Molecular Identification of Entamoeba histolytica Parasite by Using Actin and Amebapore-A Genes التشخيص الجزيئي لطفيلي Entamoeba histolytica بأستخدام المورثين Amebapore-A & Actin

Assist. Prof. Dr. Jameel Jerri Yousif Al-Hameedawi, Parasite Immunity / Education Faculty for Girls- Kufa University Jameel education@yahoo.com

الهدف: أجريت الدراسة الحالية في محافظة النجف للفترة من نيسان 2013 ولغاية تشرين الثاني 2013 تهدف الى التعرف على طفيلي الأميبا الحالة للنسيج Entamoeba histolytica ل التحري عن جينات

تقنية التفاعل التسلسلي لأنزيم البلمرة Real-Time Amebapore-A Actin في عينات البراز .PCR (104) عينة براز من المرضى الذين يعانون من الإسهال الوافدين الى المنهجية: مستشفى مدينة الصدر الطبية ومستشفى الزهراء للولا النتائج: كشفت نتائج فحص التفاعل التسلسلي لأنزيم البلمرة Real-Time PCR ، وجود طغيلي الأميبا الحالة للنسيج في (25) عينة براز من أصل (104) عينة (24%). الأستنتاجات: على ضوء النتائج التي تم الحصول عليها من خلال هذه الدراسة يستنتج بأن لتقنية Real-Time PCR كفاءة كبيرة ً وسريعة في تشخيُّص طفيلي الأميبا الحالة للنسيج في عينات البراز . التوصيات: دراسة التعبير الجيني Gene Expression لجينات Amebapore-A Actin النوعية بطغيلي الأميبا الحالة للنسيج

الخلاصة

Abstract

Objective: The present study was conducted in the province of Al- Najaf, for the period from April 2013 until November 2013, which aims to identify the Entamoeba histolytica parasites by screening for Actin and Amebapore – A genes in stool samples by using the real-time PCR technique.

Methodology: The study included, collect (104) stool samples from patients with diarrhea arrivals to Al-Sadar Medical City Hospital ,Al- Zahra Maternity and Children Hospital, Al-Manathera hospital and Al-Sajjad Hospital.

Results: The results of the real-time PCR revealed, the presence of *E. histolytica* parasites in ((25) stool samples out of the (104) samples, the rate (24%).

Conclusion: In light of the results obtained through this study we concludes that the efficiency of real-time -PCR technique large and rapid in diagnosis of E. histolytica parasites in stool samples.

Recommendation: The study of gene expression of *Actin* and *Amebapore – A* genes, specific for E. histolytica parasite.

Keywords: Real-Time PCR, Entamoeba histolytica, Actin, Amebapore – A

INTRODUCTION

Amebiasis caused by *Entamoeba histolytica* is still mentioned as one of the major health problems in tropical and subtropical areas (1). It is the cause of various infectious diseases ranging from dysentery to abscess of liver or other organs.

It is estimated that amebiasis is responsible for up to 110,000 deaths per year (2, 3). This infection is usually predominant in low socioeconomic status and poor hygienic situations that favor the indirect fecal-oral transmission of the infection (4). Previously two morphologically identical species of *Entamoeba* had been found, and was shown that only one of them is able to cause infection in kittens or human volunteers (5). However, *E. histolytica* has recently been re-described as two distinct species; the pathogenic species *E. histolytica* and the nonpathogenic species *E. dispar*. As these two species are morphologically similar, development of new methods for their rapid differentiation is currently under investigation (6).

Actin is a cytoskeletal protein involved in cellular functions like maintaining cell morphology, cell motility and division, and intracellular transport (7). *Actin* proteins are encoded by a multigene family. In *Entamoeba histolytica*, two to four *Actin* genes have been described (8).

One of the remarkable capacities of *Entamoeba histolytica* trophozoites is their ability to lyse eukaryotic cells on contact. This ability derives from *Amoebapores*, a family of three pore-forming peptides (*Amoebapore A* [*AP-A*], *AP-B*, and *AP-C*) (9, 10). *Amoebapores* insert into the membranes of bacteria or eukaryotic cells and form pores that result in lysis of the target cells. The addition of purified *Amoebapores* to eukaryotic cells results in cell necrosis and possibly apoptosis (11). Amplification of ameba DNA fragments by PCR has been proven to constitute a sensitive and specific method to detect *E. histolytica* or *E. dispar* from human feces (11, 12). The present study was aimed to identification of *E.histolytica* parasite by detection and characterization of *Actin* and *Amebapore-A* (*AP-A*) genes in stool samples by using real-time PCR techniques.

