolytica, E. dispar, and E. moshkovskii DNA, even at the minimum parasite concentration tested (1,000 parasites/0.05 g of feces). The detection limit of this PCR for E. histolytica, E. dispar, and E. moshkovskii was approximately 25 Entamoeba protozoa cells. In the present study, it was observed that the nested multiplex PCR was positive in 30 (4.75%) cases and negative in 601 (95.25%) cases for *E*. histolytica. The possible causes of PCR negative samples among the serology positive cases might be attributable to very low parasite density in stool or an extraintestinal infection. The validity indicators of PCR against microscopy showed a sensitivity of 100%, specificity of 98.36%, PPV of 66.67%, NPV of 100%, and a diagnostic accuracy of 98.42%. This is similar to the study findings by Blessmann et al.²⁹ In comparison to microscopy or coproserology, PCR detected slightly higher number of positive samples, suggesting that PCR is more sensitive than either of those techniques. The study by Weitzel et al study suggests that PCR testing would

be a great aid in the laboratory diagnosis of amoebiasis, which will surpass the limitations of conventional modes of diagnosis of this parasitic disease.³⁰ In endemic setups, which usually the developing countries are, it is advisable to perform all the three tests simultaneously, so as to diagnose all the cases and treat accordingly. But with respect to cost-effectiveness, microscopy and serology can be performed in all suspected cases and PCR can be done only for cases with strong clinical suspicion despite a negative microscopy and serology result.

Conclusion

This study throws substantial light on the diagnostic advantages of PCR over the conventional coproantigen ELISA-based kits and stool microscopy, in both sensitivity and specificity. In addition, the nested multiplex PCR has the advantage of specifically targeting and detecting *E. histolytica*, *E. dispar*, and *E. moshkovskii* in clinical stool samples. This study aimed to discuss different methods that exist for the laboratory identification of *E. histolytica*. After statistical analysis of the observations from the present study, it is thus concluded that in all cases of clinically suspected amoebiasis, preferably a combination of stool microscopy, stool coproantigen testing followed up by molecular detection of the parasite DNA from stool samples offers the best and robust approach to laboratory diagnosis of *E. histolytica*.

Informed Consent

Written informed consent was taken from all adult participants and parents of minor participants, who agreed to be enrolled in the study group, after getting approval from the JIPMER Ethical Committee, JIPMER.

Authors' Contribution

Sindhusuta Das was responsible for sample and data collection, and its processing including microscopy, coproELISA, and PCR. She was responsible for writing the manuscript and its submission. Nonika Rajkumari was responsible for the conceptualization of the study, counterchecking the results, and monitoring of the work performed and analysis of the results. She also checked and refined the manuscript and gave critical inputs to the paper. Anitha Gunalan helped in data collection and initial data processing. Dhanalakshmi Rajavelu helped in data collection and the figure refinement. Jeby Jose Olickal helped in the statistical modeling of the study selection of the study parameters as well as gave critical inputs to the study and helped in refining the final manuscript.

Conflict of Interest

The authors declare no conflicts of interest.

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