MATERIALS AND METHODS

The present study was conducted for the period from April 2013 until November 2013, where he was collecting (104) stool samples from patients suffering from diarrhea, which is between the ages of (1-58) years who arrivals to some hospitals of Al-Najaf province (Al-Sadar Medical City Hospital, Al- Zahra Maternity and Children Hospital, Al-Manathera hospital and Al-Sajjad Hospital), where placed the samples in sterile plastic containers and record them some necessary informations such as the patient's name, age, residence, general appearance of the stool, and the presence of blood, and the presence of mucus, and preserved the samples in the temperature (-20°C) until used in the DNA extraction process **.**

Extraction of DNA from stool samples

DNA was extracted from stool samples according to (Bioneer Kit, Korea). Agarose gel electrophoresis was carried out to detect the presence of DNA (13).

Amplification by real-time PCR

Table (1):show the sequence of primers, which used for amplification of *Actin* and *Ap-A* genes for detection of *E. histolytica*, whereas the real-time PCR mixture was prepared as explained in table (2).

Table (1): Primers used for real-time PCR

Primer	DNA Sequence	Reference	

AP-A	F	5'CACTAAGGGAGCTGATAAAGTAAAAGATTA-3'	(14)	
	R	5'-TCCAAAATCAAGAACTTTAGTGCAA-3'		
Actin	F	5'TGTAGATAATGGATCAGGAATGTGTAAA3'		
	R	5'-CAATGGATGGGAATACAGCTCTT-3'		

Table (2):The mixture of real – time PCR

Content	Volume (µ l)
master mix with Sybr green	20
F- Primer (10 pmole)	2
R- Primer (10 pmole)	2
DNA Template	10
DEPC-distlled water	16
Total	50

Real-Time PCR protocol was carried out as following :denaturation at 95 0 C for 5 minute and 45 cycle of 95 0 C for 20 second , annealing at 55 0 C for 40 second for *Actin* and 60 0 C for 40 second for *AP-A*.Melting at 50-95 0 C for 45 minute.

Binding of Sybr green to the real-time PCR product

Syber green is fluorescent dye and when it bind to double strand DNA, the fluorescence emission occurs which is used for the visualization of amplified product during Real-Time PCR of *Actin* and *Ap-A* genes with Sybr green. The amplification of each DNA specimen was determined by observation the fluorescence emission curve. These curves were produced due to binding Sybr green to the real-time PCR products, and the fluorescence reading were taken after each extension step of real-time PCR. The fluorescence emission increases due to the increase cycles of real-time PCR (15).

MELTING CURVE ANALYSIS

After completion of (45) cycles PCR amplification, the PCR products were melted by raising the temperature from (50 C°) to (95 C°) at rate (1C°/min.) .The Exicycler thermal block soft ware displayed the data collection during melting curve analysis as result melting temperature were derived from melting peaks by melting curve analysis of the amplified DNA specimens. Melting curve is of great importance, because the Sybr green will detect any double stranded DNA including primer dimers, contaminating DNA, and PCR product from misannealed primer. By viewing a dissociation curve, you ensure that the desired amplicon was detected (16).

RESULTS

The results of the real- time PCR showed that the amplified DNA for the genes Actin and AP-A in (25) stool samples out of (104) samples with the percentage (24%) by using specific primers and can be determined by a serving the fluorescence emission curves , the fluorescence curve which exceed the threshold line represent the amplified Actin and AP-A genes of E. *histolytica*, as shown in fig. (1) and (2). The negative samples did not amplified and the fluorescence curves remained below the threshold line as seen in fig. (3).

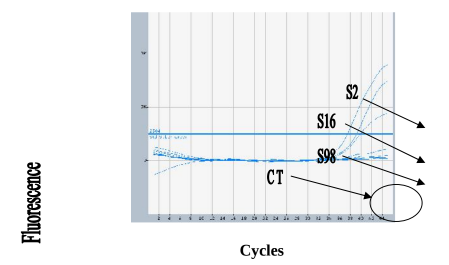
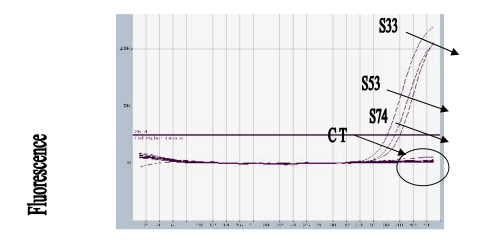


Fig.(1) Amplification of *Actin* gene of *E. histolytica* with Sybr green by using real –time PCR.

S= specimen, CT= Cycle Threshold CT for S2=38.7 ; CT for S16 = 37.8 ; CT for S98 = 39.9



Cycles

Fig.(2) : Amplification of *AP-A* gene of *E. histolytica* with Sybr green by using real- time PCR. S= specimen, CT= Cycle Threshold CT for S33 = 38.1 ; CT for S53= 39.3 ; CT for S74 = 40.0

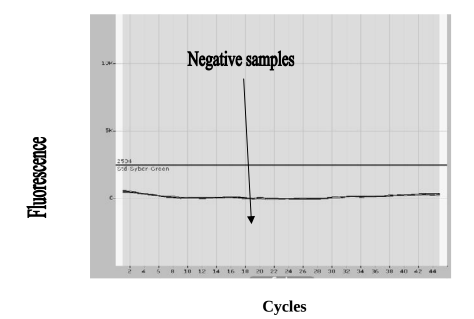
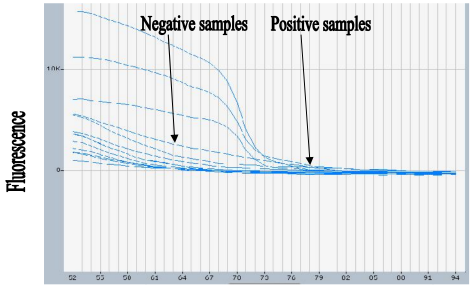


Fig.(3) Not amplified of *Actin & AP-A* genes of *E. histolytica* with Sybr green by using real –time PCR.

Melting curve

Melting curve analysis after completing the PCR. The temperature is then gradually increased and the fluorescence (syber green) is measured as function of temperature. The fluorescence decreases gradually with increasing temperature because of increased thermal motion which allows for more internal rotation in the bound dye. However, when the temperature is reached at which the double stranded DNA separates the dye comes off and the fluorescence drops abruptly this temperature, referred to as the melting temperature, as shown in fig.(4) &(5).



Temperature

Fig.(4) The melting curve product for *Actin* gene

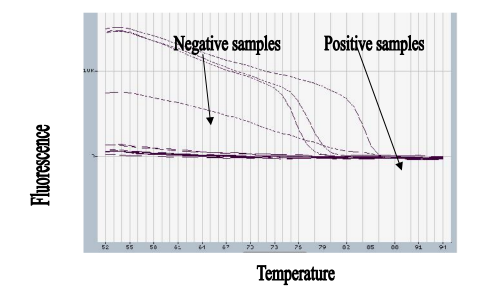


Fig.(5) The melting curve product for AP-A gene

DISCUSSION

Diagnosis of amoebiasis using molecular methods is useful not only in terms of diagnosis and but also for epidemiological studies through removing possible microscopy mistakes (17, 18). Currently, *E. histolytica* antigen-specific ELISA or DNA determination (Real-Time PCR is more sensitive) tests are considered to be the most scientific alternatives for definitive diagnosis (19, 20).

The results of the present study revealed that the amplified DNA by real-time PCR for Actin and Amebapore-A genes of E.histolytica in (25) samples out of (104) samples . The current study reinforced by (21), which found that the Actin, which occurs in the *E.histolytica* parasite encodes by two genes. DNA microarray screening also showed the upregulation of several genes encoding for proteins involved in Actin cytoskeleton dynamics during chemotaxis of E.histolytica towards Tumor Necrosis Factor (TNF) . These included proteins participating in microfilament dynamics, such as nucleation (Arp2/3, Formins), F-actin capping Cap34), microfilament network formation (ABP-120 (Cap34 . (filamin), cortexellin), Actin bundling array (actinin) (22)). Also the results of PCR showed The E. dispar Actin intergenic sequences differed in 83 nucleotides (22%) from those of E. histolytica (23). There is abundant evidence that the actin cytoskeleton of the ameba is vital for adherence to target cells, cytotoxicity and phagocytosis process (24 , 25). On the other hand Amebapore genes are important in the detection of E. histolytica parasite and in turn these genes encode Amebapore enzymes which is of great importance in the pathogenesis of the parasite (26). The present study supported by (27), which found that *E. histolytica Amoebapores* exist as mature and potentially active peptides inside cytoplasmic granules of the trophozoite. The mode of action of the Amoebapore in viable E. histolytica trophozoites was that following the lectinmediated recognition and intimate adherence between the amoeba and its target cell, the Amoebapore molecules are inserted into the membrane without depending on the interaction with a specific membrane receptor. As well as Amoebapore have antibacterial, cytotoxic and pore-forming activities (28) . Also Amoebapore expression is required for full virulence in the mouse model of amebic liver abscess, but E. histolytica trophozoites that do not express Amoebapore-A can still cause inflammation and tissue damage in infected human colonic xenografts (14).

CONCLUSION of the current study, it can be used *Actin* and *Amoebapore-A* genes in the detection of *E. histolytica* parasite by using real-time PCR.

RECOMMENDATION to study of the Gene Expression of *Actin* and *Amebapore* – A genes, specific for *E. histolytica* parasite.

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